
Spectre[®] Circuit Simulator Reference

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Preface

This manual assumes that you are familiar with the development, design, and simulation of integrated circuits and that you have some familiarity with SPICE simulation. It contains information about the Spectre[®] circuit simulator.

Spectre is an advanced circuit simulator that simulates analog and digital circuits at the differential equation level. The simulator uses improved algorithms that offer increased simulation speed and greatly improved convergence characteristics over SPICE. Besides the basic capabilities, the Spectre circuit simulator provides significant additional capabilities over SPICE. SpectreHDL (Spectre High-Level Description Language) and Verilog[®]-A use functional description text files (modules) to model the behavior of electrical circuits and other systems. SpectreRF adds several new analyses that support the efficient calculation of the operating point, transfer function, noise, and distortion of common RF and communication circuits, such as mixers, oscillators, sample holds, and switched-capacitor filters.

This preface discusses the following topics:

- [Related Documents](#) on page Preface-8
- [Typographic and Syntax Conventions](#) on page Preface-8
- [References](#) on page Preface-9

Related Documents

The following can give you more information about the Spectre circuit simulator and related products:

- To learn more about the equations used in the Spectre circuit simulator, consult the [Spectre Circuit Simulator Device Model Equations](#) manual.
- The Spectre circuit simulator is often run within the Cadence[®] analog circuit design environment, under the Cadence[®] design framework II. To see how the Spectre circuit simulator is run under the analog circuit design environment, read the [Cadence Analog Design Environment User Guide](#).
- For more information about using the Spectre circuit simulator with SpectreHDL, see the [SpectreHDL Reference](#) manual.
- For more information about using the Spectre circuit simulator with Verilog-A, see the [Verilog-A Language Reference](#) manual.
- If you want to see how SpectreRF is run under the analog circuit design environment, read [SpectreRF Help](#).
- For more information about RF theory, see [SpectreRF Theory](#).
- For more information about how you work with the design framework II interface, see [Design Framework II Help](#).
- For more information about specific applications of Spectre analyses, see *The Designer's Guide to SPICE & Spectre*¹.

Typographic and Syntax Conventions

This list describes the syntax conventions used for the Spectre circuit simulator.

literal Nonitalic words indicate keywords that you must enter literally. These keywords represent command (function, routine) or option names, file names and paths, and any other sort of type-in commands.

argument Words in italics indicate user-defined arguments for which you must substitute a name or a value. (The characters before the underscore () in the word indicate the data types that this argument can take. Names are case sensitive.

1. Kundert, Kenneth S. *The Designer's Guide to SPICE & Spectre*. Boston: Kluwer Academic Publishers, 1995.

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Preface

|Vertical bars (OR-bars) separate possible choices for a single argument. They take precedence over any other character.

[] Brackets denote optional arguments. When used with OR-bars, they enclose a list of choices. You can choose one argument from the list.

{ } Braces are used with OR-bars and enclose a list of choices. You must choose one argument from the list.

. . . Three dots (...) indicate that you can repeat the previous argument. If you use them with brackets, you can specify zero or more arguments. If they are used without brackets, you must specify at least one argument, but you can specify more.

Important

The language requires many characters not included in the preceding list. You must enter required characters exactly as shown.

References

Text within brackets ([]) are references. See [Appendix A, "References,"](#) for more detailed information.

Introducing the Spectre Circuit Simulator

This chapter discusses the following:

- [Improvements over SPICE](#) on page 10
- [Analog HDLs](#) on page 14
- [RF Capabilities](#) on page 15
- [Mixed-Signal Simulation](#) on page 17
- [Environments](#) on page 17

The Spectre[®] circuit simulator is a modern circuit simulator that uses direct methods to simulate analog and digital circuits at the differential equation level. The basic capabilities of the Spectre circuit simulator are similar in function and application to SPICE, but the Spectre circuit simulator is not descended from SPICE. The Spectre and SPICE simulators use the same basic algorithms—such as implicit integration methods, Newton-Raphson, and direct matrix solution—but every algorithm is newly implemented. Spectre algorithms, the best currently available, give you an improved simulator that is faster, more accurate, more reliable, and more flexible than previous SPICE-like simulators.

Improvements over SPICE

The Spectre circuit simulator has many improvements over SPICE.

Improved Capacity

The Spectre circuit simulator can simulate larger circuits than other simulators because its convergence algorithms are effective with large circuits, because it is fast, and because it is frugal with memory and uses dynamic memory allocation. For large circuits, the Spectre circuit simulator typically uses less than half as much memory as SPICE.

Improved Accuracy

Improved component models and core simulator algorithms make the Spectre circuit simulator more accurate than other simulators. These features improve Spectre accuracy:

- Advanced metal oxide semiconductor (MOS) and bipolar models

- The Spectre BSIM 3v3 is a physics-based metal-oxide semiconductor field effect transistor (MOSFET) model for simulating analog circuits.
- The Spectre models include the MOS0 model, which is even simpler and faster than MOS1 for simulating noncritical MOS transistors in logic circuits and behavioral models, MOS 9, EKV, BTA-HVMOS, BTA-SOI, VBIC95, TOM2, and HBT.

- Charge-conserving models

The capacitance-based nonlinear MOS capacitor models used in many SPICE derivatives can create or destroy small amounts of charge on every time step. The Spectre circuit simulator avoids this problem because all Spectre models are charge-conserving.

- Improved Fourier analyzer

The Spectre circuit simulator includes a two-channel Fourier analyzer that is similar in application to the SPICE `.FOURIER` statement but is more accurate. The Spectre simulator's Fourier analyzer has greater resolution for measuring small distortion products on a large sinusoidal signal. Resolution is normally greater than 120 dB. Furthermore, the Spectre simulator's Fourier analyzer is not subject to aliasing, a common error in Fourier analysis. As a result, the Spectre simulator can accurately compute the Fourier coefficients of highly discontinuous waveforms.

- Better control of numerical error

Many algorithms in the Spectre circuit simulator are superior to their SPICE counterparts in avoiding known sources of numerical error. The Spectre circuit simulator improves the control of local truncation error in the transient analysis by controlling error in the voltage rather than the charge.

In addition, the Spectre circuit simulator directly checks Kirchhoff's Current Law (also known as Kirchhoff's Flow Law) at each time step, improves the charge-conservation accuracy of the Spectre circuit simulator, and eliminates the possibility of false convergence.

- Superior time-step control algorithm

The Spectre circuit simulator provides an adaptive time-step control algorithm that reliably follows rapid changes in the solution waveforms. It does so without limiting assumptions about the type of circuit or the magnitude of the signals.

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- More accurate simulation techniques

Techniques that reduce reliability or accuracy, such as device bypass, simplified models, or relaxation methods, are not used in the Spectre circuit simulator.

- User control of accuracy tolerances

For some simulations, you might want to sacrifice some degree of accuracy to improve the simulation speed. For other simulations, you might accept a slower simulation to achieve greater accuracy. With the Spectre circuit simulator, you can make such adjustments easily by setting a single parameter.

Improved Speed

The Spectre circuit simulator is designed to improve simulation speed. The Spectre circuit simulator improves speed by increasing the efficiency of the simulator rather than by sacrificing accuracy.

- Faster simulation of small circuits

The average Spectre simulation time for small circuits is typically two to three times faster than SPICE. The Spectre circuit simulator can be over 10 times faster than SPICE when SPICE is hampered by discontinuity in the models or problems in the code. Occasionally, the Spectre circuit simulator is slower when it finds ringing or oscillation that goes unnoticed by SPICE. This can be improved by setting the `macromodels` option to `yes`.

- Faster simulation for large circuits

The Spectre circuit simulator is generally two to five times faster than SPICE with large circuits because it has fewer convergence difficulties and because it rapidly factors and solves large sparse matrices.

Improved Reliability

The Spectre circuit simulator offers you the following improvements in reliability:

- Improved convergence

Spectre proprietary algorithms ensure convergence of the Newton-Raphson algorithm in the DC analysis. The Spectre circuit simulator virtually eliminates the convergence problems that earlier simulators had with transient simulation.

- Helpful error and warning messages

The Spectre circuit simulator detects and notifies you of many conditions that are likely to be errors. For example, the Spectre circuit simulator warns of models used in

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forbidden operating regions, of incorrectly wired circuits, and of erroneous component parameter values. By identifying such common errors, the Spectre circuit simulator saves you the time required to find these errors with other simulators.

The Spectre circuit simulator lets you define soft parameter limits and sends you warnings if parameters exceed these limits.

- **Thorough testing**

Automated tests, which include over 1,000 test circuits, are constantly run on all hardware platforms to ensure that the Spectre circuit simulator is consistently reliable and accurate.

- **Benchmark suite**

There is an independent collection of SPICE netlists that are difficult to simulate. You can obtain these circuits from the Microelectronics Center of North Carolina (MCNC) if you have File Transfer Protocol (FTP) access on the Internet. You can also get information about the performance of several simulators with these circuits.

The Spectre circuit simulator has successfully simulated all of these circuits. Sometimes the netlists required minor syntax corrections, such as inserting balance parentheses, but circuits were never altered, and options were never changed to affect convergence.

Improved Models

The Spectre circuit simulator has MOSFET Level 0–3, BSIM1, BSIM2, BSIM3, BSIM 3v3, EKV, MOS9, JFET, TOM2, GaAs MESFET, BJT, VBIC, HBT, diode, and many other models. It also includes the temperature effects, noise, and MOSFET intrinsic capacitance models.

The Spectre Compiled Model Interface (CMI) option lets you integrate new devices into the Spectre simulator using a very powerful, efficient, and flexible C language interface. This CMI option, the same one used by Spectre developers, lets you install proprietary models.

Spectre Usability Features and Customer Service

The following features and services help you use the Spectre circuit simulator easily and efficiently:

- You can use Spectre soft limits to catch errors created by typing mistakes.
- Spectre diagnosis mode, available as an options statement parameter, gives you information to help diagnose convergence problems.

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- You can run the Spectre circuit simulator standalone or run it under the Cadence analog design environment. To see how the Spectre circuit simulator is run under the analog design environment, read the [Cadence Analog Design Environment User Guide](#). You can also run the Spectre circuit simulator in the Composer-to-Spectre direct simulation environment. The environment provides a graphical user interface for running the simulation.
- The Spectre circuit simulator gives you an online help system. With this system, you can find information about any parameter associated with any Spectre component or analysis. You can also find articles on other topics that are important to use the Spectre circuit simulator effectively.
- The Spectre circuit simulator also includes a waveform display tool, Analog Waveform Display (AWD), to use to display simulation results. For more information about AWD, see the [Analog Waveform User Guide](#).
- If you experience a stubborn convergence or accuracy problem, you can send the circuit to Customer Support to get help with the simulation. For current phone numbers and e-mail addresses, see the following web site: <http://sourcelink.cadence.com/supportcontacts.html>.

Analog HDLs

The Spectre circuit simulator works with two analog high-level description languages (AHDLS): SpectreHDL and Verilog[®]-A. These languages are part of the Spectre Verilog-A Simulation option. SpectreHDL is proprietary to Cadence and is provided for backward compatibility. The Verilog-A language is an open standard, which was based upon SpectreHDL. The Verilog-A language is preferred because it is upward compatible with Verilog-AMS, a powerful and industry-standard mixed-signal language.

Both languages use functional description text files (modules) to model the behavior of electrical circuits and other systems. Each programming language allows you to create your own models by simply writing down the equations. The AHDL lets you describe models in a simple and natural manner. This is a higher level modeling language than previous modeling languages, and you can use it without being concerned about the complexities of the simulator or the simulator algorithms. In addition, you can combine AHDL components with Spectre built-in primitives.

Both languages let designers of analog systems and integrated circuits create and use modules that encapsulate high-level behavioral descriptions of systems and components. The behavior of each module is described mathematically in terms of its terminals and external parameters applied to the module. Designers can use these behavioral descriptions in many disciplines (electrical, mechanical, optical, and so on).

Both languages borrow many constructs from Verilog and the C programming language. These features are combined with a minimum number of special constructs for behavioral simulation. These high-level constructs make it easier for designers to use a high-level description language for the first time.

RF Capabilities

SpectreRF adds several new analyses that support the efficient calculation of the operating point, transfer function, noise, and distortion of common analog and RF communication circuits, such as mixers, oscillators, sample and holds, and switched-capacitor filters.

SpectreRF adds four types of analyses to the Spectre simulator. The first is periodic steady-state (PSS) analysis, a large-signal analysis that directly computes the periodic steady-state response of a circuit. With PSS, simulation times are independent of the time constants of the circuit, so PSS can quickly compute the steady-state response of circuits with long time constants, such as high-Q filters and oscillators.

You can also embed a PSS analysis in a sweep loop (referred to as an SPSS analysis in the Cadence analog design environment), which allows you to easily determine harmonic levels as a function of input level or frequency, making it easy to measure compression points, intercept points, and voltage-controlled oscillator (VCO) linearity.

The second new type of analysis is the periodic small-signal analysis. After completing a PSS analysis, SpectreRF can predict the small-signal transfer functions and noise of frequency translation circuits, such as mixers or periodically driven circuits such as oscillators or switched-capacitor or switched-current filters. The periodic small-signal analyses—periodic AC (PAC) analysis, periodic transfer function (PXF) analysis, and periodic noise (Pnoise) analysis—are similar to Spectre's AC, XF, and Noise analyses, but the traditional small-signal analyses are limited to circuits with DC operating points. The periodic small-signal analyses can be applied to circuits with periodic operating points, such as the following:

- Mixers
- VCOs
- Switched-current filters
- Phase/frequency detectors
- Frequency multipliers
- Chopper-stabilized amplifiers
- Oscillators

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- Switched-capacitor filters
- Sample and holds
- Frequency dividers
- Narrow-band active circuits

The third SpectreRF addition to Spectre functionality is periodic distortion (PDISTO) analysis. PDISTO analysis directly computes the steady-state response of a circuit driven with a large periodic signal, such as an LO (local oscillation) or a clock, and one or more tones with moderate level. With PDISTO, you can model periodic distortion and include harmonic effects. PDISTO computes both a large signal, the periodic steady-state response of the circuit, and also the distortion effects of a specified number of moderate signals, including the distortion effects of the number of harmonics that you choose. This is a common scenario when trying to predict the intermodulation distortion of a mixer, amplifier, or a narrow-band filter. In this analysis, the tones can be large enough to create significant distortion, but not so large as to cause the circuit to switch or clip. The frequencies of the tones need not be periodically related to each other or to the large signal LO or clock. Thus, you can make the tone frequencies very close to each other without penalty, which allows efficient computation of intermodulation distortion of even very narrow band circuits.

The fourth analysis that SpectreRF adds to the Spectre circuit simulator is the envelope-following analysis. This analysis computes the envelope response of a circuit. The simulator automatically determines the clock period by looking through all the sources with the specified name. Envelope-following analysis is most efficient for circuits where the modulation bandwidth is orders of magnitude lower than the clock frequency. This is typically the case, for example, in circuits where the clock is the only fast varying signal and other input signals have a spectrum whose frequency range is orders of magnitude lower than the clock frequency. For another example, the down conversion of two closely placed frequencies can also generate a slow-varying modulation envelope. The analysis generates two types of output files, a voltage versus time (td) file, and an amplitude/phase versus time (fd) file for each specified harmonic of the clock fundamental.

In summary, with periodic small-signal analyses, you apply a small signal at a frequency that might not be harmonically related (noncommensurate) to the periodic response of the undriven system, the clock. This small signal is assumed to be small enough so that the circuit is unaffected by its presence.

With PDISTO, you can apply one or two additional signals at frequencies not harmonically related to the large signal, and these signals can be large enough to drive the circuit to behave nonlinearly.

For complex nonlinear circuits, hand calculation of noise or transfer function is virtually impossible. Without SpectreRF, these circuits must be breadboarded to determine their

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performances. The SpectreRF simulator eliminates unnecessary breadboarding, saving time.

Mixed-Signal Simulation

You can use the Spectre circuit simulator coupled with the Verilog[®]-XL simulator in the Cadence analog design environment to simulate mixed analog and digital circuits efficiently. This mixed-signal simulation solution can easily handle complex designs with tens of thousands of transistors and tens of thousands of gates. The digital Verilog data can come from the digital designer as either an RTL block or gates out of synthesis.

Environments

The Spectre circuit simulator is fully integrated into the Cadence[®] design framework II for the Cadence analog design environment and also into the Cadence analog workbench design system. You can also use the Spectre circuit simulator by itself with several different output format options.

Assura[®] interactive verification, Dracula[®] distributed multi-CPU option, and Assura hierarchical physical verification produce a netlist that can be read into the Spectre circuit simulator. However, only interactive verification when used with the Cadence analog design environment automatically attaches the stimulus file. All other situations require a stimulus file as well as device models.

Spectre Command Options

This chapter lists the options you can use with the `spectre` command and gives a brief description of each. It also discusses the following topics:

- [Default Values](#) on page 22
- [Default Parameter Values](#) on page 22

The `spectre` command takes the following syntax at the command line:

```
spectre options inputfile
```

Note: The Spectre[®] circuit simulator reads default values for all the command line arguments marked with a dagger (†) from the UNIX environment variable `%S_DEFAULTS`.

<code>-help</code>	Lists command options and available components and analyses. You can use <code>-h</code> as an abbreviation of <code>-help</code> .
<code>-help name</code>	Gives a synopsis of the device or analysis <i>name</i> . If <i>name</i> is <code>all</code> , the synopses for all components and analyses are given. You can use <code>-h</code> as an abbreviation of <code>-help</code> .
<code>-helpsort name</code>	Gives a synopsis of the device or analysis <i>name</i> and sorts all the parameters by name. You can use <code>-hs</code> as an abbreviation of <code>-helpsort</code> .
<code>-helpfull name</code>	Gives a full synopsis of the component or analysis <i>name</i> , including parameter types and range limits. You can use <code>-hf</code> as an abbreviation of <code>-helpfull</code> .

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Spectre Command Options

<code>-helpsortfull <i>name</i></code>	Gives a full synopsis of component or analysis <i>name</i> , including parameter types and range limits. Sorts all parameters by name. You can use <code>-hsf</code> as an abbreviation of <code>-helpsortfull</code> .
<code>-param†</code>	Does not read the file containing the suggested parameter range limits. You can use <code>-p</code> as an abbreviation of <code>-param</code> .
<code>+param <i>file</i>†</code>	Reads <i>file</i> for the suggested parameter range limits. You can use <code>+p</code> as an abbreviation of <code>+param</code> .
<code>-log†</code>	Does not copy all messages to a file. You can use <code>-l</code> as an abbreviation of <code>-log</code> .
<code>+log <i>file</i>†</code>	Copies all messages to <i>file</i> . You can use <code>+l</code> as an abbreviation of <code>+log</code> .
<code>=log <i>file</i>†</code>	Sends all messages to <i>file</i> . You can use <code>=l</code> as an abbreviation of <code>=log</code> .
<code>-raw <i>raw</i>†</code>	Puts results in a file or directory named <i>raw</i> . In <i>raw</i> , <code>%C</code> is replaced by a circuit name. You can use <code>-r</code> as an abbreviation of <code>-raw</code> .
<code>-format <i>fmt</i>†</code>	Produces raw data in the format <i>fmt</i> . You can use <code>-f</code> as an abbreviation of <code>-format</code> . Possible values for <i>fmt</i> are <code>nutbin</code> , <code>nutascii</code> , <code>wsfbin</code> , <code>wsfascii</code> , <code>psfbin</code> , <code>psfascii</code> , or <code>awb</code> .
<code>+checkpoint†</code>	Turns on the checkpoint capability. You can use <code>+cp</code> as an abbreviation of <code>+checkpoint</code> .
<code>-checkpoint†</code>	Turns off the checkpoint capability. You can use <code>-cp</code> as an abbreviation of <code>-checkpoint</code> .
<code>-recover†</code>	Does not restart the simulation, even if a checkpoint file exists. You can use <code>-rec</code> as an abbreviation of <code>-recover</code> .
<code>+recover†</code>	Restarts the simulation from a checkpoint file, if it exists. You can use <code>+rec</code> as an abbreviation of <code>+recover</code> .

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Spectre Command Options

<code>-cols <i>N</i>†</code>	Sets screen width in characters to <i>N</i> . You can use <code>-c</code> as an abbreviation of <code>-cols</code> . If not set, the Spectre simulator determines the screen width automatically.
<code>-env <i>env</i></code>	Calls the Spectre simulator from the <i>env</i> simulation environment. Possible values for <i>env</i> are <code>artist2</code> , <code>artist4</code> , <code>awb</code> , <code>edge</code> , <code>opus</code> , or <code>solo</code> .
<code>-%<i>X</i></code>	In quoted strings within the netlist, replaces % <i>X</i> with nothing where <i>X</i> is any uppercase or lowercase letter.
<code>+%<i>X string</i>†</code>	In quoted strings within the netlist, replaces % <i>X</i> with <i>string</i> , where <i>X</i> is an uppercase or lowercase letter. You can modify the string by using the <code>:x</code> operators.
<code>+error†</code>	Prints error messages.
<code>-error†</code>	Does not print error messages.
<code>+warn†</code>	Prints warning messages.
<code>-warn†</code>	Does not print warning messages.
<code>+note</code>	Prints notices.
<code>-note</code>	Does not print notices.
<code>+info†</code>	Prints informational messages.
<code>-info†</code>	Does not print informational messages.
<code>+debug†</code>	Prints debugging messages.
<code>-debug†</code>	Does not print debugging messages.
<code>-slave <<i>cmd</i>></code>	Starts the attached simulator using the command <i>cmd</i> .
<code>-slvhost <<i>hostname</i>></code>	Runs the attached simulator on machine <i>hostname</i> . Defaults to local machine.
<code>-V</code>	Prints version information.
<code>-W</code>	Prints subversion information.
<code>-alias <<i>name</i>>†</code>	Gives <i>name</i> to the license manager as the name of the simulator.

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Spectre Command Options

<code>-E†</code>	Runs the C preprocessor on an input file. In SPICE mode, the first line in the file must be a comment.
<code>-D<x>†</code>	Defines string <i>x</i> and runs the C preprocessor.
<code>-D<x=y>†</code>	Defines string <i>x</i> to be <i>y</i> and runs the C preprocessor.
<code>-U<x>†</code>	Undefines string <i>x</i> and runs the C preprocessor.
<code>-I<dir>†</code>	Runs the C preprocessor and searches the directory <i>dir</i> for include files.
<code>-sppt†</code>	Do not run the Spice netlist reader on the input file.
<code>+sppt†</code>	Run the Spice netlist reader on the input file. Use <code>+spp -sppbin</code> on the command line option to read other spp binaries.
<code>-sppbin file†</code>	Specify the path to nondefault spp binary. Default provided.
<code>+sensdata <file></code>	Sends the sensitivity analyses data to <i>file</i> .
<code>-interactive</code>	Run in the noninteractive mode, that is, process the input file and then return. You can use <code>-inter</code> as an abbreviation of <code>-interactive</code> .
<code>+interactive</code>	Run in the default interactive mode. You can use <code>+inter</code> as an abbreviation of <code>+interactive</code> .
<code>+interactive=type</code>	Run in the interactive mode of the type specified. You can use <code>+inter</code> as an abbreviation of <code>+interactive</code> . Possible values for <i>type</i> are <code>skill</code> or <code>mpsc</code> .
<code>+mpsession=sessionName</code>	The <i>sessionName</i> for an interactive session using multiprocess SKILL (MPS). This option is necessary for <code>+interactive=mpsc</code> and implies <code>+interactive=mpsc</code> .
<code>+mpshost=sessionHost</code>	The <i>sessionHost</i> for an interactive session using MPS.

If you do not specify an input file, the Spectre simulator reads from standard input. When `+/-` pairs of `spectre` command options are available, the default is the first value given in the previous list. For further information about the percent code options, `+%` and `-%`, see [Chapter 11, “Managing Files,”](#) in the *Spectre Circuit Simulator User Guide*.

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Spectre Command Options

Note: To remain consistent with the C preprocessor, there is no space between the preprocessor flags (D, U, I) and their arguments. The C preprocessor is available on UNIX systems only and requires that the first line of the file (the SPICE title line) begin with a comment character (* or //).

Default Values

The Spectre simulator reads default values for all the command line arguments marked with a dagger (†) from the UNIX environment variable %S_DEFAULTS. The name of the simulator as called replaces %S. Typically, this name is `spectre`, and the Spectre simulator looks for `spectre_DEFAULTS`. However, the name can be different if you move the executable to a file with a different name or if you call the Spectre simulator through a symbolic or hard link with a different name. This feature lets you set different default values for each name you use to call the Spectre simulator.

If the variable %S_DEFAULTS does not exist, SPECTRE_DEFAULTS is used instead. The command line arguments always override any specifications from the `options` statement in the circuit file. The `options` statement specifications, in turn, override any specifications in the environment variable.

Default Parameter Values

Many Spectre parameters have default values, and sometimes you will need to know them so you can determine whether they are acceptable for your simulation. You can find the default values for component, analysis, and control statement parameters by consulting the documentation for the statement in Spectre online help (`spectre -h`). Values given for parameters in the online help are the default values.

The following examples show you some defaults for different types of parameters from the Spectre online help:

<code>nf=1.0</code>	Forward emission coefficient
<code>etchc=etchm</code>	Narrowing due to etching for capacitances
<code>homotopy=all</code>	Method used when there is no convergence on initial attempt of DC analysis; possible values are <code>none</code> , <code>gmin</code> , <code>source</code> , <code>dptran</code> , <code>ptran</code> , or <code>all</code>
<code>rawfile="%C:r.raw"</code>	Output raw data filename

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Spectre Command Options

In this example, the default values for `nf`, `etchc`, `homotopy`, and `rawfile` are a real number (1.0), the value of a different parameter (`etch`), an enumerated type (`all`), and a character string with a percent code and a colon modifier that gives Spectre instructions for creating the output filename ("`%C:r.raw`").

For more information about percent codes and colon modifiers, see [“Description of Spectre Predefined Percent Codes.”](#) [“Customizing Percent Codes.”](#) and [“Creating Filenames from Parts of Input Filenames”](#) in the *Spectre Circuit Simulator User Guide*.

Component Statements Part I

This chapter discusses the following topics:

- [Analog-to-Logic Converter \(a2d\)](#) on page 25
- [device checker \(assert\)](#) on page 25
- [B3SOI-PD Transistor \(b3soipd\)](#) on page 27
- [Bipolar Junction Transistor \(bjt\)](#) on page 49
- [Lateral PNP Transistor \(bjt301\)](#) on page 62
- [Lateral PNP Transistor \(bjt500\)](#) on page 69
- [Vertical NPN/PNP Transistor \(bjt503\)](#) on page 80
- [Vertical NPN/PNP Transistor \(bjt504\)](#) on page 90
- [Vertical NPN/PNP Transistor \(bjt504t\)](#) on page 108
- [BSIM1 Field Effect Transistor \(bsim1\)](#) on page 126
- [BSIM2 Field Effect Transistor \(bsim2\)](#) on page 147
- [BSIM3 MOS Transistor \(bsim3\)](#) on page 172
- [BSIM3v3 MOS Transistor \(bsim3v3\)](#) on page 186
- [BSIM4 MOS Transistor \(bsim4\)](#) on page 208
- [BSIMSOI-PD/FD Transistor \(bsimsoi\)](#) on page 237
- [BTA SOI Transistor \(btasoi\)](#) on page 259

Analog-to-Logic Converter (a2d)

Description

The analog-to-logic converter transfers analog waveforms to a logic simulator.

This device is not supported within altergroup.

Sample Instance Statement

```
da99 (cmp_out 0) a2d dest="99991" vl=0 vh=5 timex=200u
// 99991 is a digital net in the verilog netlist.
```

Instance Definition

```
Name p n a2d parameter=value ...
```

Instance Parameters

1	dest	The foreign simulator name for the destination of the signal.
2	nestlev=0	Number of nesting levels to ignore in the hierarchical name. This should be used to skip over extra levels that do not exist in the co-simulator.
3	vl=0 V	Voltages below this will be logical 0.
4	vh=5 V	Voltages above this will be logical 1.
5	timex=1 s	Time signal can linger between vl and vh before the state becomes X.

device checker (assert)

Instance Definition

```
Name assert parameter=value ...
```

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Component Statements Part I

Instance Parameters

1	<code>max=∞</code>	Upperbound of the parameter to be checked.
2	<code>min=∞</code>	Lowerbound of the parameter to be checked.
3	<code>duration=0 s</code>	Time period over which the check has been in violation before a message is flagged.
4	<code>mod</code>	Model to be checked.
5	<code>dev</code>	Device or Subcircuit instance to be checked.
6	<code>primitive</code>	Primitive to be checked.
7	<code>sub</code>	Subcircuit master to be checked.
8	<code>param</code>	Any one of input, output, operating point or subcircuit instance parameter to be checked.
9	<code>modelparam</code>	Model parameter to be checked.
10	<code>message</code>	Message to be printed when assertion fails.
11	<code>level=warning</code>	Severity used when assertion fails. Possible values are <code>none</code> , <code>notice</code> , <code>warning</code> , or <code>error</code> .
12	<code>info=no</code>	When <code>info=yes</code> , parameter will not be checked against min/max, only its value will be printed out. Possible values are <code>no</code> or <code>yes</code> .
13	<code>expr</code>	Expression to be checked for violation (in MDL syntax).

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the

Spectre Circuit Simulator Reference

Component Statements Part I

description for that parameter. For example, a reference of M-35 means the 35th model parameter.

dev	I-5	level	I-11	mod	I-4	sub	I-7
duration	I-3	max	I-1	modelparam	I-9		
expr	I-13	message	I-10	param	I-8		
info	I-12	min	I-2	primitive	I-6		

B3SOI-PD Transistor (b3soipd)

Description

B3SOI is an SOI model developed by U.C. Berkeley based on bsim3v3. B3SOI devices require that you use a model statement. This is the B3SOI version-2.2 model. The versions supported are 2.2, 2.22 and 2.23

This device is not supported within altergroup.

Instance Definition

Name d g s e [p] [b] [t] ModelName parameter=value ...

Instance Parameters

1	w (m)	Channel width.
2	l (m)	Channel length.
3	as (m ²)	Area of source diffusion.
4	ad (m ²)	Area of drain diffusion.
5	ps (m)	Perimeter of source diffusion.
6	pd (m)	Perimeter of drain diffusion.
7	nrd (m/m)	Number of squares of drain diffusion.

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Component Statements Part I

8	<code>nrs</code> (m/m)	Number of squares of source diffusion.
9	<code>nrb</code> (m/m)	Number of body squares.
10	<code>m=1</code>	Multiplicity factor (number of MOSFETs in parallel).
11	<code>region=triode</code>	Estimated operating region. Spectre outputs number (0-3) in a rawfile. Possible values are <code>off</code> , <code>triode</code> , <code>sat</code> , or <code>subth</code> .
12	<code>rth0</code> (Ω)	Thermal resistance.
13	<code>cth0</code> (F)	Thermal capacitance.
14	<code>bjtoff=0</code>	BJT off flag.
15	<code>nbc=0</code> m/m	Number of body contact isolation edge.
16	<code>nseg=1</code> m/m	Number of segments for channel width partitioning.
17	<code>pdbcpr=0</code> m	Perimeter length for body contact parasitic at drain.
18	<code>psbcpr=0</code> m	Perimeter length for body contact parasitic at source.
19	<code>agbcpr=0</code> m	Gate to body overlap for body contact parasitic.
20	<code>aebcpr=0</code> m	Gate to body overlap for body contact parasitic.
21	<code>vbsusr=0.0</code> V	Optional initial value of V_{bs} for transient.
22	<code>tnodeout=0</code>	Temperature node flag associated with T node.

Model Definition

`model modelName b3soipd parameter=value ...`

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Component Statements Part I

Model Parameters

Device type parameters

- | | | |
|---|--------------------------|--|
| 1 | <code>type=n</code> | Transistor type.
Possible values are n or p. |
| 2 | <code>version=2.2</code> | Model version selector. The available versions are 2.2, 2.22 and 2.23. |

Threshold voltage parameters

- | | | |
|----|---|--|
| 3 | <code>vtho (V)</code> | Threshold voltage at zero body bias for long-channel devices. For enhancement-mode devices, $v_{tho} > 0$ for n-channel and $v_{th} < 0$ for p-channel. Default value is calculated from other model parameters. |
| 4 | <code>k1=0.5 \sqrt{V}</code> | Body-effect coefficient. |
| 5 | <code>k1w1=0.0 m</code> | First body effect width dependent parameter. |
| 6 | <code>k1w2=0.0 m</code> | Second body effect width dependent parameter. |
| 7 | <code>k2=-0.0186</code> | Charge-sharing parameter. |
| 8 | <code>k3=0</code> | Narrow width coefficient. |
| 9 | <code>k3b=0 1/V</code> | Narrow width coefficient. |
| 10 | <code>w0=2.5e-6 m</code> | Narrow width coefficient. |
| 11 | <code>n1x=1.74e-7 m</code> | Lateral nonuniform doping coefficient. |
| 12 | <code>gamma1 (\sqrt{V})</code> | Body-effect coefficient near the surface. |
| 13 | <code>gamma2 (\sqrt{V})</code> | Body-effect coefficient in the bulk. |
| 14 | <code>vbx (V)</code> | Threshold voltage transition body voltage. |
| 15 | <code>vbm=-3 V</code> | Maximum applied body voltage. |

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16	$dvt0=2.2$	First coefficient of short-channel effects.
17	$dvt1=0.53$	Second coefficient of short-channel effects.
18	$dvt2=-0.032$ 1/V	Body-bias coefficient of short-channel effects.
19	$dvt0w=0$	First coefficient of narrow-width effects.
20	$dvt1w=5.3e6$	Second coefficient of narrow-width effects.
21	$dvt2w=-0.032$ 1/V	Body-bias coefficient of narrow-width effects.
22	$a0=1$	Nonuniform depletion width effect coefficient.
23	$b0=0$ m	Bulk charge coefficient due to narrow width effect.
24	$b1=0$ m	Bulk charge coefficient due to narrow width effect.
25	$a1=0$	No-saturation coefficient.
26	$a2=1$	No-saturation coefficient.
27	$ags=0$ F/m ² V	Gate-bias dependence of abulk.
28	$keta=-0.6$ 1/V	Body-bias coefficient for non-uniform depletion width effect.
29	$ketas=0.0$ V	Surface Potential adjustment for bulk charge effect.

Process parameters

30	$nsub=6e16$ cm ⁻³	Substrate doping concentration.
31	$nch=1.7e17$ cm ⁻³	Peak channel doping concentration.
32	$ngate$ (cm ⁻³)	Poly-gate doping concentration.
33	$xj=0.15e-6$ m	Source/drain junction depth.
34	$lint=0$ m	Lateral diffusion for one side.
35	$wint=0$ m	Width reduction for one side.

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Component Statements Part I

36	$ll=0$ m	Length dependence of delta L.
37	$lln=1$	Length exponent of delta L.
38	$lw=0$ m	Width dependence of delta L.
39	$lwn=1$	Width exponent of delta L.
40	$lwl=0$ m ²	Area dependence of delta L.
41	$wl=0$ m	Length dependence of delta W.
42	$wln=1$	Length exponent of delta W.
43	$ww=0$ m	Width dependence of delta W.
44	$wwn=1$	Width exponent of delta W.
45	$wwl=0$ m ²	Area dependence of delta W.
46	$dwg=0$ m/v	Gate-bias dependence of channel width.
47	$dwb=0$ m/ \sqrt{v}	Body-bias dependence of channel width.
48	$dwbc=0.0$ m	Width offset for body contact isolation edge.
49	$tox=1e-8$ m	Gate oxide thickness.
50	$tbox=3e-7$ m	Buried oxide thickness.
51	$tsi=1e-7$ m	Silicon film thickness.
52	$xt=1.55e-7$ m	Doping depth.
53	$rdsw=100$ Ω μm	Width dependence of drain-source resistance.
54	$prwb=0$ 1/ \sqrt{v}	Body-effect coefficient for Rds.
55	$prwg=0$ 1/V	Gate-effect coefficient for Rds.
56	$wr=1$	Width offset for parasitic resistance.
57	$xl=0$ m	Length variation due to masking and etching.

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Component Statements Part I

- 58 `xw=0 m` Width variation due to masking and etching.
- 59 `binunit=1` Bin parameter unit selector. 1 for microns and 2 for meters.

Mobility parameters

- 60 `mobmod=1` Mobility model selector.
- 61 `u0=670 cm2/V s` Low-field surface mobility at t_{nom} . Default is 250 for PMOS.
- 62 `vsat=8e4 m/s` Carrier saturation velocity at t_{nom} .
- 63 `ua=2.25e-9 m/v` First-order mobility reduction coefficient.
- 64 `ub=5.87e-19 m2/v2` Second-order mobility reduction coefficient.
- 65 `uc=-4.65e-11 m/v2` Body-bias dependence of mobility. Default is -0.046 and unit is 1/V for `mobmod=3`.

Output resistance parameters

- 66 `drout=0.56` DIBL effect on output resistance coefficient.
- 67 `pclm=1.3` Channel length modulation coefficient.
- 68 `pdiblc1=0.39` First coefficient of drain-induced barrier lowering.
- 69 `pdiblc2=8.6e-3` Second coefficient of drain-induced barrier lowering.
- 70 `pdiblc b=0 1/V` Body-effect coefficient for DIBL.
- 71 `pvag=0` Gate dependence of Early voltage.
- 72 `delta=0.01 V` Effective drain voltage smoothing parameter.

Subthreshold parameters

- 73 `cdsc=2.4e-4 F/m2` Source/drain and channel coupling capacitance.

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Component Statements Part I

74	<code>cdscb=0 F/m² V</code>	Body-bias dependence of <code>cdsc</code> .
75	<code>cdscd=0 F/m² V</code>	Drain-bias dependence of <code>cdsc</code> .
76	<code>nfactor=1</code>	Subthreshold swing coefficient.
77	<code>cit=0 F</code>	Interface trap parameter for subthreshold swing.
78	<code>voff=-0.08 V</code>	Threshold voltage offset.
79	<code>dsub=drout</code>	DIBL effect in subthreshold region.
80	<code>eta0=0.08</code>	DIBL coefficient subthreshold region.
81	<code>etab=-0.07 1/V</code>	Body-bias dependence of <code>et0</code> .

Substrate current parameters

82	<code>alpha0=0 m/v</code>	Substrate current impact ionization coefficient.
83	<code>beta0=0 1/V</code>	First V_{ds} dependent parameter of impact ionization current.
84	<code>fbjtii=0.0</code>	Fraction of bipolar current affecting the impact ionization.
85	<code>beta1=0</code>	Second V_{ds} dependent parameter of impact ionization current.
86	<code>beta2=0 V</code>	Third V_{ds} dependent parameter of impact ionization current.
87	<code>vdsatii0=0.9 V</code>	Nominal drain saturation voltage at threshold for impact ionization current.
88	<code>tii=0</code>	Temperature dependent parameter for impact ionization current.
89	<code>lii=0</code>	Channel length dependent parameter at threshold for impact ionization current.
90	<code>esatii=1e7 V/m</code>	Saturation channel electric field for impact ionization current.
91	<code>sii0=0.5 1/V</code>	First V_{gs} dependent parameter for impact ionization current.
92	<code>sii1=0.1 1/V</code>	Second V_{gs} dependent parameter for impact ionization current.

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Component Statements Part I

- 93 `sii2=0.0 1/V` Third V_{gs} dependent parameter for impact ionization current.
- 94 `siid=0 1/V` V_{ds} dependent parameter of drain saturation voltage for impact ionization current.

Parasitic resistance parameters

- 95 `rbsh=0 Ω` Extrinsic body contact sheet resistance.
- 96 `rsh=0 Ω/sqr` Source/drain diffusion sheet resistance.
- 97 `rs=0 Ω` Source resistance.
- 98 `rd=0 Ω` Drain resistance.
- 99 `rbody=0 F` Body resistance.
- 100 `rsc=0 Ω` Source contact resistance.
- 101 `rdc=0 Ω` Drain contact resistance.
- 102 `rss=0 Ω m` Scalable source resistance.
- 103 `rdd=0 Ω m` Scalable drain resistance.
- 104 `hdif=0 m` Length of heavily doped diffusion.
- 105 `ldif=0 m` Lateral diffusion beyond the gate.
- 106 `minr=0.1 Ω` Minimum source/drain resistance.

Junction diode model parameters

- 107 `dskip=yes` Use simple piece-wise linear model for diode currents below $0.1 \cdot i_{abstol}$.
Possible values are `no` or `yes`.
- 108 `imelt=`imaxA`` Explosion current.

Spectre Circuit Simulator Reference

Component Statements Part I

Overlap capacitance parameters

109	<code>cgso</code> (F/m)	Gate-source overlap capacitance.
110	<code>cgdo</code> (F/m)	Gate-drain overlap capacitance.
111	<code>cgeo=0.0</code> F/m	Gate-substrate overlap capacitance.
112	<code>cgbo=2 Dwc Cox</code> F/m	Gate-bulk overlap capacitance..
113	<code>meto=0</code> m	Metal overlap in fringing field.
114	<code>cgs1=0</code> F/m	Gate-source overlap capacitance in LDD region.
115	<code>cgd1=0</code> F/m	Gate-drain overlap capacitance in LDD region.
116	<code>ckappa=0.6</code>	Overlap capacitance fitting parameter.

Junction capacitance model parameters

117	<code>cjswg=cjsw</code> F/m	Zero-bias gate-side junction capacitance density.
118	<code>mjswg=0.5</code>	Gate-side junction grading coefficient.
119	<code>pbswg=0.7</code> V	Gate-side junction built-in potential.
120	<code>tt=1e-12</code> s	Transit time.
121	<code>ndif=1</code>	Power coefficient of channel length dependency for diffusion capacitance.
122	<code>ldif0=1</code>	Power coefficient of channel length dependency for diffusion capacitance.

Charge model selection parameters

123	<code>capmod=2</code>	Intrinsic charge model.
124	<code>dwc=wint</code> m	Delta W for capacitance model.

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125	<code>delvt=0.0</code>	V	Threshold voltage adjustment for C-V.
126	<code>fbody=1.0</code>		Scaling factor for body charge.
127	<code>dlc=lint</code>	m	Delta L for capacitance model.
128	<code>dlcb=lint</code>	m	Length offset fitting parameter for body charge.
129	<code>dlbg=0.0</code>	m	Length offset fitting parameter for backgate charge.
130	<code>clc=1e-8</code>	m	Intrinsic capacitance fitting parameter.
131	<code>cle=0.0</code>		Intrinsic capacitance fitting parameter.
132	<code>cf</code>	(F/m)	Fringe capacitance parameter.
133	<code>vfbcv=-1</code>		Flat-band voltage for capmod=0.
134	<code>xpart=0</code>		Drain/source channel charge partition in saturation for BSIM charge model, use 0.0 for 40/60, 0.5 for 50/50, or 1.0 for 0/100.

Default instance parameters

135	<code>w=5e-6</code>	m	Default channel width.
136	<code>l=5e-6</code>	m	Default channel length.
137	<code>as=0</code>	m ²	Default area of source diffusion.
138	<code>ad=0</code>	m ²	Default area of drain diffusion.
139	<code>ps=0</code>	m	Default perimeter of source diffusion.
140	<code>pd=0</code>	m	Default perimeter of drain diffusion.
141	<code>nrd=0</code>	m/m	Default number of squares of drain diffusion.
142	<code>nrs=0</code>	m/m	Default number of squares of source diffusion.
143	<code>nrb=0</code>	m/m	Default body squares.

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Temperature effects parameters

144	t_{nom} (C)	Parameters measurement temperature. Default set by <code>options</code> .
145	$t_{max}=500$ C	Maximum device temperature above ambient.
146	$shmod=0$	Self-heating selector.
147	$t_{lev}=0$	DC temperature selector.
148	$t_{levc}=0$	AC temperature selector.
149	$eg=1.12452$ V	Energy band gap.
150	$gap1=7.02e-4$ V/C	Band gap temperature coefficient.
151	$gap2=1108$ C	Band gap temperature offset.
152	$kt1=-0.11$ V	Temperature coefficient for threshold voltage.
153	$kt1l=0$ v m	Temperature coefficient for threshold voltage.
154	$kt2=0.022$	Temperature coefficient for threshold voltage.
155	$at=3.3e4$ m/s	Temperature coefficient for v_{sat} .
156	$tcjswg=0$ 1/K	Temperature coefficient of C_{jswg} .
157	$tpbswg=0$ V/K	Temperature coefficient of P_{bswg} .
158	$ua1=4.31e-9$ m/v	Temperature coefficient for u_a .
159	$ub1=-7.61e-18$ m ² /v ²	Temperature coefficient for u_b .
160	$uc1=-5.5e-11$ m/v ²	Temperature coefficient for u_c . Default is -0.056 for <code>mobmod=3</code> .
161	$prt=0$ Ω	Temperature coefficient for R_{ds} .
162	$trs=0$ 1/C	Temperature parameter for source resistance.
163	$trd=0$ 1/C	Temperature parameter for drain resistance.

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164	<code>ute=-1.5</code>	Mobility temperature exponent.
165	<code>dt1=0</code>	First temperature coefficient for tau.
166	<code>dt2=0</code>	Second temperature coefficient for tau.
167	<code>cth0=0 F</code>	Self-heating thermal capacitance.
168	<code>rth0=0 Ω</code>	Self-heating thermal resistance.
169	<code>ntrecf=0</code>	Temperature coefficient of Ntrecf.
170	<code>ntreocr=0</code>	Temperature coefficient of Ntrecr.
171	<code>xbjt=2</code>	BJT current temperature exponent.
172	<code>xdif=2</code>	Diffusion current temperature exponent.
173	<code>xrec=20</code>	Recombination current temperature exponent.
174	<code>xtun=0</code>	Tunneling current temperature exponent.

Noise model parameters

175	<code>noimod=1</code>	Noise model selector.
176	<code>kf=0</code>	Flicker (1/f) noise coefficient.
177	<code>af=1</code>	Flicker (1/f) noise exponent.
178	<code>ef=1</code>	Flicker (1/f) noise frequency exponent.
179	<code>noia=1e20</code>	Oxide trap density coefficient. Default is 9.9e18 for pmos.
180	<code>noib=5e4</code>	Oxide trap density coefficient. Default is 2.4e3 for pmos.
181	<code>noic=-1.4e-12</code>	Oxide trap density coefficient. Default is 1.4e-8 for pmos.
182	<code>em=4.1e7 V/m</code>	Maximum electric field.

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Auto Model Selector parameters

183	<code>wmax=1 m</code>	Maximum channel width for which the model is valid.
184	<code>wmin=0 m</code>	Minimum channel width for which the model is valid.
185	<code>lmax=1 m</code>	Maximum channel length for which the model is valid.
186	<code>lmin=0 m</code>	Minimum channel length for which the model is valid.

Operating region warning control parameters

187	<code>alarm=none</code>	Forbidden operating region. Possible values are <code>none</code> , <code>off</code> , <code>triode</code> , <code>sat</code> , <code>subth</code> , or <code>rev</code> .
188	<code>imax=1 A</code>	Maximum allowable current.
189	<code>bvj=∞ V</code>	Junction reverse breakdown voltage.
190	<code>vbox=1e9 tox V</code>	Oxide breakdown voltage.
191	<code>warn=on</code>	Parameter to turn warnings on and off. Possible values are <code>off</code> or <code>on</code> .

SOI specific parameters

192	<code>vbsa=0 V</code>	Vbs0t offset voltage.
193	<code>delp=0.02</code>	Offset constant for limiting Vbseff to Phis.
194	<code>kb1=1</code>	Scaling factor for backgate charge.
195	<code>kb3=1</code>	Backgate coupling coefficient at subthreshold.
196	<code>dvbd0=0 V</code>	First coefficient of short-channel effect on Vbs0t.
197	<code>dvbd1=0</code>	First coefficient of short-channel effect on Vbs0t.
198	<code>abp=1</code>	Gate bias coefficient for Xcsat calculation.
199	<code>mx_C=-0.9</code>	A smoothing parameter for Xcsat calculation.

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200	agidl=0	GIDL constant.
201	bgidl=0 V/m	GIDL exponential coefficient.
202	ngidl=1.2 V	GIDL Vds enhancement coefficient.
203	ntun=10	Reverse tunneling non-ideality factor.
204	nrecf0=2.0	Recombination non-ideality factor at forward bias.
205	nrecr0=10	Recombination non-ideality factor at reversed bias.
206	vsdfb (F/m)	Source/Drain diffusion flatband voltage.
207	vsdth	Source/Drain diffusion threshold voltage.
208	csdmin (F)	Source/Drain diffusion bottom minimum capacitance.
209	csdesw=0	Source/drain sidewall fringing constant.
210	aii=0	First parameter for critical field.
211	bii=0	Second parameter for critical field.
212	cii=0	Gate dependence of critical field.
213	dii=-1	Body dependence of critical field.
214	ndiode=1	Diode non-ideality factor.
215	asd=0.3	Source/Drain diffusion smoothing parameter.
216	isbjt=1e-6 A	BJT saturation current.
217	isdif=0 A	Diffusion saturation current.
218	isrec=1e-5 A	Recombination saturation current.
219	istun=0 A	Tunneling saturation current.
220	ln=2e-6 m	Electron diffusion length.
221	vrec0=0 V	Voltage dependent parameter for recombination current.

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222	$v_{tun0}=0$ V	Voltage dependent parameter for tunneling current.
223	$n_{bjt}=1$	Power coefficient of channel length dependency for bipolar current.
224	$l_{bjt0}=0.20e-6$ m	Reference channel length for bipolar current.
225	$v_{abjt}=10$ V	Early voltage for bipolar current.
226	$a_{ely}=0$ V	Channel length dependency of early voltage for bipolar current.
227	$a_{hli}=0$	High level injection parameter for bipolar current.
228	$k_{bjt1}=0$ m	Parasitic bipolar base width.

Gate tunneling parameters

229	$w_{th0}=0.0$ μm	Minimum width for thermal resistance calculation..
230	$r_{halo}=1.0e15$ Ω/sqr	Body halo sheet resistance.
231	$n_{tox}=1.0$	Power term of gate current.
232	$t_{oxref}=2.5e-9$ m	Target oxide thickness.
233	$e_{bg}=1.2$ V	Effective bandgap in gate current calculation.
234	$n_{evb}=3.0$	Valence-band electron non-ideality factor.
235	$\alpha_{hagb1}=0.35$	First V_{ox} dependent parameter for gate current in inversion..
236	$\beta_{tagb1}=0.03$	Second V_{ox} dependent parameter for gate current in inversion..
237	$v_{gb1}=300$	Third V_{ox} dependent parameter for gate current in inversion..
238	$\alpha_{hagb2}=0.43$	First V_{ox} dependent parameter for gate current in accumulation..
239	$\beta_{tagb2}=0.05$	Second V_{ox} dependent parameter for gate current in accumulation..
240	$n_{ecb}=1.0$	Conduction-band electron non-ideality factor.

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241	<code>vgb2=17</code>	Third Vox dependent parameter for gate current in accumulation..
242	<code>toxqm=Tox m</code>	Effective oxide thickness considering quantum effects..
243	<code>voxh=5.0 V</code>	Limit of Vox in gate current calculation..
244	<code>deltavox=0.005 V</code>	Smoothing parameter in the Vox smoothing function..
245	<code>igmod=0</code>	Gate current model selector.

Length dependent parameters (Not listed)

Width dependent parameters (Not listed)

Cross-term dependent parameters

246	<code>paramchk=1</code>	Model parameter checking selector.
247	<code>noif=1</code>	Floating body excess noise ideality factor.
248	<code>w0flk=0 m</code>	Width constant for IBM flicker noise equation.
249	<code>frbody=1</code>	Layout dependent body-resistance coefficient.
250	<code>vevb=0.075</code>	Vaux parameter for valence-band electron tunneling.
251	<code>vecb=0.026</code>	Vaux parameters for conduction-band electron tunneling.
252	<code>dtoxcv=0.0 m</code>	Delta oxide thickness in Capmod3.
253	<code>llc=0 m</code>	Length dependence of delta LC.
254	<code>lwc=0 m</code>	Width dependence of delta LC.
255	<code>lwlc=0 m²</code>	Area dependence of delta LC.
256	<code>wlc=0 m</code>	Length dependence of delta WC.
257	<code>wwc=0 m</code>	Width dependence of delta WC.

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258 $w \cdot l \cdot c = 0 \text{ m}^2$

Area dependence of delta WC.

The `jmelt` parameter is used to aid convergence and prevent numerical overflow. The junction characteristics of the FET are accurately modeled for current (density) up to `jmelt`. For current density above `jmelt`, the junction is modeled as a linear resistor and a warning is printed.

Auto Model Selection

Many models need to be characterized for different geometries in order to obtain accurate results for model development. The model selector program automatically searches for a model with the length and width range specified in the instance statement and uses this model in the simulations.

For the auto model selector program to find a specific model, the models to be searched should be grouped together within braces. Such a group is called a model group. An opening brace is required at the end of the line defining each model group. Every model in the group is given a name followed by a colon and the list of parameters. Also, the four geometric parameters `lmax`, `lmin`, `wmax`, and `wmin` should be given. The selection criteria to choose a model is as follows:

$$l_{\min} \leq \text{inst_length} < l_{\max} \text{ and } w_{\min} \leq \text{inst_width} < w_{\max}$$

Example:

```
model ModelName ModelType {  
  1: <model parameters> lmin=2 lmax=4 wmin=1 wmax=2  
  2: <model parameters> lmin=1 lmax=2 wmin=2 wmax=4  
  3: <model parameters> lmin=2 lmax=4 wmin=4 wmax=6  
}
```

Then for a given instance

```
M1 1 2 3 4 ModelName w=3 l=1.5
```

the program would search all the models in the model group with the name `ModelName` and then pick the first model whose geometric range satisfies the selection criteria. In the preceding example, the auto model selector program would choose `ModelName.2`.

You must specify both length (`l`) and width (`w`) on the device instance line to enable automatic model selection.

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Output Parameters

1	<code>w_{eff}</code> (m)	Effective channel width.
2	<code>l_{eff}</code> (m)	Effective channel length.
3	<code>r_{th_{eff}}</code> (Ω)	Effective thermal resistance.
4	<code>C_{th_{eff}}</code> (F)	Effective thermal capacitance.
5	<code>r_{se_{eff}}</code> (Ω)	Effective source resistance.
6	<code>r_{d_{eff}}</code> (Ω)	Effective drain resistance.

Operating-Point Parameters

1	<code>type=n</code>	Transistor type. Possible values are <code>n</code> or <code>p</code> .
2	<code>region=triode</code>	Estimated operating region. Spectre outputs number (0-3) in a rawfile. Possible values are <code>off</code> , <code>triode</code> , <code>sat</code> , or <code>subth</code> .
3	<code>reversed</code>	Reverse mode indicator. Possible values are <code>no</code> or <code>yes</code> .
4	<code>v_{gs}</code> (V)	Gate-source voltage.
5	<code>v_{ds}</code> (V)	Drain-source voltage.
6	<code>v_{bs}</code> (V)	Bulk-source voltage.
7	<code>v_{bgs}</code> (V)	Back-Gate-source voltage.
8	<code>i_{ds}</code> (A)	Resistive drain-to-source current.
9	<code>i_c</code> (A)	BJT collector current.
10	<code>i_{sgidl}</code> (A)	Source GIDL current.
11	<code>i_{dgidl}</code> (A)	Drain GIDL current.
12	<code>i_{ii}</code> (A)	Impact ionization current.

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13	i_{bd} (A)	Resistive bulk-to-drain junction current.
14	i_{gbt} (A)	Gate-to-body tunneling current.
15	i_{bs} (A)	Resistive bulk-to-source junction current.
16	v_{th} (V)	Threshold voltage.
17	v_{dsat} (V)	Drain-source saturation voltage.
18	g_m (S)	Common-source transconductance.
19	g_{ds} (S)	Common-source output conductance.
20	g_{mb} (S)	Body-transconductance.
21	g_{mbg} (S)	Back-gate-transconductance.
22	μ_{eff} ($cm^2/V\ s$)	Effective mobility.
23	β_{eff} (A/V^2)	Effective β .
24	q_g (Coul)	Gate charge.
25	q_d (Coul)	Drain charge.
26	q_s (Coul)	Source charge.
27	q_b (Coul)	Body charge.
28	q_{bg} (Coul)	Back-Gate charge.
29	c_{gg} (F)	dQ_g/dV_g .
30	c_{gd} (F)	dQ_g/dV_d .
31	c_{gs} (F)	dQ_g/dV_s .
32	c_{gb} (F)	dQ_g/dV_{bk} .
33	c_{dg} (F)	dQ_d/dV_g .
34	c_{dd} (F)	dQ_d/dV_d .

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35	<code>cds</code>	(F)	<code>dQd_dVs.</code>
36	<code>cdb</code>	(F)	<code>dQd_dVb.</code>
37	<code>csg</code>	(F)	<code>dQs_dVg.</code>
38	<code>csd</code>	(F)	<code>dQs_dVd.</code>
39	<code>css</code>	(F)	<code>dQs_dVs.</code>
40	<code>csb</code>	(F)	<code>dQs_dVb.</code>
41	<code>cbg</code>	(F)	<code>dQb_dVg.</code>
42	<code>cbd</code>	(F)	<code>dQb_dVd.</code>
43	<code>cbs</code>	(F)	<code>dQb_dVs.</code>
44	<code>cbb</code>	(F)	<code>dQb_dVb.</code>
45	<code>id</code>	(A)	Total resistive drain current.
46	<code>is</code>	(A)	Total resistive source current.
47	<code>ib</code>	(A)	Total resistive bulk current.
48	<code>pwr</code>	(W)	Power at op point.
49	<code>gmoverid</code>	(1/V)	Gm/Ids.
50	<code>tdev</code>	(C)	Temperature rise from ambient.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

`Ctheff` O-4 `dt1` M-165 `lwl` M-40 `siid` M-94

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a0	M-22	dt2	M-166	lwlc	M-255	tbox	M-50
a1	M-25	dtoxcv	M-252	lwn	M-39	tcjswg	M-156
a2	M-26	dvbd0	M-196	m	I-10	tdev	OP-50
abp	M-198	dvbd1	M-197	meto	M-113	tii	M-88
ad	I-4	dvt0	M-16	minr	M-106	tlev	M-147
ad	M-138	dvt0w	M-19	mjswg	M-118	tlevc	M-148
aebcp	I-20	dvt1	M-17	mobmod	M-60	tmax	M-145
aely	M-226	dvt1w	M-20	mxo	M-199	tnodeout	I-22
af	M-177	dvt2	M-18	nbc	I-15	tnom	M-144
agbcp	I-19	dvt2w	M-21	nbjt	M-223	tox	M-49
agidl	M-200	dwb	M-47	nch	M-31	toxqm	M-242
ags	M-27	dwbc	M-48	ndif	M-121	toxref	M-232
ahli	M-227	dwc	M-124	ndiode	M-214	tpbswg	M-157
aia	M-210	dwg	M-46	necb	M-240	trd	M-163
alarm	M-187	ebg	M-233	nevb	M-234	trs	M-162
alpha0	M-82	ef	M-178	nfactor	M-76	tsi	M-51
alphagb1	M-235	eg	M-149	ngate	M-32	tt	M-120
alphagb2	M-238	em	M-182	ngidl	M-202	type	M-1
as	M-137	esatii	M-90	nlx	M-11	type	OP-1
as	I-3	eta0	M-80	noia	M-179	u0	M-61
asd	M-215	etab	M-81	noib	M-180	ua	M-63
at	M-155	fbjtii	M-84	noic	M-181	ua1	M-158
b0	M-23	fbody	M-126	noif	M-247	ub	M-64
b1	M-24	frbody	M-249	noimod	M-175	ub1	M-159
beta0	M-83	gamma1	M-12	nrb	I-9	uc	M-65
beta1	M-85	gamma2	M-13	nrb	M-143	uc1	M-160
beta2	M-86	gap1	M-150	nrd	I-7	ueff	OP-22
betaeff	OP-23	gap2	M-151	nrd	M-141	ute	M-164
betagb1	M-236	gds	OP-19	nrecf0	M-204	vabjt	M-225
betagb2	M-239	gm	OP-18	nrecr0	M-205	vbgs	OP-7
bgidl	M-201	gmb	OP-20	nrs	I-8	vbm	M-15
bii	M-211	gmbg	OP-21	nrs	M-142	vbox	M-190

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bjtoff I-14	hdif M-104	nsub M-30	vbsa M-192
bvj M-189	ib OP-47	ntox M-231	vbsusr I-21
capmod M-123	ibd OP-13	ntrecf M-169	vbxb M-14
cbb OP-44	ibs OP-15	ntrechr M-170	vds OP-5
cbd OP-42	ic OP-9	ntun M-203	vdsat OP-17
cbg OP-41	id OP-45	paramchk M-246	vdsatii0 M-87
cbs OP-43	idgidl OP-11	pbswg M-119	vecb M-251
cdb OP-36	ids OP-8	pclm M-67	version M-2
cdd OP-34	igbt OP-14	pd I-6	vevb M-250
cdg OP-33	igmod M-245	pd M-140	vfbcv M-133
cds OP-35	iii OP-12	pdbcp I-17	vgl M-237
cdsc M-73	imax M-188	pdiblc1 M-68	vgl2 M-241
cdscb M-74	imelt M-108	pdiblc2 M-69	vgs OP-4
cdscd M-75	is OP-46	pdiblc3 M-70	voff M-78
cf M-132	isbjt M-216	prt M-161	voxh M-243
cgb OP-32	isdif M-217	prwb M-54	vrec0 M-221
cgbo M-112	isgidl OP-10	prwg M-55	vsat M-62
cgd OP-30	isrec M-218	ps I-5	vsdfb M-206
cgdl M-115	istun M-219	ps M-139	vsdth M-207
cgdo M-110	k1 M-4	psbcp I-18	vth OP-16
cgeo M-111	k1w1 M-5	pvag M-71	vtho M-3
cgg OP-29	k1w2 M-6	pwr OP-48	vtun0 M-222
cgs OP-31	k2 M-7	qb OP-27	w M-135
cgs1 M-114	k3 M-8	qbg OP-28	w I-1
cgso M-109	k3b M-9	qd OP-25	w0 M-10
cii M-212	kbl M-194	qg OP-24	w0flk M-248
cit M-77	kb3 M-195	qs OP-26	warn M-191
cjswg M-117	kbjt1 M-228	rbody M-99	weff O-1
ckappa M-116	keta M-28	rbsh M-95	wint M-35
clc M-130	ketas M-29	rd M-98	wl M-41
cle M-131	kf M-176	rdc M-101	wlc M-256

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csb	OP-40	kt1	M-152	rdd	M-103	wln	M-42
csd	OP-38	kt11	M-153	rdeff	O-6	wmax	M-183
csdesw	M-209	kt2	M-154	rdsw	M-53	wmin	M-184
csdmin	M-208	l	I-2	region	OP-2	wr	M-56
csg	OP-37	l	M-136	region	I-11	wth0	M-229
css	OP-39	lbjt0	M-224	reversed	OP-3	ww	M-43
cth0	I-13	ldif	M-105	rhalo	M-230	wwc	M-257
cth0	M-167	ldif0	M-122	rs	M-97	wwl	M-45
delp	M-193	leff	O-2	rsc	M-100	wwlc	M-258
delta	M-72	lii	M-89	rseff	O-5	wwn	M-44
deltavox	M-244	lint	M-34	rsh	M-96	xbjt	M-171
delvt	M-125	ll	M-36	rss	M-102	xdif	M-172
dii	M-213	llc	M-253	rth0	I-12	xj	M-33
dlbg	M-129	lln	M-37	rth0	M-168	xl	M-57
dlc	M-127	lmax	M-185	rtheff	O-3	xpart	M-134
dlcb	M-128	lmin	M-186	shmod	M-146	xrec	M-173
drout	M-66	ln	M-220	sii0	M-91	xt	M-52
dskip	M-107	lw	M-38	sii1	M-92	xtun	M-174
dsub	M-79	lwc	M-254	sii2	M-93	xw	M-58

Bipolar Junction Transistor (bjt)

Description

The bipolar transistor model is adapted from the integral charge model of Gummel and Poon, and it includes several high bias-level effects. This model defaults to the simpler Ebers-Moll model if certain parameters are left unspecified. This model also includes a substrate junction that connects either to the collector or to the base to model vertical and lateral structures.

This model has the following enhancements over SPICE2G.6:

1. Two base resistance models are provided.
2. Nonlinear collector resistance is implemented.

Spectre Circuit Simulator Reference

Component Statements Part I

3. The integral form of the Early voltage effect is available.
4. The substrate junction includes both the diode and the capacitor.

This device is supported within altergroups.

Sample Instance Statement

```
q1 (vcc net3 minus) npn_mod region=fwd area=1 m=1
```

Sample Model Statement

```
model npn_mod bjt type=npn is=10e-13 bf=200 va=58.8 ikf=5.63e-3 rb=700 rbm=86  
re=3.2 cje=0.352e-12 pe=0.76 me=0.34 tf=249e-12 cjc=0.34e-12 pc=0.55
```

Instance Definition

```
Name c b e [s] ModelName parameter=value ...
```

You do not have to specify the substrate terminal. If you do not specify it, the substrate is connected to ground.

Instance Parameters

- | | | |
|---|------------|---|
| 1 | area=1 | Transistor area factor. |
| 2 | areab=1 | Transistor areab factor. |
| 3 | areac=1 | Transistor areac factor. |
| 4 | m=1 | Multiplicity factor. |
| 5 | trise | Temperature rise from ambient. |
| 6 | region=fwd | Estimated operating region. Spectre outputs number (0-4) in a rawfile.
Possible values are off, fwd, rev, sat, or breakdown. |

Model Definition

```
model modelName bjt parameter=value ...
```

Spectre Circuit Simulator Reference

Component Statements Part I

Model Parameters

Structural parameters

- 1 `type=npn` Transistor type.
Possible values are `npn` or `pnp`.
- 2 `struct=vertical` Transistor structure. For `pnp` default=`lateral`.
Possible values are `vertical` or `lateral`.

Saturation current parameters

- 3 `is=1e-16 A` Saturation current (*area).
- 4 `ise=0 A` B-E leakage saturation current. Set to `c2*is` if not given.
(*area).
- 5 `isc=0 A` B-C leakage saturation current. Set to `c4*is` if not given.
(*area).
- 6 `iss=0 A` Substrate leakage saturation current (*area).
- 7 `c2=0` Forward leakage saturation current coefficient.
- 8 `c4=0` Reverse leakage saturation current coefficient.

B-C leakage model parameters

- 9 `cbo=0 A` Extrapolated 0-volt B-C leakage current (*area).
- 10 `gbo=0 S` Slope of I_{cbo} vs. V_{bc} above V_{bo} (*area).
- 11 `vbo=0 V` Slope of I_{cbo} vs. V_{bc} at $V_{bc}=0$.
- 12 `tcbo=0 1/C` Temperature coefficient for `cbo`.
- 13 `tgbo=0 1/C` Temperature coefficient for `gbo`.

Spectre Circuit Simulator Reference

Component Statements Part I

Emission coefficient parameters

14	$n_f=1$	Forward emission coefficient.
15	$n_r=1$	Reverse emission coefficient.
16	$n_e=1.5$	B-E leakage emission coefficient.
17	$n_c=2$	B-C leakage emission coefficient.
18	$n_s=1$	Substrate junction emission coefficient.

Current gain parameters

19	$b_f=100$ A/A	Forward current gain (beta).
20	$b_r=1$ A/A	Reverse current gain (beta).
21	$i_{kf}=\infty$ A	High current corner for forward beta (*area).
22	$i_{kr}=\infty$ A	High current corner for reverse beta (*area).

Early voltage parameters

23	$v_{af}=\infty$ V	Forward Early voltage.
24	$v_{ar}=\infty$ V	Reverse Early voltage.
25	$k_e=0$ 1/V	B-E space-charge integral multiplier.
26	$k_c=0$ 1/V	B-C space-charge integral multiplier.

Parasitic resistance parameters

27	$r_b=0$ Ω	Zero-bias base resistance (/area).
28	$r_{bm}=r_b$ Ω	Minimum base resistance for high currents (/area).
29	$i_{rb}=\infty$ A	Current at base resistance midpoint (*area).

Spectre Circuit Simulator Reference

Component Statements Part I

30	<code>rbmod=spice</code>	Nonlinear Rb model. Possible values are <code>spectre</code> or <code>spice</code> .
31	<code>rc=0 Ω</code>	Collector resistance (/area).
32	<code>rcv=0 Ω</code>	Variable collector resistance (/area).
33	<code>rcm=0 Ω</code>	Minimum collector resistance (/area).
34	<code>dope=1e15 cm⁻³</code>	Collector background doping concentration.
35	<code>cex=1</code>	Current crowding exponent.
36	<code>cco=1 A</code>	Current crowding normalization constant (*area).
37	<code>re=0 Ω</code>	Emitter resistance (/area).
38	<code>minr=0.1 Ω</code>	Minimum parasitic resistance.

Junction capacitance parameters

39	<code>cje=0 F</code>	B-E zero-bias junction capacitance (*area).
40	<code>vje=0.75 V</code>	B-E built-in junction potential.
41	<code>mje=1/3</code>	B-E junction exponent.
42	<code>cjc=0 F</code>	B-C zero-bias junction capacitance (*area).
43	<code>vjc=0.75 V</code>	B-C built-in junction potential.
44	<code>mjc=1/3</code>	B-C junction exponent.
45	<code>xcjc=1</code>	Fraction of B-C capacitance tied to internal base node.
46	<code>xcjc2=1</code>	Fraction of B-C capacitance tied to collector and fraction of B-C tied to internal node.
47	<code>cjs=0 F</code>	B-S zero-bias junction capacitance (*area).
48	<code>vjs=0.75 V</code>	B-S built-in junction potential.

Spectre Circuit Simulator Reference

Component Statements Part I

49	$mjs=0$	B-S junction exponent.
50	$fc=0.5$	Junction capacitor forward-bias threshold.
51	$cbc_p=0$ F	B-C parasitic capacitance.
52	$cbe_p=0$ F	B-E parasitic capacitance.
53	$ccs_p=0$ F	C-S parasitic capacitance.

Transit time and excess phase parameters

54	$t_f=0$ s	Ideal forward transit time.
55	$t_d=0$ s	Intrinsic base delay time.
56	$x_{t_f}=0$	Coefficient for bias dependence of t_f .
57	$v_{t_f}=\infty$ V	Voltage describing V_{bc} dependence of t_f .
58	$i_{t_f}=0$ A	High current parameter for effect on t_f (*area).
59	$t_r=0$ s	Ideal reverse transit time.
60	$p_{t_f}=0$ °	Excess phase at $\text{freq} = 1.0/(t_f * 2 \pi)$ Hz.

Temperature effects parameters

61	t_{nom} (C)	Parameters measurement temperature. Default set by <code>options</code> .
62	$t_{rise}=0$ C	Temperature rise from ambient.
63	$e_g=1.11$ V	Band-gap.
64	$x_{t_b}=0$	Beta temperature exponent.
65	$x_{t_i}=3$	Temperature exponent for effect on i_s .
66	$t_{rb1}=0$ 1/C	Linear temperature coefficient for the base resistor.
67	$t_{rb2}=0$ C ⁻²	Quadratic temperature coefficient for the base resistor.

Spectre Circuit Simulator Reference

Component Statements Part I

68	<code>trm1=0</code>	1/C	Linear temperature coefficient for the minimum base resistor.
69	<code>trm2=0</code>	C ⁻²	Quadratic temperature coefficient for the minimum base resistor.
70	<code>trc1=0</code>	1/C	Linear temperature coefficient for the collector resistor.
71	<code>trc2=0</code>	C ⁻²	Quadratic temperature coefficient for the collector resistor.
72	<code>tre1=0</code>	1/C	Linear temperature coefficient for the emitter resistor.
73	<code>tre2=0</code>	C ⁻²	Quadratic temperature coefficient for the emitter resistor.
74	<code>tlev=0</code>		DC temperature selector.
75	<code>tlevc=0</code>		AC temperature selector.
76	<code>gap1=7.02e-4</code>	V/C	Band-gap temperature coefficient.
77	<code>gap2=1108</code>	C	Band-gap temperature offset.
78	<code>tikf1=0</code>	1/C	Linear temperature coefficient for <code>ikf</code> .
79	<code>tikf2=0</code>	C ⁻²	Quadratic temperature coefficient for <code>ikf</code> .
80	<code>tikr1=0</code>	1/C	Linear temperature coefficient for <code>ikr</code> .
81	<code>tikr2=0</code>	C ⁻²	Quadratic temperature coefficient for <code>ikr</code> .
82	<code>tirb1=0</code>	1/C	Linear temperature coefficient for <code>irb</code> .
83	<code>tirb2=0</code>	C ⁻²	Quadratic temperature coefficient for <code>irb</code> .
84	<code>tis1=0</code>	1/C	Linear temperature coefficient for <code>is</code> .
85	<code>tis2=0</code>	C ⁻²	Quadratic temperature coefficient for <code>is</code> .
86	<code>tise1=0</code>	1/C	Linear temperature coefficient for <code>ise</code> .
87	<code>tise2=0</code>	C ⁻²	Quadratic temperature coefficient for <code>ise</code> .
88	<code>tisc1=0</code>	1/C	Linear temperature coefficient for <code>isc</code> .
89	<code>tisc2=0</code>	C ⁻²	Quadratic temperature coefficient for <code>isc</code> .

Spectre Circuit Simulator Reference

Component Statements Part I

90	$t_{iss1}=0$	1/C	Linear temperature coefficient for <i>iss</i> .
91	$t_{iss2}=0$	C^{-2}	Quadratic temperature coefficient for <i>iss</i> .
92	$t_{bf1}=0$	1/C	Linear temperature coefficient for <i>bf</i> .
93	$t_{bf2}=0$	C^{-2}	Quadratic temperature coefficient for <i>bf</i> .
94	$t_{br1}=0$	1/C	Linear temperature coefficient for <i>br</i> .
95	$t_{br2}=0$	C^{-2}	Quadratic temperature coefficient for <i>br</i> .
96	$t_{vaf1}=0$	1/C	Linear temperature coefficient for <i>vaf</i> .
97	$t_{vaf2}=0$	C^{-2}	Quadratic temperature coefficient for <i>vaf</i> .
98	$t_{var1}=0$	1/C	Linear temperature coefficient for <i>var</i> .
99	$t_{var2}=0$	C^{-2}	Quadratic temperature coefficient for <i>var</i> .
100	$t_{itf1}=0$	1/C	Linear temperature coefficient for <i>itf</i> .
101	$t_{itf2}=0$	C^{-2}	Quadratic temperature coefficient for <i>itf</i> .
102	$t_{tf1}=0$	1/C	Linear temperature coefficient for <i>tf</i> .
103	$t_{tf2}=0$	C^{-2}	Quadratic temperature coefficient for <i>tf</i> .
104	$t_{tr1}=0$	1/C	Linear temperature coefficient for <i>tr</i> .
105	$t_{tr2}=0$	C^{-2}	Quadratic temperature coefficient for <i>tr</i> .
106	$t_{nf1}=0$	1/C	Linear temperature coefficient for <i>nf</i> .
107	$t_{nf2}=0$	C^{-2}	Quadratic temperature coefficient for <i>nf</i> .
108	$t_{nr1}=0$	1/C	Linear temperature coefficient for <i>nr</i> .
109	$t_{nr2}=0$	C^{-2}	Quadratic temperature coefficient for <i>nr</i> .
110	$t_{ne1}=0$	1/C	Linear temperature coefficient for <i>ne</i> .
111	$t_{ne2}=0$	C^{-2}	Quadratic temperature coefficient for <i>ne</i> .

Spectre Circuit Simulator Reference

Component Statements Part I

112	$t_{nc1}=0$	1/C	Linear temperature coefficient for n_c .
113	$t_{nc2}=0$	C^{-2}	Quadratic temperature coefficient for n_c .
114	$t_{ns1}=0$	1/C	Linear temperature coefficient for n_s .
115	$t_{ns2}=0$	C^{-2}	Quadratic temperature coefficient for n_s .
116	$t_{mje1}=0$	1/C	Linear temperature coefficient for m_{je} .
117	$t_{mje2}=0$	C^{-2}	Quadratic temperature coefficient for m_{je} .
118	$t_{mjc1}=0$	1/C	Linear temperature coefficient for m_{jc} .
119	$t_{mjc2}=0$	C^{-2}	Quadratic temperature coefficient for m_{jc} .
120	$t_{mjs1}=0$	1/C	Linear temperature coefficient for m_{js} .
121	$t_{mjs2}=0$	C^{-2}	Quadratic temperature coefficient for m_{js} .
122	$ct_e=0$	1/C	Temperature coefficient for c_{je} .
123	$ct_c=0$	1/C	Temperature coefficient for c_{jc} .
124	$ct_s=0$	1/C	Temperature coefficient for c_{js} .
125	$tv_{je}=0$	V/C	Temperature coefficient for v_{je} .
126	$tv_{jc}=0$	V/C	Temperature coefficient for v_{jc} .
127	$tv_{js}=0$	V/C	Temperature coefficient for v_{js} .
128	$t_{vtf1}=0$	1/C	Linear temperature coefficient for v_{tf} .
129	$t_{vtf2}=0$	C^{-2}	Quadratic temperature coefficient for v_{tf} .
130	$t_{xtf1}=0$	1/C	Linear temperature coefficient for x_{tf} .
131	$t_{xtf2}=0$	C^{-2}	Quadratic temperature coefficient for x_{tf} .

Spectre Circuit Simulator Reference

Component Statements Part I

Junction diode model control parameters

- 132 `dskip=yes` Skip junction calculations if they are reverse-saturated.
Possible values are `no` or `yes`.
- 133 `imelt=imax A` Junction explosion current (*area).

Operating region warning control parameters

- 134 `bvbe= ∞ V` B-E breakdown voltage.
- 135 `bvbc= ∞ V` B-C breakdown voltage.
- 136 `bvce= ∞ V` C-E breakdown voltage.
- 137 `bvsub= ∞ V` Substrate junction breakdown voltage.
- 138 `vbefwd=0.2 V` B-E forward voltage.
- 139 `vbcfwd=0.2 V` B-C forward voltage.
- 140 `vsubfwd=0.2 V` Substrate junction forward voltage.
- 141 `imax=1e3 A` Maximum allowable base current (*area).
- 142 `imax1=imax A` Maximum allowable collector current (*area).
- 143 `alarm=none` Forbidden operating region.
Possible values are `none`, `off`, `fwd`, `rev`, or `sat`.

Noise model parameters

- 144 `kf=0` Flicker (1/f) noise coefficient.
- 145 `af=1` Flicker (1/f) noise exponent.
- 146 `kb=0` Burst noise coefficient.
- 147 `bnoisefc=1` Burst noise cutoff frequency.
- 148 `rbnoi= $r_b \Omega$` Effective base noise resistance.

Spectre Circuit Simulator Reference

Component Statements Part I

Imax and Imelt

The `imax` parameter aids convergence and prevents numerical overflow. The junction characteristics of the device are accurately modeled for current up to `imax`. If `imax` is exceeded during iterations, the linear model is substituted until the current drops below `imax` or until convergence is achieved. If convergence is achieved with the current exceeding `imax`, the results are inaccurate, and Spectre prints a warning.

A separate model parameter, `imelt`, is used as a limit warning for the junction current. This parameter can be set to the maximum current rating of the device. When any component of the junction current exceeds `imelt`, note that base and collector currents are composed of many exponential terms, Spectre issues a warning and the results become inaccurate. The junction current is linearized above the value of `imelt` to prevent arithmetic exception, with the exponential term replaced by a linear equation at `imelt`.

Operating-Point Parameters

1	<code>type=npn</code>	Transistor type. Possible values are <code>npn</code> or <code>pnP</code> .
2	<code>struct=vertical</code>	Transistor structure. For <code>pnP</code> default= <code>lateral</code> . Possible values are <code>vertical</code> or <code>lateral</code> .
3	<code>region=fwd</code>	Estimated operating region. Spectre outputs number (0-4) in a rawfile. Possible values are <code>off</code> , <code>fwd</code> , <code>rev</code> , <code>sat</code> , or <code>breakdown</code> .
4	<code>vbe (V)</code>	Base-emitter voltage.
5	<code>vbc (V)</code>	Base-collector voltage.
6	<code>vce (V)</code>	Collector-emitter voltage.
7	<code>vsub (V)</code>	Substrate junction voltage.
8	<code>ic (A)</code>	Resistive collector current.
9	<code>ib (A)</code>	Resistive base current.
10	<code>isub (A)</code>	Resistive substrate current.
11	<code>pwr (W)</code>	Power dissipation.

Spectre Circuit Simulator Reference

Component Statements Part I

12	betadc (A/A)	Ratio of resistive collector current to resistive base current.
13	betaac (A/A)	Small-signal common-emitter current gain.
14	gm (S)	Common-emitter transconductance.
15	rpi (Ω)	Common-emitter input resistance.
16	ro (Ω)	Common-emitter output resistance.
17	rb (Ω)	Parasitic base resistance.
18	rc (Ω)	Parasitic collector resistance.
19	cpi (F)	Common-emitter input capacitance.
20	cmu (F)	Common-base output capacitance.
21	cmux (F)	External common-base output capacitance.
22	csub (F)	Substrate capacitance.
23	ft (Hz)	Unity small-signal current-gain frequency.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

af	M-145	imax	M-141	tbr2	M-95	tre1	M-72
alarm	M-143	imax1	M-142	tcbo	M-12	tre2	M-73
area	I-1	imelt	M-133	t d	M-55	trise	M-62
areab	I-2	irb	M-29	tf	M-54	trise	I-5
areac	I-3	is	M-3	tgbo	M-13	trm1	M-68
betaac	OP-13	isc	M-5	tikf1	M-78	trm2	M-69
betadc	OP-12	ise	M-4	tikf2	M-79	ttf1	M-102

Spectre Circuit Simulator Reference Component Statements Part I

bf	M-19	iss	M-6	tikr1	M-80	ttf2	M-103
bnoisefc	M-147	isub	OP-10	tikr2	M-81	ttr1	M-104
br	M-20	itf	M-58	tirb1	M-82	ttr2	M-105
bvbc	M-135	kb	M-146	tirb2	M-83	tvaf1	M-96
bvbe	M-134	kc	M-26	tis1	M-84	tvaf2	M-97
bvce	M-136	ke	M-25	tis2	M-85	tvar1	M-98
bvsub	M-137	kf	M-144	tisc1	M-88	tvar2	M-99
c2	M-7	m	I-4	tisc2	M-89	tvjc	M-126
c4	M-8	minr	M-38	tisel	M-86	tvje	M-125
cbcp	M-51	mjc	M-44	tise2	M-87	tvjs	M-127
cbep	M-52	mje	M-41	tiss1	M-90	tvtf1	M-128
cbo	M-9	mjs	M-49	tiss2	M-91	tvtf2	M-129
cco	M-36	nc	M-17	titf1	M-100	txtf1	M-130
ccsp	M-53	ne	M-16	titf2	M-101	txtf2	M-131
cex	M-35	nf	M-14	tlev	M-74	type	M-1
cjc	M-42	nr	M-15	tlevc	M-75	type	OP-1
cje	M-39	ns	M-18	tmjc1	M-118	vaf	M-23
cjs	M-47	ptf	M-60	tmjc2	M-119	var	M-24
cmu	OP-20	pwr	OP-11	tmje1	M-116	vbc	OP-5
cmux	OP-21	rb	M-27	tmje2	M-117	vbcfwd	M-139
cpu	OP-19	rb	OP-17	tmjs1	M-120	vbe	OP-4
csub	OP-22	rbm	M-28	tmjs2	M-121	vbefwd	M-138
ctc	M-123	rbmod	M-30	tnc1	M-112	vbo	M-11
cte	M-122	rbnoi	M-148	tnc2	M-113	vce	OP-6
cts	M-124	rc	M-31	tne1	M-110	vjc	M-43
dope	M-34	rc	OP-18	tne2	M-111	vje	M-40
dskip	M-132	rcm	M-33	tnf1	M-106	vjs	M-48
eg	M-63	rcv	M-32	tnf2	M-107	vsub	OP-7
fc	M-50	re	M-37	tnom	M-61	vsubfwd	M-140
ft	OP-23	region	OP-3	tnr1	M-108	vtf	M-57
gap1	M-76	region	I-6	tnr2	M-109	xcjc	M-45
gap2	M-77	ro	OP-16	tns1	M-114	xcjc2	M-46

Spectre Circuit Simulator Reference Component Statements Part I

gbo	M-10	rpi	OP-15	tns2	M-115	xtb	M-64
gm	OP-14	struct	OP-2	tr	M-59	xtf	M-56
ib	OP-9	struct	M-2	trb1	M-66	xti	M-65
ic	OP-8	tbf1	M-92	trb2	M-67		
ikf	M-21	tbf2	M-93	trc1	M-70		
ikr	M-22	tbr1	M-94	trc2	M-71		

Lateral PNP Transistor (bjt301)

Description

The bjt301 model provides an extensive description of a lateral integrated circuit junction-isolated PNP transistor. It is described in the Philips Bipolar Modelbook (Dec.93) as TPL level 301.

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In extension to the modelbook description a minimum conductance `gmin` is inserted between the internal base and internal collector node, between the internal base and the internal emitter node, and between the external base and the substrate node to aid convergence. The value of `gmin` is set by an options statement, default = 1e-12 S.

The `imax` parameter is used to aid convergence and to prevent numerical overflow. The junction characteristics of the transistor are accurately modeled for currents up to `imax`. For currents above `imax`, the junction is modeled as a linear resistor and a warning is printed.

This device is supported within altergroups.

This device is dynamically loaded from the shared object `/vobs/spectre_dev/tools.sun4v/spectre/lib/cmi/3.0.doc/libphilips_sh.so`

Sample Instance Statement

```
q2 (minus net3 vcc) pnp_mod region=fwd area=1 m=1
```

Sample Model Statement

```
model pnp_mod bjt301 type=pnp struct=lateral is=1e-14 bf=85 ilf=11e-9 ikf=95e-6  
re=3.2 cje=0.352e-12
```


Spectre Circuit Simulator Reference

Component Statements Part I

Instance Definition

Name c b e [s] ModelName parameter=value ...

Instance Parameters

- | | | |
|---|------------|---|
| 1 | area=1 | Area factor. |
| 2 | mult=1 | Alias of area factor. |
| 3 | m=1 | Multiplicity factor. |
| 4 | region=fwd | Estimated DC operating region, used as a convergence aid.
Possible values are <code>off</code> , <code>fwd</code> , <code>rev</code> , or <code>sat</code> . |

Model Definition

model modelName bjt301 parameter=value ...

Model Parameters

Structural parameters

- | | | |
|---|----------------|---|
| 1 | type=pnp | Transistor type.
Possible values are <code>pnp</code> or <code>pnpl</code> . |
| 2 | struct=lateral | Transistor structure.
Possible values are <code>lateral</code> . |

Current parameters

- | | | |
|---|---------------|---|
| 3 | is=1.0e-15 A | Saturation current. |
| 4 | imax=1.0 A | Explosion current. |
| 5 | bf=100.0 A/A | Ideal forward common-emitter current gain (beta). |
| 6 | ilf=10.0e-9 A | Low-level knee-current of forward beta. |
| 7 | nlf=2.0 | Emission coefficient of non-ideal forward base current. |

Spectre Circuit Simulator Reference

Component Statements Part I

8	<code>ikf=100.0e-6 A</code>	High-injection knee-current of forward beta.
9	<code>nhf=1.0</code>	Basewidening exponent.
10	<code>veaf=50.0 V</code>	Early voltage related to collector junction.
11	<code>br=10.0 A/A</code>	Ideal reverse common-collector current gain (beta).
12	<code>ilr=10.0e-9 A</code>	Low-level knee-current of reverse beta.
13	<code>nlr=2.0</code>	Emission coefficient of non-ideal reverse base current.
14	<code>ikr=100.0e-6 A</code>	High-injection knee-current of reverse beta.
15	<code>iks=100.0e-6 A</code>	High-injection current of substrate effect.
16	<code>xcs=1.0</code>	Current fraction of c-b-s transistor.
17	<code>xes=0.01</code>	Current fraction of e-b-s transistor.

Parasitic resistance parameters

18	<code>rc=1.0 Ω</code>	Collector resistance.
19	<code>rbc=10.0 Ω</code>	Constant part of base resistance.
20	<code>rbv=10.0 Ω</code>	Variable part of base resistance.
21	<code>re=1.0 Ω</code>	Emitter series resistance.

Junction capacitance parameters

22	<code>taub=25.0e-9 s</code>	Forward transit time related to neutral base.
23	<code>taune=1.0e-9 s</code>	Forward transit time related to neutral emitter in neutral e-b region.
24	<code>mtau=1.0</code>	Coefficient of current dependence of taune.
25	<code>cje=100.0e-15 F</code>	Zero bias emitter-base depletion capacitance.

Spectre Circuit Simulator Reference

Component Statements Part I

26	<code>vde=0.55 V</code>	Emitter-base diffusion voltage.
27	<code>pe=0.333</code>	Emitter-base grading coefficient.
28	<code>taur=100.0e-9 s</code>	Ideal reverse transit time.
29	<code>cjc=200.0e-15 F</code>	Zero bias collector-base depletion capacitance.
30	<code>vdc=0.55 V</code>	Collector-base diffusion voltage.
31	<code>pc=0.333</code>	Collector-base grading coefficient.
32	<code>cjs=1.0e-12 F</code>	Zero bias substrate junction depletion capacitance.
33	<code>vds=0.55 V</code>	Substrate junction diffusion voltage.
34	<code>ps=0.333</code>	Substrate junction grading coefficient.
35	<code>exphi=0.3</code>	Excess phase shift.
36	<code>fc=0.95</code>	Coefficient for forward bias capacitance.

Temperature effects parameters

37	<code>tref (C)</code>	Reference temperature. Default set by option <code>tnom</code> .
38	<code>tnom (C)</code>	Alias of <code>tref</code> . Default set by option <code>tnom</code> .
39	<code>dta=0.0 K</code>	Difference between device temperature and ambient temperature.
40	<code>trise=0.0 K</code>	Alias of <code>dta</code> .
41	<code>ptbf=0.0</code>	Power for temperature dependence of <code>bf</code> .
42	<code>ptbr=0.0</code>	Power for temperature dependence of <code>br</code> .
43	<code>ptrc=0.0</code>	Power for temperature dependence of <code>rc</code> .
44	<code>ptrb=0.0</code>	Power for temperature dependence of <code>rbc</code> and <code>rbv</code> .
45	<code>vg=1.2 V</code>	Band-gap voltage.

Spectre Circuit Simulator Reference

Component Statements Part I

46 `pt=1.2` Power for temperature dependence of diffusion coefficient.

Noise model parameters

47 `kf=0.0` Flicker noise coefficient.

48 `af=1.0` Flicker noise exponent.

Output Parameters

1 `ist (A)` Saturation current.

2 `iole (A)` Non-ideal forward base saturation current.

3 `iolc (A)` Non-ideal reverse base saturation current.

4 `bft (A/A)` Ideal forward common-emitter current gain (beta).

5 `brt (A/A)` Ideal reverse common-collector current gain (beta).

6 `rct (Ω)` Collector resistance.

7 `rbct (Ω)` Constant part of base resistance.

8 `rbvt (Ω)` Variable part of base resistance.

9 `taubt (s)` Forward transit time related to neutral base.

10 `cjet (F)` Zero bias emitter-base depletion capacitance.

11 `vdet (V)` Emitter-base diffusion voltage.

12 `taurt (s)` Ideal reverse transit time.

13 `cjct (F)` Zero bias collector-base depletion capacitance.

14 `vdct (V)` Collector-base diffusion voltage.

15 `cjst (F)` Zero bias substrate junction depletion capacitance.

16 `vdst (V)` Substrate junction diffusion voltage.

Spectre Circuit Simulator Reference

Component Statements Part I

Operating-Point Parameters

1	i_b (A)	Base current.
2	i_c (A)	Collector current.
3	i_e (A)	Emitter current.
4	i_{sub} (A)	Substrate current.
5	v_{be} (V)	Base-emitter voltage.
6	v_{bc} (V)	Base-collector voltage.
7	v_{ce} (V)	Collector-emitter voltage.
8	v_{subj} (V)	Substrate voltage.
9	β_{tadc} (A/A)	Ratio of DC collector current to DC Base current.
10	r_b (Ω)	Base resistance at operating point.
11	r_c (Ω)	Collector resistance at operating point.
12	r_e (Ω)	Emitter resistance at operating point.
13	i_{cb} (A)	Collector-Base current.
14	i_{eb} (A)	Emitter-Base current.
15	i_{csub} (A)	Collector-Substrate current.
16	i_{esub} (A)	Emitter-Substrate current.
17	pwr (W)	Power.
18	g_{pi} (S)	Conductance emitter-base junction.
19	g_{mu} (S)	Conductance collector-base junction.
20	g_f (S)	Forward transconductance.
21	g_r (S)	Reverse transconductance.

Spectre Circuit Simulator Reference

Component Statements Part I

22	<code>gs</code> (S)	Conductance substrate-base junction.
23	<code>g3</code> (S)	Transconductance (parasitic PNP) c-b-s transistor.
24	<code>g4</code> (S)	Transconductance (parasitic PNP) e-b-s transistor.
25	<code>ced</code> (F)	Emitter diffusion capacitance.
26	<code>ccd</code> (F)	Collector diffusion capacitance.
27	<code>cet</code> (F)	Emitter junction depletion capacitance.
28	<code>cct</code> (F)	Collector junction depletion capacitance.
29	<code>cst</code> (F)	Substrate junction depletion capacitance.
30	<code>betaac</code> (A/A)	Small-signal common-emitter current gain.
31	<code>ft</code> (Hz)	Unity small-signal current-gain frequency.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

<code>af</code>	M-48	<code>gf</code>	OP-20	<code>mtau</code>	M-24	<code>struct</code>	M-2
<code>area</code>	I-1	<code>gmu</code>	OP-19	<code>mult</code>	I-2	<code>taub</code>	M-22
<code>betaac</code>	OP-30	<code>gpi</code>	OP-18	<code>nhf</code>	M-9	<code>taubt</code>	O-9
<code>betadc</code>	OP-9	<code>gr</code>	OP-21	<code>nlf</code>	M-7	<code>taune</code>	M-23
<code>bf</code>	M-5	<code>gs</code>	OP-22	<code>nlr</code>	M-13	<code>taur</code>	M-28
<code>bft</code>	O-4	<code>ib</code>	OP-1	<code>pc</code>	M-31	<code>taurt</code>	O-12
<code>br</code>	M-11	<code>ic</code>	OP-2	<code>pe</code>	M-27	<code>tnom</code>	M-38
<code>brt</code>	O-5	<code>icb</code>	OP-13	<code>ps</code>	M-34	<code>tref</code>	M-37
<code>ccd</code>	OP-26	<code>icsub</code>	OP-15	<code>pt</code>	M-46	<code>trise</code>	M-40
<code>cct</code>	OP-28	<code>ie</code>	OP-3	<code>ptbf</code>	M-41	<code>type</code>	M-1

Spectre Circuit Simulator Reference Component Statements Part I

ced	OP-25	ieb	OP-14	ptbr	M-42	vbc	OP-6
cet	OP-27	iesub	OP-16	ptrb	M-44	vbe	OP-5
cjc	M-29	ikf	M-8	ptrc	M-43	vce	OP-7
cjct	O-13	ikr	M-14	pwr	OP-17	vdc	M-30
cje	M-25	iks	M-15	rb	OP-10	vdct	O-14
cjet	O-10	ilf	M-6	rbc	M-19	vde	M-26
cjs	M-32	ilr	M-12	rbct	O-7	vdet	O-11
cjst	O-15	imax	M-4	rbv	M-20	vds	M-33
cst	OP-29	iolc	O-3	rbvt	O-8	vdst	O-16
dta	M-39	iole	O-2	rc	M-18	veaf	M-10
exphi	M-35	is	M-3	rc	OP-11	vg	M-45
fc	M-36	ist	O-1	rct	O-6	vsubj	OP-8
ft	OP-31	isub	OP-4	re	OP-12	xcs	M-16
g3	OP-23	kf	M-47	re	M-21	xes	M-17
g4	OP-24	m	I-3	region	I-4		

Lateral PNP Transistor (bjt500)

Description

The `bjt500` model provides an extensive description of a lateral integrated circuit junction-isolated PNP transistor. It is described in the Philips Bipolar Modelbook (Dec.93) as TPL-level-500. Information on how to obtain this document can be found on Source Link by searching for Philips.

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In extension to the modelbook description a minimum conductance `gmin` is inserted between the internal base and internal collector node, between the internal base and the internal emitter node, and between the external base and the substrate node to aid convergence. The value of `gmin` is set by an options statement, default is `gmin = 1.0e-12 S`.

The `imax` parameter is used to aid convergence and to prevent numerical overflow. The junction characteristics of the transistor are accurately modeled for currents up to `imax`. For currents above `imax`, the junction is modeled as a linear resistor, and a warning is printed.

This device is supported within altergroups.

Spectre Circuit Simulator Reference

Component Statements Part I

This device is dynamically loaded from the shared object /vobs/spectre_dev/tools.sun4v/spectre/lib/cmi/3.0.doc/libphilips_sh.so

Sample Instance Statement

```
q3 (minus net3 vcc) pnp_mod region=fwd area=1 m=1
```

Sample Model Statement

```
model pnp_mod bjt500 type=pnp struct=lateral is=1e-14 bf=85 ik=95e-6 reex=3.2  
cje=0.352e-12
```

Instance Definition

```
Name c b e [s] ModelName parameter=value ...
```

Instance Parameters

- | | | |
|---|------------|---|
| 1 | area=1 | Area factor. |
| 2 | mult=1 | Alias of area factor. |
| 3 | m=1 | Multiplicity factor. |
| 4 | region=fwd | Estimated DC operating region, used as a convergence aid.
Possible values are off, fwd, rev, or sat. |

Model Definition

```
model modelName bjt500 parameter=value ...
```

Model Parameters

Structural parameters

- | | | |
|---|----------------|---|
| 1 | type=pnp | Transistor type.
Possible values are pnp or pnp1. |
| 2 | struct=lateral | Transistor structure.
Possible values are lateral. |

Spectre Circuit Simulator Reference

Component Statements Part I

Current parameters

3	$is=1.8e-16$ A	Collector-emitter saturation current.
4	$imax=1.0$ A	Explosion current.
5	$bf=131.0$ A/A	Ideal forward common-emitter current gain (beta).
6	$ibf=2.6e-14$ A	Saturation current of non-ideal forward base current.
7	$vlf=0.54$ V	Cross-over voltage of non-ideal forward base current.
8	$ik=1.1e-4$ A	High injection knee current.
9	$xifv=0.43$	Vertical fraction of forward current.
10	$eafl=20.5$ V	Early voltage of the lateral forward current component.
11	$eafv=75.0$ V	Early voltage of the vertical forward current component.
12	$br=25.0$ A/A	Ideal reverse common-emitter current gain.
13	$ibr=1.2e-13$ A	Saturation current of non-ideal reverse base current.
14	$vlr=0.48$ V	Cross-over voltage of non-ideal reverse base current.
15	$xirv=0.43$	Vertical fraction of reverse current.
16	$earl=13.1$ V	Early voltage of the lateral reverse current component.
17	$earv=104.0$ V	Early voltage of the vertical reverse current component.
18	$xes=2.7e-3$	Ratio between saturation current of e-b-s transistor and e-b-c transistor.
19	$xhes=0.7$	Fraction of substrate current of e-b-s transistor subject to high injection.
20	$xcs=3.0$	Ratio between saturation current of c-b-s transistor and c-b-e transistor.
21	$xhcs=1.0$	Fraction of substrate current of c-b-s transistor subject to high injection.

Spectre Circuit Simulator Reference

Component Statements Part I

22 `iss=4.0e-13 A` Saturation current of substrate-base diode.

Parasitic resistance parameters

23 `rcex=5.0 Ω` External part of the collector resistance.

24 `rcin=47.0 Ω` Internal part of the collector resistance.

25 `rbcc=10.0 Ω` Constant part of the base resistance `rbc`.

26 `rbcv=10.0 Ω` Variable part of the base resistance `rbc`.

27 `rbec=10.0 Ω` Constant part of the base resistance `rbe`.

28 `rbev=50.0 Ω` Variable part of the base resistance `rbe`.

29 `reex=27.0 Ω` External part of the emitter resistance.

30 `rein=66.0 Ω` Internal part of the emitter resistance.

31 `rsb=1.0e15 Ω` Substrate-base leakage resistance.

Junction capacitance parameters

32 `tlat=2.4e-9 s` Low injection (forward and reverse) transit time of charge stored in the epilayer between emitter and collector.

33 `tfvr=3.0e-8 s` Low injection forward transit time due to charge stored in the epilayer under the emitter.

34 `tfn=2.0e-10 s` Low injection forward transit time due to charge stored in the emitter and the buried layer under the emitter.

35 `cje=6.1e-14 F` Zero-bias emitter-base depletion capacitance.

36 `vde=0.52 V` Emitter-base diffusion voltage.

37 `pe=0.3` Emitter-base grading coefficient.

38 `trvr=1.0e-9 s` Low injection reverse transit time due to charge stored in the epilayer under the collector.

Spectre Circuit Simulator Reference

Component Statements Part I

39	<code>trn=3.0e-9 s</code>	Low injection reverse transit time due to charge stored in the collector and the buried layer under the collector.
40	<code>cjc=3.9e-13 F</code>	Zero-bias collector-base depletion capacitance.
41	<code>vdc=0.57 V</code>	Collector-base diffusion voltage.
42	<code>pc=0.36</code>	Collector-base grading coefficient.
43	<code>cjs=1.3e-12 F</code>	Zero-bias substrate-base depletion capacitance.
44	<code>vds=0.52 V</code>	Substrate-base diffusion voltage.
45	<code>ps=0.35</code>	Substrate-base grading coefficient.
46	<code>exphi</code>	Not used in model bjt500.

Temperature effects parameters

47	<code>tref (C)</code>	Reference temperature. Default set by option <code>tnom</code> .
48	<code>tnom (C)</code>	Alias of <code>tref</code> .
49	<code>tr (C)</code>	Alias of <code>tref</code> .
50	<code>dta=0.0 K</code>	Difference between the device temperature and the ambient analysis temperature.
51	<code>trise=0.0 K</code>	Alias of <code>dta</code> .
52	<code>vgeb=1.206 V</code>	Bandgap voltage of the emitter-base depletion region.
53	<code>vgcb=1.206 V</code>	Bandgap voltage of the collector-base depletion region.
54	<code>vg sb=1.206 V</code>	Bandgap voltage of the substrate-base depletion region.
55	<code>vgb=1.206 V</code>	Bandgap voltage of the base between emitter and collector.
56	<code>vge=1.206 V</code>	Bandgap voltage of the emitter.
57	<code>vgje=1.123 V</code>	Bandgap voltage recombination emitter-base junction.

Spectre Circuit Simulator Reference

Component Statements Part I

58	<code>ae=4.48</code>	Temperature coefficient of <code>bf</code> .
59	<code>spb=2.853</code>	Temperature coefficient of the epitaxial base hole mobility.
60	<code>snb=2.6</code>	Temperature coefficient of the epitaxial base electron mobility.
61	<code>snbn=0.3</code>	Temperature coefficient of buried layer electron mobility.
62	<code>spe=0.73</code>	Temperature coefficient of emitter hole mobility.
63	<code>spc=0.73</code>	Temperature coefficient of collector hole mobility.
64	<code>sx=1.0</code>	Temperature coefficient of combined minority carrier mobility in emitter and buried layer.

Noise model parameters

65	<code>kf=0.0</code>	Flicker noise coefficient.
66	<code>af=1.0</code>	Flicker noise exponent.

Output Parameters

1	<code>ist (A)</code>	Collector-emitter saturation current.
2	<code>bft (A/A)</code>	Ideal forward common-emitter current gain (beta).
3	<code>ibft (A)</code>	Saturation current of non-ideal forward base current.
4	<code>ikt (A)</code>	High injection knee current.
5	<code>eaflt (V)</code>	Early voltage of the lateral forward current component.
6	<code>eafvt (V)</code>	Early voltage of the vertical forward current component.
7	<code>brt (A/A)</code>	Ideal reverse common-emitter current gain.
8	<code>ibr (A)</code>	Saturation current of non-ideal reverse base current.
9	<code>earlt (V)</code>	Early voltage of the lateral reverse current component.
10	<code>earvt (V)</code>	Early voltage of the vertical reverse current component.

Spectre Circuit Simulator Reference

Component Statements Part I

11	<code>isst</code> (A)	Saturation current of substrate-base diode.
12	<code>rcint</code> (Ω)	Internal part of the collector resistance.
13	<code>rbcct</code> (Ω)	Constant part of the base resistance <code>rbc</code> .
14	<code>rbcvt</code> (Ω)	Variable part of the base resistance <code>rbc</code> .
15	<code>rbect</code> (Ω)	Constant part of the base resistance <code>rbe</code> .
16	<code>rbev</code> (Ω)	Variable part of the base resistance <code>rbe</code> .
17	<code>reint</code> (Ω)	Internal part of the emitter resistance.
18	<code>tlatt</code> (s)	Low injection (forward and reverse) transit time of charge stored in the epilayer between emitter and collector.
19	<code>tfvrt</code> (s)	Low injection forward transit time due to charge stored in the epilayer under the emitter.
20	<code>tfnt</code> (s)	Low injection forward transit time due to charge stored in the emitter and the buried layer under the emitter.
21	<code>cjet</code> (F)	Zero-bias emitter-base depletion capacitance.
22	<code>vdet</code> (V)	Emitter-base diffusion voltage.
23	<code>trvrt</code> (s)	Low injection reverse transit time due to charge stored in the epilayer under the collector.
24	<code>trnt</code> (s)	Low injection reverse transit time due to charge stored in the collector and the buried layer under the collector.
25	<code>cjct</code> (F)	Zero-bias collector-base depletion capacitance.
26	<code>vdct</code> (V)	Collector-base diffusion voltage.
27	<code>cjst</code> (F)	Zero-bias substrate-base depletion capacitance.
28	<code>vdst</code> (V)	Substrate-base diffusion voltage.

Spectre Circuit Simulator Reference

Component Statements Part I

Operating-Point Parameters

1	ic (A)	Resistive collector current.
2	ib (A)	Resistive base current.
3	ie (A)	Resistive emitter current.
4	isub (A)	Resistive substrate current.
5	iflat (A)	Lateral forward current.
6	irlat (A)	Lateral reverse current.
7	ifver (A)	Vertical forward current.
8	irver (A)	Vertical reverse current.
9	ire (A)	Ideal forward base current.
10	ile (A)	Non-ideal forward base current.
11	ise (A)	Forward substrate current.
12	irc (A)	Ideal reverse base current.
13	ilc (A)	Non-ideal reverse base current.
14	isc (A)	Reverse substrate current.
15	isf (A)	Reverse leakage current of the substrate-base junction.
16	ip (A)	Main current.
17	betadc (A/A)	Ratio of DC collector current to DC base current.
18	vbc (V)	Base-collector voltage.
19	vbe (V)	Base-emitter voltage.
20	vce (V)	Collector-emitter voltage.
21	vsb (V)	Substrate-base voltage.

Spectre Circuit Simulator Reference

Component Statements Part I

22	<code>rcex</code> (Ω)	External part of the collector resistance.
23	<code>rcint</code> (Ω)	Internal part of the collector resistance.
24	<code>reex</code> (Ω)	External part of the emitter resistance.
25	<code>reint</code> (Ω)	Internal part of the emitter resistance.
26	<code>rbc</code> (Ω)	Base resistance under the collector.
27	<code>rbe</code> (Ω)	Base resistance under the emitter.
28	<code>rsb</code> (Ω)	Ohmic leakage across the substrate-base junction.
29	<code>pwr</code> (W)	Power.
30	<code>gfl</code> (S)	Forward conductance, lateral path.
31	<code>grl</code> (S)	Reverse conductance, lateral path.
32	<code>g11v</code> (S)	Forward conductance, vertical path.
33	<code>g12v</code> (S)	Collector Early-effect on I_{fver} .
34	<code>g21v</code> (S)	Emitter Early-effect on I_{rver} .
35	<code>g22v</code> (S)	Reverse conductance, vertical path.
36	<code>gpiv</code> (S)	Conductance emitter-base junction.
37	<code>gmuv</code> (S)	Conductance collector-base junction.
38	<code>gbe</code> (S)	Emitter-side: base conductance B1-B.
39	<code>gibe</code> (S)	Emitter Early-effect on I_{b1b} .
40	<code>gbc</code> (S)	Collector-side: base conductance B2-B.
41	<code>gibc</code> (S)	Collector Early-effect on I_{b2b} .
42	<code>gise</code> (S)	Transconductance (parasitic PNP) e-b-s transistor.
43	<code>gisc</code> (S)	Transconductance (parasitic PNP) c-b-s transistor.

Spectre Circuit Simulator Reference

Component Statements Part I

44	gsb (S)	Conductance substrate-base junction.
45	cpil (F)	Forward diffusion capacitance, lateral path.
46	cipil (F)	Collector Early-effect on Qflat.
47	cpiv (F)	Forward total capacitance, vertical path.
48	cmul (F)	Reverse diffusion capacitance, lateral path.
49	cimul (F)	Emitter Early-effect on Qrlat.
50	cmuv (F)	Reverse total capacitance, vertical path.
51	csb (F)	Total capacitance substrate-base junction.
52	irbe (A)	Ideal total forward base current.
53	irbc (A)	Ideal total reverse base current.
54	irsb (A)	Substrate base leakage resistance current.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

ae	M-58	gibc	OP-41	kf	M-65	tfnt	O-20
af	M-66	gibe	OP-39	m	I-3	tfvr	M-33
area	I-1	gisc	OP-43	mult	I-2	tfvrt	O-19
betadc	OP-17	gise	OP-42	pc	M-42	tlat	M-32
bf	M-5	gmuv	OP-37	pe	M-37	tlatt	O-18
bft	O-2	gpiv	OP-36	ps	M-45	tnom	M-48
br	M-12	grl	OP-31	pwr	OP-29	tr	M-49
brt	O-7	gsb	OP-44	rbc	OP-26	tref	M-47

Spectre Circuit Simulator Reference Component Statements Part I

cimul	OP-49	ib	OP-2	rbcc	M-25	trise	M-51
cipil	OP-46	ibf	M-6	rbcct	O-13	trn	M-39
cjc	M-40	ibft	O-3	rbcv	M-26	trnt	O-24
cjct	O-25	ibr	M-13	rbcvt	O-14	trvr	M-38
cje	M-35	ibrtr	O-8	rbe	OP-27	trvrt	O-23
cjet	O-21	ic	OP-1	rbec	M-27	type	M-1
cjs	M-43	ie	OP-3	rbect	O-15	vbc	OP-18
cjst	O-27	iflat	OP-5	rbev	M-28	vbe	OP-19
cmul	OP-48	ifver	OP-7	rbevt	O-16	vce	OP-20
cmuv	OP-50	ik	M-8	rcex	OP-22	vdc	M-41
cpil	OP-45	ikt	O-4	rcex	M-23	vdct	O-26
cpiv	OP-47	ilc	OP-13	rcin	M-24	vde	M-36
csb	OP-51	ile	OP-10	rcint	O-12	vdet	O-22
dta	M-50	imax	M-4	rcint	OP-23	vds	M-44
eaf1	M-10	ip	OP-16	reex	OP-24	vdst	O-28
eaf1t	O-5	irbc	OP-53	reex	M-29	vgb	M-55
eafv	M-11	irbe	OP-52	region	I-4	vgcb	M-53
eafvt	O-6	irc	OP-12	rein	M-30	vge	M-56
ear1	M-16	ire	OP-9	reint	OP-25	vgeb	M-52
ear1t	O-9	irlat	OP-6	reint	O-17	vgje	M-57
earv	M-17	irsb	OP-54	rsb	OP-28	vgsb	M-54
earvt	O-10	irver	OP-8	rsb	M-31	vlf	M-7
exphi	M-46	is	M-3	snb	M-60	vlr	M-14
g11v	OP-32	isc	OP-14	snbn	M-61	vsb	OP-21
g12v	OP-33	ise	OP-11	spb	M-59	xcs	M-20
g21v	OP-34	isf	OP-15	spc	M-63	xes	M-18
g22v	OP-35	iss	M-22	spe	M-62	xhcs	M-21
gbc	OP-40	isst	O-11	struct	M-2	xhes	M-19
gbe	OP-38	ist	O-1	sx	M-64	xifv	M-9
gfl	OP-30	isub	OP-4	tfn	M-34	xirv	M-15

Vertical NPN/PNP Transistor (bjt503)

Description

The bjt503 model provides a detailed description of a vertical integrated NPN and PNP transistor. It is described in the Philips Bipolar Modelbook (Dec.95) as TN/TNS and TP/TPS level 503.

The NPN is also described in Nat.Lab. Unclassified Report Nr. 006/94 as Mextram Bipolar Transistor Model. Information on how to obtain this document can be found on Source Link by searching for Philips.

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In addition to the model description a `level` parameter is added. Via the `level` parameter the user can switch between Philips Bipolar Modelbook (Dec.95) and Philips Bipolar Modelbook (Dec.94).

The `imax` parameter is used to aid convergence and to prevent numerical overflow. The junction characteristics of the transistor are accurately modeled for currents up to `imax`. For currents above `imax`, the junction is modeled as a linear resistor and a warning is printed.

The descriptions of the operating point derivatives are given for the NPN type. For the PNP type the terminal voltage in the descriptions has to be exchanged. E.g.:

NPN: $g_x = dI_n/dV_{b2e1}$

PNP: $g_x = dI_n/dV_{e1b2}$

This device is supported within altergroups.

This device is dynamically loaded from the shared object `/vobs/spectre_dev/tools.sun4v/spectre/lib/cmi/3.0.doc/libphilips_sh.so`

Sample Instance Statement

```
q4 (vcc net3 minus) npn_mod region=fwd m=1 mult=1
```

Sample Model Statement

```
model npn_mod bjt503 type=npn level=2 exmod=1 is=1e-14 bf=85 ik=95e-6 rbc=50  
cje=0.352e-12
```

Spectre Circuit Simulator Reference

Component Statements Part I

Instance Definition

Name c b e [s] ModelName parameter=value ...

Instance Parameters

- | | | |
|---|------------|---|
| 1 | area=1 | Area factor. |
| 2 | mult=1 | Alias of area factor. |
| 3 | m=1 | Multiplication factor. |
| 4 | region=fwd | Estimated DC operating region, used as a convergence aid.
Possible values are <code>off</code> , <code>fwd</code> , <code>rev</code> , or <code>sat</code> . |

Model Definition

model modelName bjt503 parameter=value ...

Model Parameters

- | | | |
|----|---------------|--|
| 1 | type=npn | Transistor type.
Possible values are <code>npn</code> , <code>npnv</code> , <code>pnv</code> , or <code>pnpv</code> . |
| 2 | level=2.0 | Transistor Level. Possible values are 1 (Philips Bipolar Modelbook Dec.94) or 2 (Philips Bipolar Modelbook Dec.95). |
| 3 | exmod=0 | Flag for extended modeling of the reverse current gain. |
| 4 | exphi=0 | Flag for distributed high frequency effects. |
| 5 | exavl=1 | Flag for extended modeling of avalanche currents. |
| 6 | is=5.0e-17 A | Collector-emitter saturation current. |
| 7 | bf=140.0 A/A | Ideal forward current gain. |
| 8 | xibi=0.0 | Fraction of ideal base current that belongs to the sidewall. |
| 9 | ibf=2.0e-14 A | Saturation current of the non-ideal forward base current. |
| 10 | vlf=0.5 V | Cross-over voltage of the non-ideal forward base current. |

Spectre Circuit Simulator Reference

Component Statements Part I

11	$ik=15.0e-3$ A	High-injection knee current.
12	$bri=16.0$ A/A	Ideal reverse current gain.
13	$ibr=8.0e-15$ A	Saturation current of the non-ideal reverse base current.
14	$vlr=0.5$ V	Cross-over voltage of the non-ideal reverse base current.
15	$xext=0.5$	Part of I_{ex} , Q_{ex} , Q_{tex} and I_{sub} that depends on V_{bc1} .
16	$qbo=1.2e-12$ Coul	Base charge at zero bias.
17	$eta=4.0$	Factor of the built-in field of the base.
18	$avl=50.0$	Weak avalanche parameter.
19	$efi=0.7$	Electric field intercept (with $exavl=1$).
20	$ihc=3.0e-3$ A	Critical current for hot carriers.
21	$rcc=25.0$ Ω	Constant part of the collector resistance.
22	$rcv=750.0$ Ω	Resistance of the unmodulated epilayer.
23	$sccv=1000.0$ Ω	Space charge resistance of the epilayer.
24	$sfh=0.6$	Current spreading factor epilayer.
25	$rbc=50.0$ Ω	Constant part of the base resistance.
26	$rbv=100.0$ Ω	Variable part of the base resistance at zero bias.
27	$re=2.0$ Ω	Emitter series resistance.
28	$taune=3.0e-10$ s	Minimum delay time of neutral and emitter charge.
29	$mtau=1.18$	Non-ideality factor of the neutral and emitter charge.
30	$cje=2.5e-13$ F	Zero bias emitter-base depletion capacitance.
31	$vde=0.9$ V	Emitter-base diffusion voltage.
32	$pe=0.33$	Emitter-base grading coefficient.

Spectre Circuit Simulator Reference

Component Statements Part I

33	<code>xcje=0.5</code>	Fraction of the e-b depletion cap. that belongs to the sidewall.
34	<code>cjc=1.3e-13 F</code>	Zero bias collector-base depletion capacitance.
35	<code>vdc=0.6 V</code>	Collector-base diffusion voltage.
36	<code>pc=0.4</code>	Collector-base grading coefficient variable part.
37	<code>xp=0.2</code>	Constant part of <code>cjc</code> .
38	<code>mc=0.5</code>	Collector current modulation coefficient.
39	<code>xcjc=0.1</code>	Fraction of the collector-base depletion cap. under the emitter area.
40	<code>tref (C)</code>	Reference temperature. Default set by option <code>tnom</code> .
41	<code>tnom (C)</code>	Alias of <code>tref</code> . Default set by option <code>tnom</code> .
42	<code>tr (C)</code>	Alias of <code>tref</code> . Default set by option <code>tnom</code> .
43	<code>dta=0.0 K</code>	Difference of the device temperature to the ambient temperature.
44	<code>trise=0.0 K</code>	Alias of <code>dta</code> .
45	<code>vge=1.01 V</code>	Band-gap voltage of the emitter.
46	<code>vgb=1.18 V</code>	Band-gap voltage of the base.
47	<code>vgc=1.205 V</code>	Band-gap voltage of the collector.
48	<code>vgj=1.1 V</code>	Band-gap voltage recombination emitter-base junction.
49	<code>vi=0.04 V</code>	Ionization voltage base dope.
50	<code>na=3.0e17 cm⁻³</code>	Maximum base dope concentration.
51	<code>er=2.0e-3</code>	Temperature coefficient of <code>vlf</code> and <code>vlr</code> .
52	<code>ab=1.35</code>	Temperature coefficient resistivity base.
53	<code>aepi=2.15</code>	Temperature coefficient resistivity of the epilayer.

Spectre Circuit Simulator Reference

Component Statements Part I

54	<code>aex=1.0</code>	Temperature coefficient resistivity of the extrinsic base.
55	<code>ac=0.4</code>	Temperature coefficient resistivity of the buried layer.
56	<code>kf=2.0e-16</code>	Flickernoise coefficient ideal base current.
57	<code>kfn=2.0e-16</code>	Flickernoise coefficient non-ideal base current.
58	<code>af=1.0</code>	Flickernoise exponent.
59	<code>iss=6.0e-16 A</code>	Base-substrate saturation current.
60	<code>iks=5.0e-6 A</code>	Knee current of the substrate.
61	<code>cjs=1.0e-12 F</code>	Zero bias collector-substrate depletion capacitance.
62	<code>vds=0.5 V</code>	Collector-substrate diffusion voltage.
63	<code>ps=0.33</code>	Collector-substrate grading coefficient.
64	<code>vgs=1.15 V</code>	Band-gap voltage of the substrate.
65	<code>as=2.15</code>	For a closed buried layer: $as=ac$. For an open buried layer: $as=ae\pi$.
66	<code>imax=1.0 A</code>	Explosion current.

Output Parameters

1	<code>ist (A)</code>	Collector-Emitter saturation current.
2	<code>bft (A/A)</code>	Ideal forward current gain.
3	<code>ibft (A)</code>	Saturation current of the non-ideal forward base current.
4	<code>vlft (V)</code>	Cross-over voltage of the non-ideal forward base current.
5	<code>ikt (A)</code>	High-injection knee current.
6	<code>ibr (A)</code>	Saturation current of the non-ideal reverse base current.
7	<code>vlr (V)</code>	Cross-over voltage of the non-ideal reverse base current.

Spectre Circuit Simulator Reference

Component Statements Part I

8	qbot (Coul)	Base charge at zero bias.
9	avlt	Weak avalanche parameter.
10	rcct (Ω)	Constant part of the collector resistance.
11	rcvt (Ω)	Resistance of the unmodulated epilayer.
12	rbct (Ω)	Constant part of the base resistance.
13	rbvt (Ω)	Variable part of the base resistance at zero bias.
14	taunet (s)	Minimum delay time of neutral and emitter charge.
15	mtaut	Non-ideality factor of the neutral and emitter charge.
16	cjet (F)	Zero bias emitter-base depletion capacitance.
17	vdet (V)	Emitter-base diffusion voltage.
18	cjct (F)	Zero bias collector-base depletion capacitance.
19	vdct (V)	Collector-base diffusion voltage.
20	xpt	Constant part of c_{jc} .
21	isst (A)	Base-substrate saturation current.
22	ikst (A)	Knee current of the substrate.
23	cjst (F)	Zero bias collector-substrate depletion capacitance.
24	vdst (V)	Collector-substrate diffusion voltage.

Operating-Point Parameters

1	ib (A)	Base current.
2	ic (A)	Collector current.
3	ie (A)	Emitter current.
4	is (A)	Substrate current.

Spectre Circuit Simulator Reference

Component Statements Part I

5	vbe (V)	Base-emitter voltage.
6	vbc (V)	Base-collector voltage.
7	vce (V)	Collector-emitter voltage.
8	vsc (V)	Substrate voltage.
9	re (Ω)	Constant emitter resistance.
10	rcc (Ω)	Constant collector resistance.
11	rbc (Ω)	Constant part of base resistance.
12	betadc (A/A)	DC current gain.
13	pwr (W)	Power.
14	Vb1e1 (V)	Internal voltage.
15	Vb2e1 (V)	Internal voltage.
16	Vb2c1 (V)	Internal voltage.
17	Vb2c2 (V)	Internal voltage.
18	Vb1b2 (V)	Internal voltage.
19	Vb1c1 (V)	Internal voltage.
20	Vbc1 (V)	Internal voltage.
21	in (A)	Main current.
22	ic1c2 (A)	Variable collector resistance current.
23	ib1 (A)	Bulk component of ideal base current.
24	ib1s (A)	Sidewall component of ideal base current.
25	ib2 (A)	Non-ideal base current.
26	iav1 (A)	Weak avalanche current.

Spectre Circuit Simulator Reference

Component Statements Part I

27	ib1b2 (A)	Variable base resistance current.
28	ib3 (A)	Non-ideal reverse base current.
29	iex (A)	Internal extrinsic base current.
30	isub (A)	Internal base-substrate current.
31	isf (A)	Substrate-collector current.
32	xiex (A)	External extrinsic base current.
33	xisub (A)	External base-substrate current.
34	gx (S)	dI_n/dV_{b2e1} .
35	gy (S)	dI_n/dV_{b2c2} .
36	gz (S)	dI_n/dV_{b2c1} .
37	grcvy (S)	dI_{c1c2}/dV_{b2c2} .
38	grcvz (S)	dI_{c1c2}/dV_{b2c1} .
39	gpi (S)	Conductance floor base-emitter junction: $dI_{b1}/dV_{b2e1} + dI_{b2}/dV_{b2e1}$.
40	sgpi (S)	Conductance sidewall base-emitter junction: dI_{b1S}/dV_{b1e1} .
41	gmux (S)	Dependence avalanche multiplication on internal b-e junction: $-dI_{avl}/dV_{b2e1}$.
42	gmu (S)	Dependence avalanche multiplication on internal b-c junction: $-dI_{avl}/dV_{b2c2}$.
43	gmuz (S)	Dependence avalanche multiplication on external b-c junction: $-dI_{avl}/dV_{b2c1}$.
44	grbv (S)	dI_{b1b2}/dV_{b1b2} .
45	grbvX (S)	Emitter Early-effect on I_{b1b2} : dI_{b1b2}/dV_{b2e1} .
46	grbvY (S)	Internal collector Early-effect on I_{b1b2} : dI_{b1b2}/dV_{b2c2} .

Spectre Circuit Simulator Reference

Component Statements Part I

47	<code>grbvz</code> (S)	External collector Early effect on I_{b1b2} : dI_{b1b2}/dV_{b2c1} .
48	<code>gmuex</code> (S)	Conductance floor extrinsic b-c junction: $dI_{lex}/dV_{b1c1} + dI_{sub}/dV_{b1c1} + dI_{b3}/dV_{b1c1}$.
49	<code>xgmuex</code> (S)	Conductance sidewall extrinsic b-c junction: $dXI_{lex}/dV_{bc1} + dXI_{sub}/dV_{bc1}$.
50	<code>gsub</code> (S)	Conductance s-c junction: dI_{sf}/dV_{sc1} .
51	<code>gpnp</code> (S)	Transconductance floor extrinsic PNP transistor: dI_{sub}/dV_{b1c1} .
52	<code>xgpnp</code> (S)	Transconductance sidewall extrinsic PNP transistor: dXI_{sub}/dV_{bc1} .
53	<code>cbex</code> (F)	Capacitance floor b-e junction: $dQ_{te}/dV_{b2e1} + dQ_{be}/dV_{b2e1} + dQ_n/dV_{b2e1}$.
54	<code>cbey</code> (F)	Internal collector Early-effect on Q_{be} : dQ_{be}/dV_{b2c2} .
55	<code>cbez</code> (F)	External collector Early-effect on Q_{be} : dQ_{be}/dV_{b2c1} .
56	<code>scte</code> (F)	Dependence of Q_{teS} on internal b-e junction: dQ_{teS}/dV_{b2e1} .
57	<code>cbcx</code> (F)	Emitter Early-effect on Q_{bc} : dQ_{bc}/dV_{b2e1} .
58	<code>cbcy</code> (F)	Capacitance intrinsic b-c junction: $dQ_{tc}/dV_{b2c2} + dQ_{bc}/dV_{b2c2} + dQ_{epi}/dV_{b2c2}$.
59	<code>cbcZ</code> (F)	Collector Early-effect on Q_{tc} : $dQ_{tc}/dV_{b2c1} + dQ_{bc}/dV_{b2c1} + dQ_{epi}/dV_{b2c1}$.
60	<code>cb1b2</code> (F)	Capacitance AC current crowding: $dQ_{b1b2}/dV_{b1b2} = C_b$.
61	<code>cb1b2x</code> (F)	Dependence of Q_{b1b2} on internal b-e junction voltage: dQ_{b1b2}/dV_{b2e1} .
62	<code>cbceX</code> (F)	Capacitance floor extrinsic b-c junction: $dQ_{tex}/dV_{b1c1} + dQ_{ex}/dV_{b1c1}$.
63	<code>xcbceX</code> (F)	Capacitance sidewall extrinsic b-c junction: $dXQ_{tex}/dV_{bc1} + dXQ_{ex}/dV_{bc1}$.

Spectre Circuit Simulator Reference

Component Statements Part I

64 cts (F)

Capacitance s-c junction: $dQ_{tex}/dV_{b1c1} + dQ_{ex}/dV_{b1c1}$.

Parameter Index

In the following index, I refers to instance parameters, M refers to the model parameters section, O refers to the output parameters section, and OP refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of M-35 means the 35th model parameter.

Vb1b2	OP-18	eta	M-17	imax	M-66	sgpi	OP-40
Vb1c1	OP-19	exavl	M-5	in	OP-21	taune	M-28
Vb1e1	OP-14	exmod	M-3	is	OP-4	taunet	O-14
Vb2c1	OP-16	exphi	M-4	is	M-6	tnom	M-41
Vb2c2	OP-17	gmu	OP-42	isf	OP-31	tr	M-42
Vb2e1	OP-15	gmux	OP-48	iss	M-59	tref	M-40
Vbc1	OP-20	gmux	OP-41	isst	O-21	trise	M-44
Xisub	OP-33	gmuz	OP-43	ist	O-1	type	M-1
ab	M-52	gpi	OP-39	isub	OP-30	vbc	OP-6
ac	M-55	gpnv	OP-51	kf	M-56	vbe	OP-5
aepi	M-53	grbv	OP-44	kfn	M-57	vce	OP-7
aex	M-54	grbvz	OP-45	level	M-2	vdc	M-35
af	M-58	grbvz	OP-46	m	I-3	vdct	O-19
area	I-1	grcvz	OP-47	mc	M-38	vde	M-31
as	M-65	grcvz	OP-37	mtau	M-29	vdet	O-17
avl	M-18	grcvz	OP-38	mtaut	O-15	vds	M-62
avlt	O-9	gsub	OP-50	mult	I-2	vdst	O-24
betadc	OP-12	gx	OP-34	na	M-50	vgb	M-46
bf	M-7	gy	OP-35	pc	M-36	vgc	M-47
bft	O-2	gz	OP-36	pe	M-32	vge	M-45
bri	M-12	iavl	OP-26	ps	M-63	vgj	M-48
cb1b2	OP-60	ib	OP-1	pwr	OP-13	vgs	M-64
cb1b2x	OP-61	ib1	OP-23	qbo	M-16	vi	M-49

Spectre Circuit Simulator Reference Component Statements Part I

cbcx	OP-62	ib1b2	OP-27	qbot	O-8	vlf	M-10
cbcx	OP-57	ib1s	OP-24	rbc	OP-11	vlft	O-4
cbcy	OP-58	ib2	OP-25	rbc	M-25	vlr	M-14
cbcz	OP-59	ib3	OP-28	rbct	O-12	vlrt	O-7
cbex	OP-53	ibf	M-9	rbv	M-26	vsc	OP-8
cbey	OP-54	ibft	O-3	rbvt	O-13	xcbcex	OP-63
cbez	OP-55	ibr	M-13	rcc	OP-10	xcjc	M-39
cjc	M-34	ibr	O-6	rcc	M-21	xcje	M-33
cjct	O-18	ic	OP-2	rcct	O-10	xext	M-15
cje	M-30	ic1c2	OP-22	rcv	M-22	xgmux	OP-49
cjet	O-16	ie	OP-3	rcvt	O-11	xgpn	OP-52
cjs	M-61	iex	OP-29	re	M-27	xibi	M-8
cjst	O-23	ihc	M-20	re	OP-9	xiex	OP-32
cts	OP-64	ik	M-11	region	I-4	xp	M-37
dta	M-43	iks	M-60	scrcv	M-23	xpt	O-20
efi	M-19	ikst	O-22	scte	OP-56		
er	M-51	ikt	O-5	sfh	M-24		

Vertical NPN/PNP Transistor (bjt504)

Description

The bjt504T model provides a detailed description of a vertical integrated NPN and PNP transistor. It is described in the Philips Bipolar Modelbook (Dec.95) as TN/TNS and TP/TPS level 504.

The NPN is also described in Nat.Lab. Unclassified Report Nr. 006/94 as Mextram Bipolar Transistor Model. Information on how to obtain this document can be found on Source Link by searching for Philips.

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In addition to the model description a `level` parameter is added. Via the `level` parameter the user can switch between Philips Bipolar Modelbook (Dec.95) and Philips Bipolar Modelbook (Dec.94).

Spectre Circuit Simulator Reference

Component Statements Part I

The `imax` parameter is used to aid convergence and to prevent numerical overflow. The junction characteristics of the transistor are accurately modeled for currents up to `imax`. For currents above `imax`, the junction is modeled as a linear resistor and a warning is printed.

The descriptions of the operating point derivatives are given for the NPN type. For the PNP type the terminal voltage in the descriptions has to be exchanged. E.g.:

NPN: $g_x = dI_n/dV_{b2e1}$

PNP: $g_x = dI_n/dV_{e1b2}$

This device is supported within altergroups.

This device is dynamically loaded from the shared object `/vobs/spectre_dev/tools.sun4v/spectre/lib/cmi/3.0.doc/libphilips_sh.so`

Instance Definition

Name `c b e [s] ... ModelName parameter=value ...`

Options:

[s] When `s` is not specified, it is bound to ground;

`dt` not used (only used in the selfheating version of the model). Do not specify.

Instance Parameters

1	<code>level=X</code>	<code>level</code> .
2	<code>mult=X</code>	<code>mult</code> .
3	<code>tref=X</code>	<code>tref</code> .
4	<code>dta=X</code>	<code>dta</code> .
5	<code>exmod=X</code>	<code>exmod</code> .
6	<code>exphi=X</code>	<code>exphi</code> .
7	<code>exavl=X</code>	<code>exavl</code> .

Spectre Circuit Simulator Reference

Component Statements Part I

8	is=X	is.
9	ik=X	ik.
10	ver=X	ver.
11	vef=X	vef.
12	bf=X	bf.
13	ibf=X	ibf.
14	mlf=X	mlf.
15	xibi=X	xibi.
16	bri=X	bri.
17	ibr=X	ibr.
18	vlr=X	vlr.
19	xext=X	xext.
20	wavl=X	wavl.
21	vavl=X	vavl.
22	sfh=X	sfh.
23	re=X	re.
24	rbc=X	rbc.
25	rbv=X	rbv.
26	rcc=X	rcc.
27	rcv=X	rcv.
28	scrcv=X	scrcv.
29	ihc=X	ihc.

Spectre Circuit Simulator Reference

Component Statements Part I

30	axi=X	axi.
31	cje=X	cje.
32	vde=X	vde.
33	pe=X	pe.
34	xcje=X	xcje.
35	cbeo=X	cbeo.
36	cjc=X	cjc.
37	vdc=X	vdc.
38	pc=X	pc.
39	xp=X	xp.
40	mc=X	mc.
41	xcjc=X	xcjc.
42	cbco=X	cbco.
43	mtau=X	mtau.
44	taue=X	taue.
45	taub=X	taub.
46	tepi=X	tepi.
47	taur=X	taur.
48	deg=X	deg.
49	xrec=X	xrec.
50	aqbo=X	aqbo.
51	ae=X	ae.

Spectre Circuit Simulator Reference

Component Statements Part I

52	ab=X	ab.
53	aepi=X	aepi.
54	aex=X	aex.
55	ac=X	ac.
56	dvgbf=X	dvgbf.
57	dvgbr=X	dvgbr.
58	vgb=X	vgb.
59	vgc=X	vgc.
60	vgj=X	vgj.
61	dvgte=X	dvgte.
62	af=X	af.
63	kf=X	kf.
64	kfn=X	kfn.
65	iss=X	iss.
66	iks=X	iks.
67	cjs=X	cjs.
68	vds=X	vds.
69	ps=X	ps.
70	vgs=X	vgs.
71	as=X	as.
72	region=fwd	Estimated DC operating region, used as a convergence aid. Possible values are off, fwd, rev, or sat.

Spectre Circuit Simulator Reference

Component Statements Part I

- 73 `m=1` Multiplication factor.
- 74 `area=1` Alias of mult.

Model Definition

```
model modelName bjt504 parameter=value ...
```

Model Parameters

- 1 `level=NULL` level.
- 2 `mult=NULL` mult.
- 3 `tref=NULL` tref.
- 4 `dta=NULL` dta.
- 5 `exmod=NULL` exmod.
- 6 `exphi=NULL` exphi.
- 7 `exavl=NULL` exavl.
- 8 `is=NULL` is.
- 9 `ik=NULL` ik.
- 10 `ver=NULL` ver.
- 11 `vef=NULL` vef.
- 12 `bf=NULL` bf.
- 13 `ibf=NULL` ibf.
- 14 `mlf=NULL` mlf.
- 15 `xibi=NULL` xibi.
- 16 `bri=NULL` bri.
- 17 `ibr=NULL` ibr.

Spectre Circuit Simulator Reference

Component Statements Part I

18	vlr=NULL	vlr.
19	xext=NULL	xext.
20	wavl=NULL	wavl.
21	vavl=NULL	vavl.
22	sfh=NULL	sfh.
23	re=NULL	re.
24	rbc=NULL	rbc.
25	rbv=NULL	rbv.
26	rcc=NULL	rcc.
27	rcv=NULL	rcv.
28	scrcv=NULL	scrcv.
29	ihc=NULL	ihc.
30	axi=NULL	axi.
31	cje=NULL	cje.
32	vde=NULL	vde.
33	pe=NULL	pe.
34	xcje=NULL	xcje.
35	cbeo=NULL	cbeo.
36	cjc=NULL	cjc.
37	vdc=NULL	vdc.
38	pc=NULL	pc.
39	xp=NULL	xp.

Spectre Circuit Simulator Reference

Component Statements Part I

40	mc=NULL	mc.
41	xcjc=NULL	xcjc.
42	cbco=NULL	cbco.
43	mtau=NULL	mtau.
44	taue=NULL	taue.
45	taub=NULL	taub.
46	tepi=NULL	tepi.
47	taur=NULL	taur.
48	deg=NULL	deg.
49	xrec=NULL	xrec.
50	aqbo=NULL	aqbo.
51	ae=NULL	ae.
52	ab=NULL	ab.
53	aepi=NULL	aepi.
54	aex=NULL	aex.
55	ac=NULL	ac.
56	dvgbf=NULL	dvgbf.
57	dvgbr=NULL	dvgbr.
58	vgb=NULL	vgb.
59	vgc=NULL	vgc.
60	vgj=NULL	vgj.
61	dvgte=NULL	dvgte.

Spectre Circuit Simulator Reference

Component Statements Part I

62	af=NULL	af.
63	kf=NULL	kf.
64	kfn=NULL	kfn.
65	iss=NULL	iss.
66	iks=NULL	iks.
67	cjs=NULL	cjs.
68	vds=NULL	vds.
69	ps=NULL	ps.
70	vgs=NULL	vgs.
71	as=NULL	as.
72	type=npn	Transistor type. Possible values are npn or pnp.
73	imax=1.0 A	Explosion current.
74	tnom=25 C	alias of tref. Default set by option tnom.
75	tr (C)	Alias of tref. Default set by option tnom.

Operating-Point Parameters

1	level=X	level.
2	mult=X	mult.
3	tref=X	tref.
4	dta=X	dta.
5	exmod=X	exmod.
6	exphi=X	exphi.

Spectre Circuit Simulator Reference

Component Statements Part I

7	exavl=X	exavl.
8	is=X	is.
9	ik=X	ik.
10	ver=X	ver.
11	vef=X	vef.
12	bf=X	bf.
13	ibf=X	ibf.
14	mlf=X	mlf.
15	xibi=X	xibi.
16	bri=X	bri.
17	ibr=X	ibr.
18	vlr=X	vlr.
19	xext=X	xext.
20	wavl=X	wavl.
21	vavl=X	vavl.
22	sfh=X	sfh.
23	re=X	re.
24	rbc=X	rbc.
25	rbv=X	rbv.
26	rcc=X	rcc.
27	rcv=X	rcv.
28	scrcv=X	scrcv.

Spectre Circuit Simulator Reference

Component Statements Part I

29	ihc=X	ihc.
30	axi=X	axi.
31	cje=X	cje.
32	vde=X	vde.
33	pe=X	pe.
34	xcje=X	xcje.
35	cbeo=X	cbeo.
36	cjc=X	cjc.
37	vdc=X	vdc.
38	pc=X	pc.
39	xp=X	xp.
40	mc=X	mc.
41	xcjc=X	xcjc.
42	cbco=X	cbco.
43	mtau=X	mtau.
44	taue=X	taue.
45	taub=X	taub.
46	tepi=X	tepi.
47	taur=X	taur.
48	deg=X	deg.
49	xrec=X	xrec.
50	aqbo=X	aqbo.

Spectre Circuit Simulator Reference

Component Statements Part I

51	ae=X	ae.
52	ab=X	ab.
53	aepi=X	aepi.
54	aex=X	aex.
55	ac=X	ac.
56	dvgbf=X	dvgbf.
57	dvgbr=X	dvgbr.
58	vgb=X	vgb.
59	vgc=X	vgc.
60	vgj=X	vgj.
61	dvgte=X	dvgte.
62	af=X	af.
63	kf=X	kf.
64	kfn=X	kfn.
65	iss=X	iss.
66	iks=X	iks.
67	cjs=X	cjs.
68	vds=X	vds.
69	ps=X	ps.
70	vgs=X	vgs.
71	as=X	as.
72	Vb2e1	Vb2e1.

Spectre Circuit Simulator Reference

Component Statements Part I

73	Vb2c2	Vb2c2.
74	Vb2c1	Vb2c1.
75	Vb1c1	Vb1c1.
76	Ve1e	Ve1e.
77	In	In.
78	Ic1c2	Ic1c2.
79	Ib1b2	Ib1b2.
80	Ib1	Ib1.
81	SIb1	SIb1.
82	Ib2	Ib2.
83	Ib3	Ib3.
84	Iex	Iex.
85	XIex	XIex.
86	Iavl	Iavl.
87	IRE	IRE.
88	IRBC	IRBC.
89	IRCC	IRCC.
90	Qe	Qe.
91	Qte	Qte.
92	SQte	SQte.
93	Qbe	Qbe.
94	Qbc	Qbc.

Spectre Circuit Simulator Reference

Component Statements Part I

95	Qtc	Qtc.
96	Qepi	Qepi.
97	Qb1b2	Qb1b2.
98	Qtex	Qtex.
99	XQtex	XQtex.
100	Qex	Qex.
101	XQex	XQex.
102	gx	gx.
103	gy	gy.
104	gz	gz.
105	Sgpi	Sgpi.
106	gpix	gpix.
107	gpiy	gpiy.
108	gpiz	gpiz.
109	gmux	gmux.
110	gmuy	gmuy.
111	gmuz	gmuz.
112	gmuex	gmuex.
113	Xgmuex	Xgmuex.
114	grcvy	grcvy.
115	grcvz	grcvz.
116	Rbv	Rbv.

Spectre Circuit Simulator Reference

Component Statements Part I

117	grbv _x	grbv _x .
118	grbv _y	grbv _y .
119	grbv _z	grbv _z .
120	RE	RE.
121	RBC	RBC.
122	RCC	RCC.
123	SCbe	SCbe.
124	Cbex	Cbex.
125	Cbey	Cbey.
126	Cbez	Cbez.
127	Cbc _x	Cbc _x .
128	Cbc _y	Cbc _y .
129	Cbc _z	Cbc _z .
130	Cbcex	Cbcex.
131	XCbcex	XCbcex.
132	Cb1b2	Cb1b2.
133	Cb1b2 _x	Cb1b2 _x .
134	Cb1b2 _y	Cb1b2 _y .
135	Cb1b2 _z	Cb1b2 _z .
136	gm	gm.
137	beta	beta.
138	gout	gout.

Spectre Circuit Simulator Reference

Component Statements Part I

139	gmu	gmu.
140	RB	RB.
141	Cbe	Cbe.
142	Cbc	Cbc.
143	fT	fT.
144	Iqs	Iqs.
145	XiWepi	XiWepi.
146	Vb2c2star	Vb2c2star.
147	Pdiss	Pdiss.
148	TK	TK.
149	Isub	Isub.
150	XIsub	XIsub.
151	I _{sf}	I _{sf} .
152	Q _{ts}	Q _{ts} .
153	g _S	g _S .
154	Xg _S	Xg _S .
155	g _{Sf}	g _{Sf} .
156	C _{ts}	C _{ts} .

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the

Spectre Circuit Simulator Reference Component Statements Part I

description for that parameter. For example, a reference of M-35 means the 35th model parameter.

Cb1b2	OP-132	aex	I-54	grcvz	OP-115	sfh	M-22
Cb1b2x	OP-133	af	I-62	gx	OP-102	sfh	OP-22
Cb1b2y	OP-134	af	OP-62	gy	OP-103	taub	OP-45
Cb1b2z	OP-135	af	M-62	gz	OP-104	taub	I-45
Cbc	OP-142	aqbo	OP-50	ibf	I-13	taub	M-45
Cbcex	OP-130	aqbo	M-50	ibf	M-13	taue	I-44
Cbcx	OP-127	aqbo	I-50	ibf	OP-13	taue	OP-44
Cbcy	OP-128	area	I-74	ibr	I-17	taue	M-44
Cbcz	OP-129	as	OP-71	ibr	M-17	taur	M-47
Cbe	OP-141	as	I-71	ibr	OP-17	taur	OP-47
Cbex	OP-124	as	M-71	ihc	M-29	taur	I-47
Cbey	OP-125	axi	M-30	ihc	OP-29	tepi	OP-46
Cbez	OP-126	axi	OP-30	ihc	I-29	tepi	I-46
Cts	OP-156	axi	I-30	ik	OP-9	tepi	M-46
IRBC	OP-88	beta	OP-137	ik	M-9	tnom	M-74
IRCC	OP-89	bf	M-12	ik	I-9	tr	M-75
IRE	OP-87	bf	I-12	iks	OP-66	tref	I-3
Iavl	OP-86	bf	OP-12	iks	I-66	tref	OP-3
Ib1	OP-80	bri	M-16	iks	M-66	tref	M-3
Ib1b2	OP-79	bri	I-16	imax	M-73	type	M-72
Ib2	OP-82	bri	OP-16	is	I-8	vavl	M-21
Ib3	OP-83	cbco	M-42	is	M-8	vavl	I-21
Ic1c2	OP-78	cbco	I-42	is	OP-8	vavl	OP-21
Iex	OP-84	cbco	OP-42	iss	I-65	vdc	I-37
In	OP-77	cbeo	OP-35	iss	M-65	vdc	OP-37
Iqs	OP-144	cbeo	M-35	iss	OP-65	vdc	M-37
Isf	OP-151	cbeo	I-35	kf	OP-63	vde	I-32
Isub	OP-149	cjc	I-36	kf	M-63	vde	M-32
Pdiss	OP-147	cjc	OP-36	kf	I-63	vde	OP-32

Spectre Circuit Simulator Reference Component Statements Part I

Qb1b2	OP-97	cjc	M-36	kfn	M-64	vds	I-68
Qbc	OP-94	cje	M-31	kfn	I-64	vds	OP-68
Qbe	OP-93	cje	OP-31	kfn	OP-64	vds	M-68
Qe	OP-90	cje	I-31	level	OP-1	vef	I-11
Qepi	OP-96	cjs	M-67	level	I-1	vef	M-11
Qex	OP-100	cjs	I-67	level	M-1	vef	OP-11
Qtc	OP-95	cjs	OP-67	m	I-73	ver	OP-10
Qte	OP-91	deg	OP-48	mc	OP-40	ver	I-10
Qtex	OP-98	deg	I-48	mc	M-40	ver	M-10
Qts	OP-152	deg	M-48	mc	I-40	vgb	M-58
RB	OP-140	dta	I-4	mlf	OP-14	vgb	I-58
RBC	OP-121	dta	M-4	mlf	I-14	vgb	OP-58
RCC	OP-122	dta	OP-4	mlf	M-14	vgc	M-59
RE	OP-120	dvgbf	M-56	mtau	M-43	vgc	I-59
Rbv	OP-116	dvgbf	OP-56	mtau	OP-43	vgc	OP-59
SCbe	OP-123	dvgbf	I-56	mtau	I-43	vgj	OP-60
SIb1	OP-81	dvgbr	OP-57	mult	OP-2	vgj	M-60
SQte	OP-92	dvgbr	I-57	mult	I-2	vgj	I-60
Sgpi	OP-105	dvgbr	M-57	mult	M-2	vgs	OP-70
TK	OP-148	dvgte	I-61	pc	I-38	vgs	I-70
Vb1c1	OP-75	dvgte	OP-61	pc	M-38	vgs	M-70
Vb2c1	OP-74	dvgte	M-61	pc	OP-38	vlr	OP-18
Vb2c2	OP-73	exavl	I-7	pe	I-33	vlr	I-18
Vb2c2star	OP-146	exavl	OP-7	pe	M-33	vlr	M-18
Vb2e1	OP-72	exavl	M-7	pe	OP-33	wavl	I-20
Ve1e	OP-76	exmod	I-5	ps	I-69	wavl	M-20
XCbcex	OP-131	exmod	OP-5	ps	M-69	wavl	OP-20
XIex	OP-85	exmod	M-5	ps	OP-69	xcjc	I-41
XIsub	OP-150	exphi	OP-6	rbc	M-24	xcjc	M-41
XQex	OP-101	exphi	M-6	rbc	I-24	xcjc	OP-41
XQtex	OP-99	exphi	I-6	rbc	OP-24	xcje	M-34
XgS	OP-154	fT	OP-143	rbv	I-25	xcje	I-34

Spectre Circuit Simulator Reference Component Statements Part I

Xgmuex	OP-113	gS	OP-153	rbv	M-25	xcje	OP-34
XiWepi	OP-145	gSf	OP-155	rbv	OP-25	xext	I-19
ab	M-52	gm	OP-136	rcc	I-26	xext	M-19
ab	I-52	gmu	OP-139	rcc	M-26	xext	OP-19
ab	OP-52	gmuex	OP-112	rcc	OP-26	xibi	M-15
ac	OP-55	gmux	OP-109	rcv	I-27	xibi	I-15
ac	M-55	gmuy	OP-110	rcv	M-27	xibi	OP-15
ac	I-55	gmuz	OP-111	rcv	OP-27	xp	M-39
ae	OP-51	gout	OP-138	re	I-23	xp	OP-39
ae	M-51	gpix	OP-106	re	M-23	xp	I-39
ae	I-51	gpiy	OP-107	re	OP-23	xrec	M-49
aepi	I-53	gpiz	OP-108	region	I-72	xrec	OP-49
aepi	OP-53	grbvz	OP-117	scrcv	I-28	xrec	I-49
aepi	M-53	grbvy	OP-118	scrcv	OP-28		
aex	M-54	grbvz	OP-119	scrcv	M-28		
aex	OP-54	grcvy	OP-114	sfh	I-22		

Vertical NPN/PNP Transistor (bjt504t)

Description

The bjt504T model provides a detailed description of a vertical integrated NPN and PNP transistor. It is described in the Philips Bipolar Modelbook (Dec.95) as TN/TNS and TP/TPS level 504.

The NPN is also described in Nat.Lab. Unclassified Report Nr. 006/94 as Mextram Bipolar Transistor Model. Information on how to obtain this document can be found on Source Link by searching for Philips.

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In addition to the model description a `level` parameter is added. Via the `level` parameter the user can switch between Philips Bipolar Modelbook (Dec.95) and Philips Bipolar Modelbook (Dec.94).

Spectre Circuit Simulator Reference

Component Statements Part I

The `imax` parameter is used to aid convergence and to prevent numerical overflow. The junction characteristics of the transistor are accurately modeled for currents up to `imax`. For currents above `imax`, the junction is modeled as a linear resistor and a warning is printed.

The descriptions of the operating point derivatives are given for the NPN type. For the PNP type the terminal voltage in the descriptions has to be exchanged. E.g.:

NPN: $g_x = dI_n/dV_{b2e1}$

PNP: $g_x = dI_n/dV_{e1b2}$

This device is supported within altergroups.

This device is dynamically loaded from the shared object `/vobs/spectre_dev/tools.sun4v/spectre/lib/cmi/3.0.doc/libphilips_sh.so`

Instance Definition

```
Name c b e [s] [dt] ModelName parameter=value ...
```

Options:

[s] When s is not specified, it is bound to ground;

dt dt must be specified, it is NOT optional.

Instance Parameters

1	<code>level=X</code>	level.
2	<code>mult=X</code>	mult.
3	<code>tref=X</code>	tref.
4	<code>dta=X</code>	dta.
5	<code>exmod=X</code>	exmod.
6	<code>exphi=X</code>	exphi.
7	<code>exavl=X</code>	exavl.
8	<code>is=X</code>	is.

Spectre Circuit Simulator Reference

Component Statements Part I

9	ik=X	ik.
10	ver=X	ver.
11	vef=X	vef.
12	bf=X	bf.
13	ibf=X	ibf.
14	mlf=X	mlf.
15	xibi=X	xibi.
16	bri=X	bri.
17	ibr=X	ibr.
18	vlr=X	vlr.
19	xext=X	xext.
20	wavl=X	wavl.
21	vavl=X	vavl.
22	sfh=X	sfh.
23	re=X	re.
24	rbc=X	rbc.
25	rbv=X	rbv.
26	rcc=X	rcc.
27	rcv=X	rcv.
28	scrcv=X	scrcv.
29	ihc=X	ihc.
30	axi=X	axi.

Spectre Circuit Simulator Reference

Component Statements Part I

31	cje=X	cje.
32	vde=X	vde.
33	pe=X	pe.
34	xcje=X	xcje.
35	cbeo=X	cbeo.
36	cjc=X	cjc.
37	vdc=X	vdc.
38	pc=X	pc.
39	xp=X	xp.
40	mc=X	mc.
41	xcjc=X	xcjc.
42	cbco=X	cbco.
43	mtau=X	mtau.
44	taue=X	taue.
45	taub=X	taub.
46	tepi=X	tepi.
47	taur=X	taur.
48	deg=X	deg.
49	xrec=X	xrec.
50	aqbo=X	aqbo.
51	ae=X	ae.
52	ab=X	ab.

Spectre Circuit Simulator Reference

Component Statements Part I

53	aepi=X	aepi.
54	aex=X	aex.
55	ac=X	ac.
56	dvgbf=X	dvgbf.
57	dvgbr=X	dvgbr.
58	vgb=X	vgb.
59	vgc=X	vgc.
60	vgj=X	vgj.
61	dvgte=X	dvgte.
62	af=X	af.
63	kf=X	kf.
64	kfn=X	kfn.
65	iss=X	iss.
66	iks=X	iks.
67	cjs=X	cjs.
68	vds=X	vds.
69	ps=X	ps.
70	vgs=X	vgs.
71	as=X	as.
72	rth=X	rth.
73	cth=X	cth.

Spectre Circuit Simulator Reference

Component Statements Part I

74	<code>region=fwd</code>	Estimated DC operating region, used as a convergence aid. Possible values are <code>off</code> , <code>fwd</code> , <code>rev</code> , or <code>sat</code> .
75	<code>m=1</code>	Multiplication factor.
76	<code>area=1</code>	Alias of <code>mult</code> .

Model Definition

```
model modelName bjt504t parameter=value ...
```

Model Parameters

1	<code>level=NULL</code>	<code>level</code> .
2	<code>mult=NULL</code>	<code>mult</code> .
3	<code>tref=NULL</code>	<code>tref</code> .
4	<code>dta=NULL</code>	<code>dta</code> .
5	<code>exmod=NULL</code>	<code>exmod</code> .
6	<code>exphi=NULL</code>	<code>exphi</code> .
7	<code>exavl=NULL</code>	<code>exavl</code> .
8	<code>is=NULL</code>	<code>is</code> .
9	<code>ik=NULL</code>	<code>ik</code> .
10	<code>ver=NULL</code>	<code>ver</code> .
11	<code>vef=NULL</code>	<code>vef</code> .
12	<code>bf=NULL</code>	<code>bf</code> .
13	<code>ibf=NULL</code>	<code>ibf</code> .
14	<code>mlf=NULL</code>	<code>mlf</code> .
15	<code>xibi=NULL</code>	<code>xibi</code> .

Spectre Circuit Simulator Reference

Component Statements Part I

16	bri=NULL	bri.
17	ibr=NULL	ibr.
18	vlr=NULL	vlr.
19	xext=NULL	xext.
20	wavl=NULL	wavl.
21	vavl=NULL	vavl.
22	sfh=NULL	sfh.
23	re=NULL	re.
24	rbc=NULL	rbc.
25	rbv=NULL	rbv.
26	rcc=NULL	rcc.
27	rcv=NULL	rcv.
28	scrcv=NULL	scrcv.
29	ihc=NULL	ihc.
30	axi=NULL	axi.
31	cje=NULL	cje.
32	vde=NULL	vde.
33	pe=NULL	pe.
34	xcje=NULL	xcje.
35	cbeo=NULL	cbeo.
36	cjc=NULL	cjc.
37	vdc=NULL	vdc.

Spectre Circuit Simulator Reference

Component Statements Part I

38	pc=NULL	pc.
39	xp=NULL	xp.
40	mc=NULL	mc.
41	xcjc=NULL	xcjc.
42	cbco=NULL	cbco.
43	mtau=NULL	mtau.
44	taue=NULL	taue.
45	taub=NULL	taub.
46	tepi=NULL	tepi.
47	taur=NULL	taur.
48	deg=NULL	deg.
49	xrec=NULL	xrec.
50	aqbo=NULL	aqbo.
51	ae=NULL	ae.
52	ab=NULL	ab.
53	aepi=NULL	aepi.
54	aex=NULL	aex.
55	ac=NULL	ac.
56	dvgbf=NULL	dvgbf.
57	dvgbr=NULL	dvgbr.
58	vgb=NULL	vgb.
59	vgc=NULL	vgc.

Spectre Circuit Simulator Reference

Component Statements Part I

60	vgj=NULL	vgj.
61	dvgte=NULL	dvgte.
62	af=NULL	af.
63	kf=NULL	kf.
64	kfn=NULL	kfn.
65	iss=NULL	iss.
66	iks=NULL	iks.
67	cjs=NULL	cjs.
68	vds=NULL	vds.
69	ps=NULL	ps.
70	vgs=NULL	vgs.
71	as=NULL	as.
72	rth=NULL	rth.
73	cth=NULL	cth.
74	type=npn	Transistor type. Possible values are npn or pnp.
75	imax=1.0 A	Explosion current.
76	tnom=25 C	alias of tref. Default set by option tnom.
77	tr (C)	Alias of tref. Default set by option tnom.

Operating-Point Parameters

1	level=X	level.
2	mult=X	mult.

Spectre Circuit Simulator Reference

Component Statements Part I

3	tref=X	tref.
4	dta=X	dta.
5	exmod=X	exmod.
6	exphi=X	exphi.
7	exavl=X	exavl.
8	is=X	is.
9	ik=X	ik.
10	ver=X	ver.
11	vef=X	vef.
12	bf=X	bf.
13	ibf=X	ibf.
14	mlf=X	mlf.
15	xibi=X	xibi.
16	bri=X	bri.
17	ibr=X	ibr.
18	vlr=X	vlr.
19	xext=X	xext.
20	wavl=X	wavl.
21	vavl=X	vavl.
22	sfh=X	sfh.
23	re=X	re.
24	rbc=X	rbc.

Spectre Circuit Simulator Reference

Component Statements Part I

25	rbv=X	rbv.
26	rcc=X	rcc.
27	rcv=X	rcv.
28	scrcv=X	scrcv.
29	ihc=X	ihc.
30	axi=X	axi.
31	cje=X	cje.
32	vde=X	vde.
33	pe=X	pe.
34	xcje=X	xcje.
35	cbeo=X	cbeo.
36	cjc=X	cjc.
37	vdc=X	vdc.
38	pc=X	pc.
39	xp=X	xp.
40	mc=X	mc.
41	xcjc=X	xcjc.
42	cbco=X	cbco.
43	mtau=X	mtau.
44	taue=X	taue.
45	taub=X	taub.
46	tepi=X	tepi.

Spectre Circuit Simulator Reference

Component Statements Part I

47	taur=X	taur.
48	deg=X	deg.
49	xrec=X	xrec.
50	aqbo=X	aqbo.
51	ae=X	ae.
52	ab=X	ab.
53	aepi=X	aepi.
54	aex=X	aex.
55	ac=X	ac.
56	dvgbf=X	dvgbf.
57	dvgbr=X	dvgbr.
58	vgb=X	vgb.
59	vgc=X	vgc.
60	vgj=X	vgj.
61	dvgte=X	dvgte.
62	af=X	af.
63	kf=X	kf.
64	kfn=X	kfn.
65	iss=X	iss.
66	iks=X	iks.
67	cjs=X	cjs.
68	vds=X	vds.

Spectre Circuit Simulator Reference

Component Statements Part I

69	ps=X	ps.
70	vgs=X	vgs.
71	as=X	as.
72	rth=X	rth.
73	cth=X	cth.
74	Vb2e1	Vb2e1.
75	Vb2c2	Vb2c2.
76	Vb2c1	Vb2c1.
77	Vb1c1	Vb1c1.
78	Ve1e	Ve1e.
79	In	In.
80	Ic1c2	Ic1c2.
81	Ib1b2	Ib1b2.
82	Ib1	Ib1.
83	SIb1	SIb1.
84	Ib2	Ib2.
85	Ib3	Ib3.
86	Iex	Iex.
87	XIex	XIex.
88	Iavl	Iavl.
89	IRE	IRE.
90	IRBC	IRBC.

Spectre Circuit Simulator Reference

Component Statements Part I

91	IRCC	IRCC.
92	Qe	Qe.
93	Qte	Qte.
94	SQte	SQte.
95	Qbe	Qbe.
96	Qbc	Qbc.
97	Qtc	Qtc.
98	Qepi	Qepi.
99	Qb1b2	Qb1b2.
100	Qtex	Qtex.
101	XQtex	XQtex.
102	Qex	Qex.
103	XQex	XQex.
104	gx	gx.
105	gy	gy.
106	gz	gz.
107	Sgpi	Sgpi.
108	gpix	gpix.
109	gpiy	gpiy.
110	gpiz	gpiz.
111	gmux	gmux.
112	gmuy	gmuy.

Spectre Circuit Simulator Reference

Component Statements Part I

113 gmuz	gmuz.
114 gmueX	gmueX.
115 XgmueX	XgmueX.
116 grcvy	grcvy.
117 grcvz	grcvz.
118 Rbv	Rbv.
119 grbvX	grbvX.
120 grbvy	grbvy.
121 grbvz	grbvz.
122 RE	RE.
123 RBC	RBC.
124 RCC	RCC.
125 SCbe	SCbe.
126 Cbex	Cbex.
127 Cbey	Cbey.
128 Cbez	Cbez.
129 Cbcx	Cbcx.
130 Cbcy	Cbcy.
131 Cbcz	Cbcz.
132 Cbcex	Cbcex.
133 XCbcex	XCbcex.
134 Cb1b2	Cb1b2.

Spectre Circuit Simulator Reference

Component Statements Part I

135 Cb1b2x	Cb1b2x.
136 Cb1b2y	Cb1b2y.
137 Cb1b2z	Cb1b2z.
138 gm	gm.
139 beta	beta.
140 gout	gout.
141 gmU	gmU.
142 RB	RB.
143 Cbe	Cbe.
144 Cbc	Cbc.
145 fT	fT.
146 Iqs	Iqs.
147 XiWepi	XiWepi.
148 Vb2c2star	Vb2c2star.
149 Pdiss	Pdiss.
150 TK	TK.
151 Isub	Isub.
152 XIsub	XIsub.
153 Isf	Isf.
154 Qts	Qts.
155 gS	gS.
156 XgS	XgS.

Spectre Circuit Simulator Reference Component Statements Part I

157 gSf

gSf.

158 Cts

Cts.

Parameter Index

In the following index, I refers to instance parameters, M refers to the model parameters section, O refers to the output parameters section, and OP refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of M-35 means the 35th model parameter.

Cb1b2	OP-134	af	M-62	grcvy	OP-116	scrcv	OP-28
Cb1b2x	OP-135	af	OP-62	grcvz	OP-117	scrcv	M-28
Cb1b2y	OP-136	af	I-62	gx	OP-104	sfh	OP-22
Cb1b2z	OP-137	aqbo	M-50	gy	OP-105	sfh	I-22
Cbc	OP-144	aqbo	OP-50	gz	OP-106	sfh	M-22
Cbcex	OP-132	aqbo	I-50	ibf	I-13	taub	OP-45
Cbcx	OP-129	area	I-76	ibf	M-13	taub	I-45
Cbcy	OP-130	as	OP-71	ibf	OP-13	taub	M-45
Cbcz	OP-131	as	M-71	ibr	OP-17	taue	I-44
Cbe	OP-143	as	I-71	ibr	I-17	taue	M-44
Cbex	OP-126	axi	I-30	ibr	M-17	taue	OP-44
Cbey	OP-127	axi	OP-30	ihc	M-29	taur	M-47
Cbez	OP-128	axi	M-30	ihc	OP-29	taur	I-47
Cts	OP-158	beta	OP-139	ihc	I-29	taur	OP-47
IRBC	OP-90	bf	OP-12	ik	OP-9	tepi	I-46
IRCC	OP-91	bf	M-12	ik	M-9	tepi	M-46
IRE	OP-89	bf	I-12	ik	I-9	tepi	OP-46
Iavl	OP-88	bri	M-16	iks	OP-66	tnom	M-76
Ib1	OP-82	bri	OP-16	iks	M-66	tr	M-77
Ib1b2	OP-81	bri	I-16	iks	I-66	tref	I-3
Ib2	OP-84	cbco	I-42	imax	M-75	tref	M-3
Ib3	OP-85	cbco	OP-42	is	OP-8	tref	OP-3

Spectre Circuit Simulator Reference Component Statements Part I

Ic1c2	OP-80	cbco	M-42	is	I-8	type	M-74
Iex	OP-86	cbeo	I-35	is	M-8	vavl	OP-21
In	OP-79	cbeo	OP-35	iss	I-65	vavl	M-21
Iqs	OP-146	cbeo	M-35	iss	M-65	vavl	I-21
Isf	OP-153	cjc	M-36	iss	OP-65	vdc	OP-37
Sub	OP-151	cjc	OP-36	kf	OP-63	vdc	I-37
Pdiss	OP-149	cjc	I-36	kf	I-63	vdc	M-37
Qb1b2	OP-99	cje	OP-31	kf	M-63	vde	M-32
Qbc	OP-96	cje	M-31	kfn	OP-64	vde	I-32
Qbe	OP-95	cje	I-31	kfn	I-64	vde	OP-32
Qe	OP-92	cjs	M-67	kfn	M-64	vds	M-68
Qepi	OP-98	cjs	I-67	level	M-1	vds	OP-68
Qex	OP-102	cjs	OP-67	level	OP-1	vds	I-68
Qtc	OP-97	cth	M-73	level	I-1	vef	I-11
Qte	OP-93	cth	OP-73	m	I-75	vef	OP-11
Qtex	OP-100	cth	I-73	mc	M-40	vef	M-11
Qts	OP-154	deg	I-48	mc	OP-40	ver	OP-10
RB	OP-142	deg	M-48	mc	I-40	ver	I-10
RBC	OP-123	deg	OP-48	mlf	I-14	ver	M-10
RCC	OP-124	dta	I-4	mlf	OP-14	vgb	OP-58
RE	OP-122	dta	OP-4	mlf	M-14	vgb	I-58
Rbv	OP-118	dta	M-4	mtau	I-43	vgb	M-58
SCbe	OP-125	dvgbf	M-56	mtau	OP-43	vgc	I-59
SIb1	OP-83	dvgbf	I-56	mtau	M-43	vgc	OP-59
SQte	OP-94	dvgbf	OP-56	mult	M-2	vgc	M-59
Sgpi	OP-107	dvgbr	OP-57	mult	OP-2	vgj	I-60
TK	OP-150	dvgbr	I-57	mult	I-2	vgj	OP-60
Vb1c1	OP-77	dvgbr	M-57	pc	I-38	vgj	M-60
Vb2c1	OP-76	dvgte	OP-61	pc	M-38	vgs	I-70
Vb2c2	OP-75	dvgte	I-61	pc	OP-38	vgs	M-70
Vb2c2star	OP-148	dvgte	M-61	pe	OP-33	vgs	OP-70
Vb2e1	OP-74	exavl	I-7	pe	M-33	vlr	I-18

Spectre Circuit Simulator Reference Component Statements Part I

Ve1e	OP-78	exavl	M-7	pe	I-33	v1r	OP-18
XCbcex	OP-133	exavl	OP-7	ps	M-69	v1r	M-18
XIex	OP-87	exmod	M-5	ps	I-69	wavl	I-20
XIsub	OP-152	exmod	I-5	ps	OP-69	wavl	M-20
XQex	OP-103	exmod	OP-5	rbc	OP-24	wavl	OP-20
XQtex	OP-101	exphi	M-6	rbc	M-24	xcjc	M-41
XgS	OP-156	exphi	I-6	rbc	I-24	xcjc	I-41
Xgmux	OP-115	exphi	OP-6	rbv	I-25	xcjc	OP-41
XiWepi	OP-147	fT	OP-145	rbv	M-25	xcje	M-34
ab	OP-52	gS	OP-155	rbv	OP-25	xcje	I-34
ab	I-52	gSf	OP-157	rcc	OP-26	xcje	OP-34
ab	M-52	gm	OP-138	rcc	M-26	xext	OP-19
ac	M-55	gmu	OP-141	rcc	I-26	xext	M-19
ac	I-55	gmux	OP-114	rcv	I-27	xext	I-19
ac	OP-55	gmux	OP-111	rcv	M-27	xibi	M-15
ae	I-51	gmuy	OP-112	rcv	OP-27	xibi	OP-15
ae	OP-51	gmuz	OP-113	re	OP-23	xibi	I-15
ae	M-51	gout	OP-140	re	I-23	xp	M-39
aepi	M-53	gpix	OP-108	re	M-23	xp	OP-39
aepi	I-53	gpiy	OP-109	region	I-74	xp	I-39
aepi	OP-53	gpiz	OP-110	rth	OP-72	xrec	I-49
aex	M-54	grbvz	OP-119	rth	M-72	xrec	M-49
aex	OP-54	grbvy	OP-120	rth	I-72	xrec	OP-49
aex	I-54	grbvz	OP-121	scrcv	I-28		

BSIM1 Field Effect Transistor (bsim1)

Description

BSIM1 is a semiempirical MOSFET model developed at the University of California, Berkeley. All the model parameters are extracted directly from physical devices. Three charge models are available. In SPICE mode, you can refer to BSIM1 as MOS level 4 or BSIM level 1. BSIM1 transistors require that you use a model statement.

Spectre Circuit Simulator Reference

Component Statements Part I

This device is supported within altergroups.

Sample Instance Statement

```
m1 (1 2 0 0) nchmod l=5u w=10u as=40u ad=40u pd=28u ps=28u m=1
```

Sample Model Statement

```
model nchmod bsim1 vfb0=-0.5 lvfb=0.5 wvfb=0.3 phi0=0.8 eta0=0.056 k1=0.5 muz=454  
eg=0.99 gap1=5.5e-04 trs=1e-3 trd=1e-3 xpart=0.5 rs=10 rd=10
```

Instance Definition

```
Name d g s b ModelName parameter=value ...
```

Instance Parameters

1	w (m)	Channel width.
2	l (m)	Channel length.
3	as (m ²)	Area of source diffusion.
4	ad (m ²)	Area of drain diffusion.
5	ps (m)	Perimeter of source diffusion.
6	pd (m)	Perimeter of drain diffusion.
7	nrd (m/m)	Number of squares of drain diffusion.
8	nrs (m/m)	Number of squares of source diffusion.
9	ld (m)	Drain diffusion length.
10	ls (m)	Source diffusion length.
11	m=1	Multiplicity factor (number of MOSFETs in parallel).
12	region=triode	Estimated operating region. Spectre outputs number (0-4) in a rawfile. Possible values are off, triode, sat, subth, or breakdown.

Spectre Circuit Simulator Reference

Component Statements Part I

- 13 `trise (C)` Temperature rise from ambient.
- 14 `degradation=no` Hot-electron degradation flag.
Possible values are `no` or `yes`.

Model Definition

```
model modelName bsim1 parameter=value ...
```

Model Parameters

Device type parameters

- 1 `type=n` Transistor type.
Possible values are `n` or `p`.

Threshold voltage parameters

- 2 `vfb0=-0.8 V` Flat-band voltage.
- 3 `lvfb=0 V μm` Length dependence of `vfb`.
- 4 `wvfb=0 V μm` Width dependence of `vfb`.
- 5 `pvfb=0 V μm` Width-length dependence of `vfb`.
- 6 `phi0=0.75 V` Surface potential.
- 7 `lphi=0 V μm` Length dependence of `phi`.
- 8 `wphi=0 V μm` Width dependence of `phi`.
- 9 `pphi=0 V μm` Width-length dependence of `phi`.
- 10 `k1=0.7 √V` Body-effect coefficient.
- 11 `lk1=0 √V μm` Length dependence of `k1`.
- 12 `wk1=0 √V μm` Width dependence of `k1`.
- 13 `pk1=0 √V μm` Width-length dependence of `k1`.

Spectre Circuit Simulator Reference

Component Statements Part I

14	$k2=0$	Charge-sharing parameter.
15	$lk2=0 \mu\text{m}$	Length dependence of $k2$.
16	$wk2=0 \mu\text{m}$	Width dependence of $k2$.
17	$pk2=0 \mu\text{m}$	Width-length dependence of $k2$.
18	$\text{eta}0=0$	Drain-induced barrier-lowering coefficient.
19	$l\text{eta}=0 \mu\text{m}$	Length dependence of eta .
20	$w\text{eta}=0 \mu\text{m}$	Width dependence of eta .
21	$p\text{eta}=0 \mu\text{m}$	Width-length dependence of eta .
22	$x2e=0 1/V$	Body-bias dependence of eta .
23	$lx2e=0 \mu\text{m}/V$	Length dependence of $x2e$.
24	$wx2e=0 \mu\text{m}/V$	Width dependence of $x2e$.
25	$px2e=0 \mu\text{m}/V$	Width-length dependence of $x2e$.
26	$x3e=0 1/V$	Drain-bias dependence of eta .
27	$lx3e=0 \mu\text{m}/V$	Length dependence of $x3e$.
28	$wx3e=0 \mu\text{m}/V$	Width dependence of $x3e$.
29	$px3e=0 \mu\text{m}/V$	Width-length dependence of $x3e$.

Mobility parameters

30	$\text{muz}=400 \text{ cm}^2/V \text{ s}$	Low-field mobility.
31	$lmuz=0 \text{ cm}^2/V \text{ s}$	Length dependence of muz .
32	$wmuz=0 \text{ cm}^2/V \text{ s}$	Width dependence of muz .
33	$pmuz=0 \text{ cm}^2/V \text{ s}$	Width-length dependence of muz .

Spectre Circuit Simulator Reference Component Statements Part I

- 34 $x_{2mz}=0$ cm^2/V^2 s Body-bias dependence of μ_{uz} .
- 35 $l_{x2mz}=0$ $\text{cm}^2 \mu\text{m}/\text{V}^2$ s Length dependence of x_{2mz} .
- 36 $w_{x2mz}=0$ $\text{cm}^2 \mu\text{m}/\text{V}^2$ s Width dependence of x_{2mz} .
- 37 $p_{x2mz}=0$ $\text{cm}^2 \mu\text{m}/\text{V}^2$ s Width-length dependence of x_{2mz} .
- 38 $\mu_{us}=450$ cm^2/V s Mobility in the saturation region.
- 39 $l_{\mu us}=0$ $\text{cm}^2 \mu\text{m}/\text{V}$ s Length dependence of μ_{us} .
- 40 $w_{\mu us}=0$ $\text{cm}^2 \mu\text{m}/\text{V}$ s Width dependence of μ_{us} .
- 41 $p_{\mu us}=0$ $\text{cm}^2 \mu\text{m}/\text{V}$ s Width-length dependence of μ_{us} .
- 42 $x_{2ms}=0$ cm^2/V^2 s Body-bias dependence of μ_{us} .
- 43 $l_{x2ms}=0$ $\text{cm}^2 \mu\text{m}/\text{V}^2$ s Length dependence of x_{2ms} .
- 44 $w_{x2ms}=0$ $\text{cm}^2 \mu\text{m}/\text{V}^2$ s Width dependence of x_{2ms} .
- 45 $p_{x2ms}=0$ $\text{cm}^2 \mu\text{m}/\text{V}^2$ s Width-length dependence of x_{2ms} .
- 46 $x_{3ms}=0$ cm^2/V^2 s Drain-bias dependence of μ_{us} .
- 47 $l_{x3ms}=0$ $\text{cm}^2 \mu\text{m}/\text{V}^2$ s Length dependence of x_{3ms} .
- 48 $w_{x3ms}=0$ $\text{cm}^2 \mu\text{m}/\text{V}^2$ s Width dependence of x_{3ms} .
- 49 $p_{x3ms}=0$ $\text{cm}^2 \mu\text{m}/\text{V}^2$ s Width-length dependence of x_{3ms} .

Spectre Circuit Simulator Reference

Component Statements Part I

Mobility modulation parameters

50	$u_{00}=0$	$1/V$	Gate voltage dependence of mobility.
51	$l_{u0}=0$	$\mu m/V$	Length dependence of u_0 .
52	$w_{u0}=0$	$\mu m/V$	Width dependence of u_0 .
53	$p_{u0}=0$	$\mu m/V$	Width-length dependence of u_0 .
54	$x_{2u0}=0$	$1/V^2$	Body-bias dependence of u_0 .
55	$l_{x2u0}=0$	$\mu m/V^2$	Length dependence of x_{2u0} .
56	$w_{x2u0}=0$	$\mu m/V^2$	Width dependence of x_{2u0} .
57	$p_{x2u0}=0$	$\mu m/V^2$	Width-length dependence of x_{2u0} .

Velocity saturation parameters

58	$u_{10}=0$	$1/V$	Velocity saturation coefficient.
59	$l_{u1}=0$	$\mu m/V$	Length dependence of u_1 .
60	$w_{u1}=0$	$\mu m/V$	Width dependence of u_1 .
61	$p_{u1}=0$	$\mu m/V$	Width-length dependence of u_1 .
62	$x_{2u1}=0$	$1/V^2$	Body-bias dependence of u_1 .
63	$l_{x2u1}=0$	$\mu m/V^2$	Length dependence of x_{2u1} .
64	$w_{x2u1}=0$	$\mu m/V^2$	Width dependence of x_{2u1} .
65	$p_{x2u1}=0$	$\mu m/V^2$	Width-length dependence of x_{2u1} .
66	$x_{3u1}=0$	$1/V^2$	Drain-bias dependence of u_1 .
67	$l_{x3u1}=0$	$\mu m/V^2$	Length dependence of x_{3u1} .
68	$w_{x3u1}=0$	$\mu m/V^2$	Width dependence of x_{3u1} .

Spectre Circuit Simulator Reference

Component Statements Part I

69 $px3u1=0$ $\mu\text{m}/\text{V}^2$ Width-length dependence of $x3u1$.

Subthreshold parameters

70 $n0=0$ Subthreshold swing parameter.

71 $ln0=0$ μm Length dependence of subthreshold swing parameter.

72 $wn0=0$ μm Width dependence of subthreshold swing parameter.

73 $pn0=0$ μm Width-length dependence of subthreshold swing parameter.

74 $nb=0$ $\sqrt{\text{V}}$ Body-bias dependence of $n0$.

75 $lnb=0$ $\sqrt{\text{V}}$ μm Length dependence of nb .

76 $wnb=0$ $\sqrt{\text{V}}$ μm Width dependence of nb .

77 $pnb=0$ $\sqrt{\text{V}}$ μm Width-length dependence of nb .

78 $nd=0$ $1/\text{V}$ Drain-bias dependence of $n0$.

79 $lnd=0$ $\mu\text{m}/\text{V}$ Length dependence of nd .

80 $wnd=0$ $\mu\text{m}/\text{V}$ Width dependence of nd .

81 $pnd=0$ $\mu\text{m}/\text{V}$ Width-length dependence of nd .

82 $subthmod=2$ Subthreshold model selector.

Impact ionization parameters

83 $ai0=0$ $1/\text{V}$ Hot-electron effect on R_{out} parameter.

84 $lai0=0$ $\mu\text{m}/\text{V}$ Length dependence of $ai0$.

85 $wai0=0$ $\mu\text{m}/\text{V}$ Width dependence of $ai0$.

86 $pai0=0$ $\mu\text{m}/\text{V}$ Width-length dependence of $ai0$.

87 $bi0=0$ V Hot-electron effect on R_{out} exponent.

Spectre Circuit Simulator Reference

Component Statements Part I

- 88 $l_{bi0}=0$ V μm Length dependence of b_{i0} .
- 89 $w_{bi0}=0$ V μm Width dependence of b_{i0} .
- 90 $p_{bi0}=0$ V μm Width-length dependence of b_{i0} .

Length and width modulation parameters

- 91 $d_{l0}=0$ μm Lateral diffusion.
- 92 $d_{w0}=0$ μm Field oxide encroachment.
- 93 $l_{ref}=\infty$ m Reference channel length.
- 94 $w_{ref}=\infty$ m Reference channel width.
- 95 $x_w=0$ m Width variation due to masking and etching.
- 96 $x_l=0$ m Length variation due to masking and etching.

Temperature effects parameters

- 97 $temp$ (C) Parameters measurement temperature. Default set by options.
- 98 $trise=0$ C Temperature rise from ambient.
- 99 $tempmod=432$ Temperature model selector.
- 100 $version=432$ Version selector.
- 101 $uto=0$ C Mobility temperature offset.
- 102 $ute=-1.5$ Mobility temperature exponent.
- 103 $tlev=0$ DC temperature selector.
- 104 $tlevc=0$ AC temperature selector.
- 105 $eg=1.12452$ V Energy band gap.

Spectre Circuit Simulator Reference

Component Statements Part I

106	$\text{gap1}=7.02\text{e-}4 \text{ V/C}^2$	Band gap temperature coefficient.
107	$\text{gap2}=1108 \text{ K}$	Band gap temperature offset.
108	$\text{trs}=0 \text{ 1/C}$	Temperature coefficient for source resistance.
109	$\text{trd}=0 \text{ 1/C}$	Temperature coefficient for drain resistance.
110	$\text{xti}=3$	Saturation current temperature exponent.

Overlap capacitance parameters

111	$\text{cgso}=0 \text{ F/m}$	Gate-source overlap capacitance.
112	$\text{cgdo}=0 \text{ F/m}$	Gate-drain overlap capacitance.
113	$\text{cgbo}=0 \text{ F/m}$	Gate-bulk overlap capacitance.
114	$\text{meto}=0 \text{ m}$	Metal overlap in fringing field.

Charge model selection parameters

115	$\text{capmod}=\text{bsim}$	Intrinsic charge model. Possible values are <i>none</i> , <i>meyer</i> , <i>yang</i> , or <i>bsim</i> .
116	$\text{xpart}=1$	Drain/source channel charge partition in saturation for BSIM charge model, use 0.0 for 40/60, 0.5 for 50/50, or 1.0 for 0/100.
117	$\text{xqc}=0$	Drain/source channel charge partition in saturation for charge models, e.g. use 0.4 for 40/60, 0.5 for 50/50, or 0 for 0/100.

Parasitic resistance parameters

118	$\text{rs}=0 \text{ }\Omega$	Source resistance.
119	$\text{rd}=0 \text{ }\Omega$	Drain resistance.
120	$\text{rsh}=0 \text{ }\Omega/\text{sqr}$	Source/drain diffusion sheet resistance.

Spectre Circuit Simulator Reference

Component Statements Part I

121	$r_{sc}=0 \ \Omega$	Source contact resistance.
122	$r_{dc}=0 \ \Omega$	Drain contact resistance.
123	$r_{ss}=0 \ \Omega \ m$	Scalable source resistance.
124	$r_{dd}=0 \ \Omega \ m$	Scalable drain resistance.
125	$minr=0.1 \ \Omega$	Minimum source/drain resistance.
126	$hdif=0 \ m$	Length of heavily doped diffusion.
127	$ldif=0 \ m$	Lateral diffusion beyond the gate.
128	$lgcs=0 \ m$	Gate-to-contact length of source side.
129	$lgcd=0 \ m$	Gate-to-contact length of drain side.
130	$sc=\infty \ m$	Spacing between contacts.

Junction diode parameters

131	$j_s \ (A/m^2)$	Bulk junction reverse saturation current density.
132	$i_s=1e-14 \ A$	Bulk junction reverse saturation current.
133	$n=1$	Junction emission coefficient.
134	$imelt='imaxA'$	Explosion current, diode is linearized beyond this current to aid convergence.
135	$jmelt='jmaxA/m'^2$	Explosion current density, diode is linearized beyond this current to aid convergence.
136	$dskip=yes$	Use simple piece-wise linear model for diode currents below $0.1*iabstol$. Possible values are <code>no</code> or <code>yes</code> .

Spectre Circuit Simulator Reference

Component Statements Part I

Junction capacitance model parameters

137	$c_{bs}=0$ F	Bulk-source zero-bias junction capacitance.
138	$c_{bd}=0$ F	Bulk-drain zero-bias junction capacitance.
139	$c_{j}=0$ F/m ²	Zero-bias junction bottom capacitance density.
140	$m_j=1/2$	Bulk junction bottom grading coefficient.
141	$p_b=0.8$ V	Bulk junction potential.
142	$f_c=0.5$	Forward-bias depletion capacitance threshold.
143	$c_{jsw}=0$ F/m	Zero-bias junction sidewall capacitance density.
144	$m_{jsw}=1/3$	Bulk junction sidewall grading coefficient.
145	$p_{bsw}=0.8$ V	Side-wall junction potential.
146	$f_{csw}=0.5$	Side-wall forward-bias depletion capacitance threshold.

Operating region warning control parameters

147	$alarm=none$	Forbidden operating region. Possible values are <i>none</i> , <i>off</i> , <i>triode</i> , <i>sat</i> , <i>subth</i> , or <i>rev</i> .
148	$i_{max}=1$ A	Maximum current, currents above this limit generate a warning.
149	$j_{max}=1e8$ A/m ²	Maximum current density, currents above this limit generate a warning.
150	$b_{vj}=\infty$ V	Junction reverse breakdown voltage.
151	$v_{box}=1e9$ t_{ox} V	Oxide breakdown voltage.

Process and power supply parameters

152	$t_{ox}=4e-8$ m	Gate oxide thickness.
153	$v_{dd}=5$ V	Drain voltage at which parameters are extracted.

Spectre Circuit Simulator Reference

Component Statements Part I

Default device parameters

154	$w=3e-6$ m	Channel width.
155	$l=3e-6$ m	Channel length.
156	$as=0$ m ²	Area of source diffusion.
157	$ad=0$ m ²	Area of drain diffusion.
158	$ps=0$ m	Perimeter of source diffusion.
159	$pd=0$ m	Perimeter of drain diffusion.
160	$nrd=0$ m/m	Number of squares of drain diffusion.
161	$nrs=0$ m/m	Number of squares of source diffusion.
162	$l_{dd}=0$ m	Drain diffusion length.
163	$l_{ds}=0$ m	Source diffusion length.

Noise model parameters

164	$noisemod=1$	Noise model selector.
165	$kf=0$	Flicker (1/f) noise coefficient.
166	$af=1$	Flicker (1/f) noise exponent.
167	$ef=1$	Flicker (1/f) noise frequency exponent.
168	$w_{noi}=1e-5$ m	Channel width at which noise parameters were extracted.

Auto Model Selector parameters

169	$w_{max}=1.0$ m	Maximum channel width for which the model is valid.
170	$w_{min}=0.0$ m	Minimum channel width for which the model is valid.
171	$l_{max}=1.0$ m	Maximum channel length for which the model is valid.

Spectre Circuit Simulator Reference

Component Statements Part I

172 `lmin=0.0 m` Minimum channel length for which the model is valid.

Degradation parameters

173 `degramod=spectre` Degradation model selector.
Possible values are `spectre` or `bert`.

174 `degradation=no` Hot-electron degradation flag.
Possible values are `no` or `yes`.

175 `dvthc=1 V` Degradation coefficient for threshold voltage.

176 `dvthe=1` Degradation exponent for threshold voltage.

177 `duoc=1 S` Degradation coefficient for transconductance.

178 `duoe=1` Degradation exponent for transconductance.

179 `crivth=0.1 V` Maximum allowable threshold voltage shift.

180 `criuo=10%` Maximum allowable normalized mobility change.

181 `crigm=10%` Maximum allowable normalized transconductance change.

182 `criids=10%` Maximum allowable normalized drain current change.

183 `wnom=5e-6 m` Nominal device width in degradation calculation.

184 `lnom=1e-6 m` Nominal device length in degradation calculation.

185 `vbsn=0 V` Substrate voltage in degradation calculation.

186 `vdsni=0.1 V` Drain voltage in I_{ds} degradation calculation.

187 `vgsni=5 V` Gate voltage in I_{ds} degradation calculation.

188 `vdsng=0.1 V` Drain voltage in G_m degradation calculation.

189 `vgsng=5 V` Gate voltage in G_m degradation calculation.

Spectre Circuit Simulator Reference

Component Statements Part I

Spectre stress parameters

190	$esat=1.1e7$	V/m	Critical field in v_{dsat} calculation.
191	$esatg=2.5e6$	1/m	Gate voltage dependence of $esat$.
192	$vpg=-0.25$		Gate voltage modifier.
193	$vpb=-0.13$		Gate voltage modifier.
194	$subc1=2.24e-5$		Substrate current coefficient.
195	$subc2=-0.1e-5$	1/V	Substrate current coefficient.
196	$sube=6.4$		Substrate current exponent.
197	$strc=1$		Stress coefficient.
198	$stre=1$		Stress exponent.

BERT stress parameters

199	$h0=1$		Aging coefficient.
200	$hgd=0$	1/V	Bias dependence of $h0$.
201	$m0=1$		Aging exponent.
202	$mgd=0$	1/V	Bias dependence of $m0$.
203	$ecrit0=1.1e5$	V/cm	Critical electric field.
204	$lecrit0=0$	μm V/cm	Length dependence of $ecrit0$.
205	$wecrit0=0$	μm V/cm	Width dependence of $ecrit0$.
206	$ecritg=0$	1/cm	Gate voltage dependence of $ecrit0$.
207	$lecritg=0$	$\mu\text{m}/\text{cm}$	Length dependence of $ecritg$.
208	$wecritg=0$	$\mu\text{m}/\text{cm}$	Width dependence of $ecritg$.

Spectre Circuit Simulator Reference

Component Statements Part I

209	$ecritb=0$	$1/cm$	Substrate voltage dependence of $ecrit0$.
210	$lecritb=0$	$\mu m/cm$	Length dependence of $ecritb$.
211	$wecritb=0$	$\mu m/cm$	Width dependence of $ecritb$.
212	$lc0=1$		Substrate current coefficient.
213	$llc0=0$	μm	Length dependence of $lc0$.
214	$wlc0=0$	μm	Width dependence of $lc0$.
215	$lc1=1$		Substrate current coefficient.
216	$llc1=0$	μm	Length dependence of $lc1$.
217	$wlc1=0$	μm	Width dependence of $lc1$.
218	$lc2=1$		Substrate current coefficient.
219	$llc2=0$	μm	Length dependence of $lc2$.
220	$wlc2=0$	μm	Width dependence of $lc2$.
221	$lc3=1$		Substrate current coefficient.
222	$llc3=0$	μm	Length dependence of $lc3$.
223	$wlc3=0$	μm	Width dependence of $lc3$.
224	$lc4=1$		Substrate current coefficient.
225	$llc4=0$	μm	Length dependence of $lc4$.
226	$wlc4=0$	μm	Width dependence of $lc4$.
227	$lc5=1$		Substrate current coefficient.
228	$llc5=0$	μm	Length dependence of $lc5$.
229	$wlc5=0$	μm	Width dependence of $lc5$.
230	$lc6=1$		Substrate current coefficient.

Spectre Circuit Simulator Reference

Component Statements Part I

231	$llc6=0 \mu\text{m}$	Length dependence of $lc6$.
232	$wlc6=0 \mu\text{m}$	Width dependence of $lc6$.
233	$lc7=1$	Substrate current coefficient.
234	$llc7=0 \mu\text{m}$	Length dependence of $lc7$.
235	$wlc7=0 \mu\text{m}$	Width dependence of $lc7$.

I_{max} and I_{melt}

The i_{max} parameter aids convergence and prevents numerical overflow. The junction characteristics of the device are accurately modeled for current up to i_{max} . If i_{max} is exceeded during iterations, the linear model is substituted until the current drops below i_{max} or until convergence is achieved. If convergence is achieved with the current exceeding i_{max} , the results are inaccurate, and Spectre prints a warning.

A separate model parameter, i_{melt} , is used as a limit warning for the junction current. This parameter can be set to the maximum current rating of the device. When any component of the junction current exceeds i_{melt} , note that base and collector currents are composed of many exponential terms, Spectre issues a warning and the results become inaccurate. The junction current is linearized above the value of i_{melt} to prevent arithmetic exception, with the exponential term replaced by a linear equation at i_{melt} .

Both of these parameters have current density counterparts, j_{max} and j_{melt} , that you can specify if you want the absolute current values to depend on the device area.

Auto Model Selection

Many models need to be characterized for different geometries in order to obtain accurate results for model development. The model selector program automatically searches for a model with the length and width range specified in the instance statement and uses this model in the simulations.

For the auto model selector program to find a specific model, the models to be searched should be grouped together within braces. Such a group is called a model group. An opening brace is required at the end of the line defining each model group. Every model in the group is given a name followed by a colon and the list of parameters. Also, the four geometric parameters l_{max} , l_{min} , w_{max} , and w_{min} should be given. The selection criteria to choose a model is as follows:

$$l_{\text{min}} \leq \text{inst_length} < l_{\text{max}} \quad \text{and} \quad w_{\text{min}} \leq \text{inst_width} < w_{\text{max}}$$

Spectre Circuit Simulator Reference

Component Statements Part I

Example:

```
model ModelName ModelType {  
    1: <model parameters> lmin=2 lmax=4 wmin=1 wmax=2  
    2: <model parameters> lmin=1 lmax=2 wmin=2 wmax=4  
    3: <model parameters> lmin=2 lmax=4 wmin=4 wmax=6  
}
```

Then for a given instance

```
M1 1 2 3 4 ModelName w=3 l=1.5
```

the program would search all the models in the model group with the name `ModelName` and then pick the first model whose geometric range satisfies the selection criteria. In the preceding example, the auto model selector program would choose `ModelName.2`.

You must specify both length (*l*) and width (*w*) on the device instance line to enable automatic model selection.

Output Parameters

1	<code>w_{eff}</code> (m)	Effective channel width.
2	<code>l_{eff}</code> (m)	Effective channel length.
3	<code>r_{seff}</code> (Ω)	Effective source resistance.
4	<code>r_{deff}</code> (Ω)	Effective drain resistance.
5	<code>a_{seff}</code> (m ²)	Effective area of source diffusion.
6	<code>a_{deff}</code> (m ²)	Effective area of drain diffusion.
7	<code>p_{seff}</code> (m)	Effective perimeter of source diffusion.
8	<code>p_{deff}</code> (m)	Effective perimeter of source diffusion.
9	<code>i_{sseff}</code> (A)	Effective source-bulk junction reverse saturation current.
10	<code>i_{sdeff}</code> (A)	Effective drain-bulk junction reverse saturation current.

Spectre Circuit Simulator Reference

Component Statements Part I

11	<code>cbseff (F)</code>	Effective zero-bias source-bulk junction capacitance.
12	<code>cbdeff (F)</code>	Effective zero-bias drain-bulk junction capacitance.
13	<code>vto (V)</code>	Effective zero-bias threshold voltage.
14	<code>vfb (V)</code>	Effective flat-band voltage.
15	<code>phi (V)</code>	Effective surface potential.
16	<code>k1 (\sqrt{V})</code>	Effective body-effect coefficient.
17	<code>k2</code>	Effective charge-sharing parameter.
18	<code>eta</code>	Effective DIBL coefficient.

Operating-Point Parameters

1	<code>type=n</code>	Transistor type. Possible values are <code>n</code> or <code>p</code> .
2	<code>region=triode</code>	Estimated operating region. Spectre outputs number (0-4) in a rawfile. Possible values are <code>off</code> , <code>triode</code> , <code>sat</code> , <code>subth</code> , or <code>breakdown</code> .
3	<code>degradation=no</code>	Hot-electron degradation flag. Possible values are <code>no</code> or <code>yes</code> .
4	<code>reversed</code>	Reverse mode indicator. Possible values are <code>no</code> or <code>yes</code> .
5	<code>ids (A)</code>	Resistive drain-to-source current.
6	<code>vgs (V)</code>	Gate-source voltage.
7	<code>vds (V)</code>	Drain-source voltage.
8	<code>vbs (V)</code>	Bulk-source voltage.
9	<code>vth (V)</code>	Threshold voltage.
10	<code>vdsat (V)</code>	Drain-source saturation voltage.

Spectre Circuit Simulator Reference

Component Statements Part I

11	betaeff (A/V ²)	Effective beta.
12	gm (S)	Common-source transconductance.
13	gds (S)	Common-source output conductance.
14	gmbs (S)	Body-transconductance.
15	cbd (F)	Drain-bulk junction capacitance.
16	cbs (F)	Source-bulk junction capacitance.
17	cgs (F)	Gate-source capacitance.
18	cgd (F)	Gate-drain capacitance.
19	cgb (F)	Gate-bulk capacitance.
20	ron (Ω)	ON-resistance.
21	id (A)	Resistive drain current.
22	ibulk (A)	Resistive bulk current.
23	pwr (W)	Power at op point.
24	gmoverid (1/V)	Gm/Ids.
25	isub (A)	Substrate current.
26	stress	Hot-electron stress.
27	age (s)	Device age.
28	he_vdsat (V)	hot electron vdsat.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the

Spectre Circuit Simulator Reference

Component Statements Part I

description for that parameter. For example, a reference of M-35 means the 35th model parameter.

ad	I-4	jmelt	M-135	nd	M-78	uto	M-101
ad	M-157	js	M-131	noisemod	M-164	vbox	M-151
adefeff	O-6	k1	O-16	nrd	M-160	vbs	OP-8
af	M-166	k1	M-10	nrd	I-7	vbsn	M-185
age	OP-27	k2	M-14	nrs	M-161	vdd	M-153
ai0	M-83	k2	O-17	nrs	I-8	vds	OP-7
alarm	M-147	kf	M-165	pai0	M-86	vdsat	OP-10
as	I-3	l	M-155	pb	M-141	vdsng	M-188
as	M-156	l	I-2	pbi0	M-90	vdsni	M-186
aseff	O-5	lai0	M-84	pbsw	M-145	version	M-100
betaeff	OP-11	lbi0	M-88	pd	M-159	vfb	O-14
bi0	M-87	lc0	M-212	pd	I-6	vfb0	M-2
bvj	M-150	lc1	M-215	pdeff	O-8	vgs	OP-6
capmod	M-115	lc2	M-218	peta	M-21	vgsng	M-189
cbd	OP-15	lc3	M-221	phi	O-15	vgsni	M-187
cbd	M-138	lc4	M-224	phi0	M-6	vpb	M-193
cbdeff	O-12	lc5	M-227	pk1	M-13	vpg	M-192
cbs	M-137	lc6	M-230	pk2	M-17	vth	OP-9
cbs	OP-16	lc7	M-233	pmus	M-41	vto	O-13
cbseff	O-11	ld	I-9	pmuz	M-33	w	I-1
cgb	OP-19	ldd	M-162	pn0	M-73	w	M-154
cgbo	M-113	ldif	M-127	pnb	M-77	wai0	M-85
cgd	OP-18	lds	M-163	pnd	M-81	wbi0	M-89
cgdo	M-112	lecrit0	M-204	pphi	M-9	wecrit0	M-205
cgs	OP-17	lecritb	M-210	ps	M-158	wecritb	M-211
cgso	M-111	lecritg	M-207	ps	I-5	wecritg	M-208
cj	M-139	leff	O-2	pseff	O-7	weff	O-1
cjsw	M-143	leta	M-19	pu0	M-53	weta	M-20
crigm	M-181	lgcd	M-129	pu1	M-61	wk1	M-12

Spectre Circuit Simulator Reference Component Statements Part I

criids M-182	lgcs M-128	pvfb M-5	wk2 M-16
criuo M-180	lk1 M-11	pwr OP-23	wlc0 M-214
crivth M-179	lk2 M-15	px2e M-25	wlc1 M-217
degradation OP-3	llc0 M-213	px2ms M-45	wlc2 M-220
degradation I-14	llc1 M-216	px2mz M-37	wlc3 M-223
degradation M-174	llc2 M-219	px2u0 M-57	wlc4 M-226
degramod M-173	llc3 M-222	px2u1 M-65	wlc5 M-229
dl0 M-91	llc4 M-225	px3e M-29	wlc6 M-232
dskip M-136	llc5 M-228	px3ms M-49	wlc7 M-235
duoc M-177	llc6 M-231	px3u1 M-69	wmax M-169
duoe M-178	llc7 M-234	rd M-119	wmin M-170
dvthc M-175	lmax M-171	rdc M-122	wmus M-40
dvthe M-176	lmin M-172	rdd M-124	wmuz M-32
dw0 M-92	lmus M-39	rdeff O-4	wn0 M-72
ecrit0 M-203	lmuz M-31	region OP-2	wnb M-76
ecritb M-209	ln0 M-71	region I-12	wnd M-80
ecritg M-206	lnb M-75	reversed OP-4	wnoi M-168
ef M-167	lnd M-79	ron OP-20	wnom M-183
eg M-105	lnom M-184	rs M-118	wphi M-8
esat M-190	lphi M-7	rsc M-121	wref M-94
esatg M-191	lref M-93	rseff O-3	wu0 M-52
eta O-18	ls I-10	rsh M-120	wu1 M-60
eta0 M-18	lu0 M-51	rss M-123	wvfb M-4
fc M-142	lu1 M-59	sc M-130	wx2e M-24
fcsw M-146	lvfb M-3	strc M-197	wx2ms M-44
gap1 M-106	lx2e M-23	stre M-198	wx2mz M-36
gap2 M-107	lx2ms M-43	stress OP-26	wx2u0 M-56
gds OP-13	lx2mz M-35	subc1 M-194	wx2u1 M-64
gm OP-12	lx2u0 M-55	subc2 M-195	wx3e M-28
gmbs OP-14	lx2u1 M-63	sube M-196	wx3ms M-48
gmoverid OP-24	lx3e M-27	subthmod M-82	wx3u1 M-68
h0 M-199	lx3ms M-47	temp M-97	x2e M-22

Spectre Circuit Simulator Reference Component Statements Part I

hdif	M-126	lx3u1	M-67	tempmod	M-99	x2ms	M-42
he_vdsat	OP-28	m	I-11	tlev	M-103	x2mz	M-34
hgd	M-200	m0	M-201	tlevc	M-104	x2u0	M-54
ibulk	OP-22	meto	M-114	tox	M-152	x2u1	M-62
id	OP-21	mgd	M-202	trd	M-109	x3e	M-26
ids	OP-5	minr	M-125	trise	M-98	x3ms	M-46
imax	M-148	mj	M-140	trise	I-13	x3u1	M-66
imelt	M-134	mjsw	M-144	trs	M-108	x1	M-96
is	M-132	mus	M-38	type	OP-1	xpart	M-116
isdeff	O-10	muz	M-30	type	M-1	xqc	M-117
isseff	O-9	n	M-133	u00	M-50	xti	M-110
isub	OP-25	n0	M-70	u10	M-58	xw	M-95
jmax	M-149	nb	M-74	ute	M-102		

BSIM2 Field Effect Transistor (bsim2)

Description

BSIM2 is a semiempirical MOSFET model developed at the University of California, Berkeley. All the model parameters are extracted directly from physical devices. Both the drain current and output resistance are accurately modeled. Three charge models are available. In SPICE mode, you can refer to BSIM2 as MOS level 5 or BSIM level 2. BSIM2 transistors require that you use a model statement.

This device is supported within altergroups.

Sample Instance Statement

```
m2 (0 2 1 1) pchmod l=5u w=10u as=40u ad=40u pd=28u ps=28u m=1
```

Sample Model Statement

```
model pchmod bsim2 type=p vfb0=-0.5 lvfb=0.5 wvfb=0.3 phi0=0.8 eta0=0.056 k1=0.5  
eg=0.99 gap1=5.5e-04 trs=1e-3 trd=1e-3 xpart=0.5 rs=10 rd=10
```

Spectre Circuit Simulator Reference

Component Statements Part I

Instance Definition

Name d g s b ModelName parameter=value ...

Instance Parameters

1	w (m)	Channel width.
2	l (m)	Channel length.
3	as (m ²)	Area of source diffusion.
4	ad (m ²)	Area of drain diffusion.
5	ps (m)	Perimeter of source diffusion.
6	pd (m)	Perimeter of drain diffusion.
7	nrd (m/m)	Number of squares of drain diffusion.
8	nrs (m/m)	Number of squares of source diffusion.
9	ld (m)	Drain diffusion length.
10	ls (m)	Source diffusion length.
11	m=1	Multiplicity factor (number of MOSFETs in parallel).
12	region=triode	Estimated operating region. Spectre outputs number (0-4) in a rawfile. Possible values are off, triode, sat, subth, or breakdown.
13	trise (C)	Temperature rise from ambient.
14	degradation=no	Hot-electron degradation flag. Possible values are no or yes.

Model Definition

model modelName bsim2 parameter=value ...=6

Spectre Circuit Simulator Reference

Component Statements Part I

Model Parameters

Device type parameters

1 `type=n` Transistor type.
Possible values are n or p.

Threshold voltage parameters

2 `vfb0=-0.8 V` Flat-band voltage.

3 `lvfb=0 V μ m` Length dependence of `vfb`.

4 `wvfb=0 V μ m` Width dependence of `vfb`.

5 `pvfb=0 V μ m` Width-length dependence of `vfb`.

6 `phi0=0.75 V` Surface potential.

7 `lphi=0 V μ m` Length dependence of `phi`.

8 `wphi=0 V μ m` Width dependence of `phi`.

9 `pphi=0 V μ m` Width-length dependence of `phi`.

10 `k1=0.7 \sqrt{V}` Body-effect coefficient.

11 `lk1=0 \sqrt{V} μ m` Length dependence of `k1`.

12 `wk1=0 \sqrt{V} μ m` Width dependence of `k1`.

13 `pk1=0 \sqrt{V} μ m` Width-length dependence of `k1`.

14 `k2=0` Charge-sharing parameter.

15 `lk2=0 μ m` Length dependence of `k2`.

16 `wk2=0 μ m` Width dependence of `k2`.

17 `pk2=0 μ m` Width-length dependence of `k2`.

Spectre Circuit Simulator Reference

Component Statements Part I

- 18 $\text{eta0}=0$ Drain-induced barrier-lowering coefficient.
- 19 $\text{l eta0}=0 \mu\text{m}$ Length dependence of eta0 .
- 20 $\text{w eta0}=0 \mu\text{m}$ Width dependence of eta0 .
- 21 $\text{p eta0}=0 \mu\text{m}$ Width-length dependence of eta0 .
- 22 $\text{etab}=0 \text{ 1/V}$ Body-bias dependence of eta0 .
- 23 $\text{l etab}=0 \mu\text{m/V}$ Length dependence of etab .
- 24 $\text{w etab}=0 \mu\text{m/V}$ Width dependence of etab .
- 25 $\text{p etab}=0 \mu\text{m/V}$ Width-length dependence of etab .

Mobility parameters

- 26 $\text{mu0}=400 \text{ cm}^2/\text{V s}$ Low-field mobility.
- 27 $\text{l mu0}=0 \text{ cm}^2/\text{V s}$ Length dependence of mu0 .
- 28 $\text{w mu0}=0 \text{ cm}^2/\text{V s}$ Width dependence of mu0 .
- 29 $\text{p mu0}=0 \text{ cm}^2/\text{V s}$ Width-length dependence of mu0 .
- 30 $\text{mu0b}=0 \text{ cm}^2/\text{V}^2 \text{ s}$ Body-bias dependence of mu0 .
- 31 $\text{l mu0b}=0 \text{ cm}^2 \mu\text{m}/\text{V}^2 \text{ s}$
Length dependence of x2mz .
- 32 $\text{w mu0b}=0 \text{ cm}^2 \mu\text{m}/\text{V}^2 \text{ s}$
Width dependence of x2mz .
- 33 $\text{p mu0b}=0 \text{ cm}^2 \mu\text{m}/\text{V}^2 \text{ s}$
Width-length dependence of x2mz .
- 34 $\text{mus0}=450 \text{ cm}^2/\text{V s}$ Mobility in the saturation region.
- 35 $\text{l mus0}=0 \text{ cm}^2 \mu\text{m}/\text{V s}$
Length dependence of mus0 .

Spectre Circuit Simulator Reference

Component Statements Part I

- 36 $w_{\mu s0}=0$ $\text{cm}^2 \mu\text{m}/\text{V s}$ Width dependence of μ_{s0} .
- 37 $p_{\mu s0}=0$ $\text{cm}^2 \mu\text{m}/\text{V s}$ Width-length dependence of μ_{s0} .
- 38 $\mu_{sb}=0$ $\text{cm}^2/\text{V}^2 \text{ s}$ Body-bias dependence of μ_{s0} .
- 39 $l_{\mu sb}=0$ $\text{cm}^2 \mu\text{m}/\text{V}^2 \text{ s}$ Length dependence of μ_{s0} .
- 40 $w_{\mu sb}=0$ $\text{cm}^2 \mu\text{m}/\text{V}^2 \text{ s}$ Length dependence of μ_{s0} .
- 41 $p_{\mu sb}=0$ $\text{cm}^2 \mu\text{m}/\text{V}^2 \text{ s}$ Length dependence of μ_{s0} .
- 42 $\mu_{20}=1$ Empirical channel length modulation parameter.
- 43 $l_{\mu 20}=0$ μm Length dependence of μ_{20} .
- 44 $w_{\mu 20}=0$ μm Width dependence of μ_{20} .
- 45 $p_{\mu 20}=0$ μm Width-length dependence of μ_{20} .
- 46 $\mu_{2b}=0$ $1/\text{V}$ Body-bias dependence of μ_{20} .
- 47 $l_{\mu 2b}=0$ $\mu\text{m}/\text{V}$ Length dependence of μ_{2b} .
- 48 $w_{\mu 2b}=0$ $\mu\text{m}/\text{V}$ Width dependence of μ_{2b} .
- 49 $p_{\mu 2b}=0$ $\mu\text{m}/\text{V}$ Width-length dependence of μ_{2b} .
- 50 $\mu_{2g}=0$ $1/\text{V}$ Gate-bias dependence of μ_{20} .
- 51 $l_{\mu 2g}=0$ $\mu\text{m}/\text{V}$ Length dependence of μ_{2g} .
- 52 $w_{\mu 2g}=0$ $\mu\text{m}/\text{V}$ Width dependence of μ_{2g} .
- 53 $p_{\mu 2g}=0$ $\mu\text{m}/\text{V}$ Width-length dependence of μ_{2g} .
- 54 $\mu_{30}=5$ $\text{cm}^2/\text{V}^2 \text{ s}$ Empirical output resistance parameter.

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- 55 $l_{\mu 30}=0$ $\text{cm}^2 \mu\text{m}/\text{V}^2 \text{ s}$ Length dependence of μ_{30} .
- 56 $w_{\mu 30}=0$ $\text{cm}^2 \mu\text{m}/\text{V}^2 \text{ s}$ Width dependence of μ_{30} .
- 57 $p_{\mu 30}=0$ $\text{cm}^2 \mu\text{m}/\text{V}^2 \text{ s}$ Width-length dependence of μ_{30} .
- 58 $\mu_{3b}=0$ $\text{cm}^2/\text{V}^3 \text{ s}$ Body-bias dependence of μ_{30} .
- 59 $l_{\mu 3b}=0$ $\text{cm}^2 \mu\text{m}/\text{V}^3 \text{ s}$ Length dependence of μ_{3b} .
- 60 $w_{\mu 3b}=0$ $\text{cm}^2 \mu\text{m}/\text{V}^3 \text{ s}$ Width dependence of μ_{3b} .
- 61 $p_{\mu 3b}=0$ $\text{cm}^2 \mu\text{m}/\text{V}^3 \text{ s}$ Width-length dependence of μ_{3b} .
- 62 $\mu_{3g}=0$ $\text{cm}^2/\text{V}^3 \text{ s}$ Gate-bias dependence of μ_{30} .
- 63 $l_{\mu 3g}=0$ $\text{cm}^2 \mu\text{m}/\text{V}^3 \text{ s}$ Length dependence of μ_{3g} .
- 64 $w_{\mu 3g}=0$ $\text{cm}^2 \mu\text{m}/\text{V}^3 \text{ s}$ Width dependence of μ_{3g} .
- 65 $p_{\mu 3g}=0$ $\text{cm}^2 \mu\text{m}/\text{V}^3 \text{ s}$ Width-length dependence of μ_{3g} .
- 66 $\mu_{40}=0$ $\text{cm}^2/\text{V}^3 \text{ s}$ Empirical output resistance parameter.
- 67 $l_{\mu 40}=0$ $\text{cm}^2 \mu\text{m}/\text{V}^3 \text{ s}$ Length dependence of μ_{40} .
- 68 $w_{\mu 40}=0$ $\text{cm}^2 \mu\text{m}/\text{V}^3 \text{ s}$ Width dependence of μ_{40} .
- 69 $p_{\mu 40}=0$ $\text{cm}^2 \mu\text{m}/\text{V}^3 \text{ s}$ Width-length dependence of μ_{40} .
- 70 $\mu_{4b}=0$ $\text{cm}^2/\text{V}^3 \text{ s}$ Empirical output resistance parameter.

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Component Statements Part I

- 71 $l_{\mu 4b=0}$ $\text{cm}^2 \mu\text{m}/\text{V}^3 \text{ s}$ Length dependence of μ_{4b} .
- 72 $w_{\mu 4b=0}$ $\text{cm}^2 \mu\text{m}/\text{V}^3 \text{ s}$ Width dependence of μ_{4b} .
- 73 $p_{\mu 4b=0}$ $\text{cm}^2 \mu\text{m}/\text{V}^3 \text{ s}$ Width-length dependence of μ_{4b} .
- 74 $\mu_{4g=0}$ $\text{cm}^2/\text{V}^3 \text{ s}$ Gate-bias dependence of μ_{4g} .
- 75 $l_{\mu 4g=0}$ $\text{cm}^2 \mu\text{m}/\text{V}^3 \text{ s}$ Length dependence of μ_{4g} .
- 76 $w_{\mu 4g=0}$ $\text{cm}^2 \mu\text{m}/\text{V}^3 \text{ s}$ Width dependence of μ_{4g} .
- 77 $p_{\mu 4g=0}$ $\text{cm}^2 \mu\text{m}/\text{V}^3 \text{ s}$ Width-length dependence of μ_{4g} .

Mobility modulation parameters

- 78 $u_{a0=0}$ $1/\text{V}$ Gate voltage dependence of mobility.
- 79 $l_{u_{a0=0}}$ $\mu\text{m}/\text{V}$ Length dependence of u_{a0} .
- 80 $w_{u_{a0=0}}$ $\mu\text{m}/\text{V}$ Width dependence of u_{a0} .
- 81 $p_{u_{a0=0}}$ $\mu\text{m}/\text{V}$ Width-length dependence of u_{a0} .
- 82 $u_{ab=0}$ $1/\text{V}^2$ Body-bias dependence of u_a .
- 83 $l_{u_{ab=0}}$ $\mu\text{m}/\text{V}^2$ Length dependence of u_{ab} .
- 84 $w_{u_{ab=0}}$ $\mu\text{m}/\text{V}^2$ Width dependence of u_{ab} .
- 85 $p_{u_{ab=0}}$ $\mu\text{m}/\text{V}^2$ Width-length dependence of u_{ab} .
- 86 $u_{b0=0}$ $1/\text{V}^2$ Second-order effect of gate voltage dependence of mobility.
- 87 $l_{u_{b0=0}}$ $\mu\text{m}/\text{V}^2$ Length dependence of u_{b0} .

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- 88 $w_{ub0}=0 \mu\text{m}/\text{V}^2$ Width dependence of u_{b0} .
- 89 $p_{ub0}=0 \mu\text{m}/\text{V}^2$ Width-length dependence of u_{b0} .
- 90 $u_{bb}=0 1/\text{V}^3$ Body-bias dependence of u_b .
- 91 $l_{ubb}=0 \mu\text{m}/\text{V}^3$ Length dependence of u_{bb} .
- 92 $w_{ubb}=0 \mu\text{m}/\text{V}^3$ Width dependence of u_{bb} .
- 93 $p_{ubb}=0 \mu\text{m}/\text{V}^3$ Width-length dependence of u_{bb} .

Velocity saturation parameters

- 94 $u_{10}=0 1/\text{V}$ Velocity saturation coefficient.
- 95 $l_{u10}=0 \mu\text{m}/\text{V}$ Length dependence of u_1 .
- 96 $w_{u10}=0 \mu\text{m}/\text{V}$ Width dependence of u_1 .
- 97 $p_{u10}=0 \mu\text{m}/\text{V}$ Width-length dependence of u_1 .
- 98 $u_{1b}=0 1/\text{V}^2$ Body-bias dependence of u_1 .
- 99 $l_{u1b}=0 \mu\text{m}/\text{V}^2$ Length dependence of u_{1b} .
- 100 $w_{u1b}=0 \mu\text{m}/\text{V}^2$ Width dependence of u_{1b} .
- 101 $p_{u1b}=0 \mu\text{m}/\text{V}^2$ Width-length dependence of u_{1b} .
- 102 $u_{1d}=0 1/\text{V}^2$ Drain-bias dependence of u_1 .
- 103 $l_{u1d}=0 \mu\text{m}/\text{V}^2$ Length dependence of u_{1d} .
- 104 $w_{u1d}=0 \mu\text{m}/\text{V}^2$ Width dependence of u_{1d} .
- 105 $p_{u1d}=0 \mu\text{m}/\text{V}^2$ Width-length dependence of u_{1d} .

Subthreshold parameters

- 106 $n_0=0$ Subthreshold swing parameter.

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107	$ln0=0$	μm	Length dependence of subthreshold swing parameter.
108	$wn0=0$	μm	Width dependence of subthreshold swing parameter.
109	$pnb=0$	μm	Width-length dependence of subthreshold swing parameter.
110	$nb=0$	\sqrt{V}	Body-bias dependence of $n0$.
111	$lnb=0$	\sqrt{V} μm	Length dependence of nb .
112	$wnb=0$	\sqrt{V} μm	Width dependence of nb .
113	$pnb=0$	\sqrt{V} μm	Width-length dependence of nb .
114	$nd=0$	$1/V$	Drain-bias dependence of $n0$.
115	$lnd=0$	$\mu\text{m}/V$	Length dependence of nd .
116	$wnd=0$	$\mu\text{m}/V$	Width dependence of nd .
117	$pnd=0$	$\mu\text{m}/V$	Width-length dependence of nd .
118	$vof0=1$	V	Threshold voltage offset in the subthreshold region.
119	$lvof0=0$	V μm	Length dependence of vof .
120	$wvof0=0$	V μm	Width dependence of vof .
121	$pvof0=0$	V μm	Width-length dependence of vof .
122	$vofb=0$		Body-bias dependence of $vof0$.
123	$lvofb=0$	μm	Length dependence of $vofb$.
124	$wvofb=0$	μm	Width dependence of $vofb$.
125	$pvofb=0$	μm	Width-length dependence of $vofb$.
126	$vofd=0$		Drain-bias dependence of $vof0$.
127	$lvofd=0$	μm	Length dependence of $vofd$.
128	$wvofd=0$	μm	Width dependence of $vofd$.

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- 129 $p_{vofd}=0 \mu\text{m}$ Width-length dependence of $vofd$.
130 $subthmod=2$ Subthreshold model selector.

Impact ionization parameters

- 131 $ai0=0 \text{ 1/V}$ Hot-electron effect on R_{out} parameter.
132 $lai0=0 \mu\text{m/V}$ Length dependence of $ai0$.
133 $wai0=0 \mu\text{m/V}$ Width dependence of $ai0$.
134 $pai0=0 \mu\text{m/V}$ Width-length dependence of $ai0$.
135 $aib=0 \text{ 1/V}^2$ Body-bias dependence of $ai0$.
136 $laib=0 \mu\text{m/V}^2$ Length dependence of aib .
137 $waib=0 \mu\text{m/V}^2$ Width dependence of aib .
138 $paib=0 \mu\text{m/V}^2$ Width-length dependence of aib .
139 $bi0=0 \text{ V}$ Hot-electron effect on R_{out} exponent.
140 $lbi0=0 \text{ V } \mu\text{m}$ Length dependence of $bi0$.
141 $wbi0=0 \text{ V } \mu\text{m}$ Width dependence of $bi0$.
142 $pbi0=0 \text{ V } \mu\text{m}$ Width-length dependence of $bi0$.
143 $bib=0$ Body-bias dependence of $bi0$.
144 $lbib=0 \mu\text{m}$ Length dependence of bib .
145 $wbib=0 \mu\text{m}$ Width dependence of bib .
146 $pbib=0 \mu\text{m}$ Width-length dependence of bib .

Transition region bound parameters

- 147 $vghigh=0.2 \text{ V}$ Upper bound of the transition region.

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148	lvghigh=0 V μm	Length dependence of V_{ghigh} .
149	wvghigh=0 V μm	Width dependence of V_{ghigh} .
150	pvgghigh=0 V μm	Width-length dependence of V_{ghigh} .
151	vglow=-0.15 V	Lower bound of the transition region.
152	lvglow=0 V μm	Length dependence of V_{glow} .
153	wvglow=0 V μm	Width dependence of V_{glow} .
154	pvgglow=0 V μm	Width-length dependence of V_{glow} .

Length and width modulation parameters

155	dl0=0 μm	Lateral diffusion.
156	dw0=0 μm	Field oxide encroachment.
157	lref= ∞ m	Reference channel length.
158	wref= ∞ m	Reference channel width.
159	xw=0 m	Width variation due to masking and etching.
160	xl=0 m	Length variation due to masking and etching.

Temperature effects parameters

161	temp (C)	Parameters measurement temperature. Default set by options.
162	trise=0 C	Temperature rise from ambient.
163	tempmod=432	Temperature model selector.
164	version=432	Version selector.
165	uto=0 C	Mobility temperature offset.
166	ute=-1.5	Mobility temperature exponent.

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167	<code>tlev=0</code>	DC temperature selector.
168	<code>tlevc=0</code>	AC temperature selector.
169	<code>ptc=0 V/C</code>	Surface potential temperature coefficient.
170	<code>eg=1.12452 V</code>	Energy band gap.
171	<code>gap1=7.02e-4 V/C²</code>	Band gap temperature coefficient.
172	<code>gap2=1108 K</code>	Band gap temperature offset.
173	<code>trs=0 1/C</code>	Temperature coefficient for source resistance.
174	<code>trd=0 1/C</code>	Temperature coefficient for drain resistance.
175	<code>x_{ti}=3</code>	Saturation current temperature exponent.

Overlap capacitance parameters

176	<code>cgso=0 F/m</code>	Gate-source overlap capacitance.
177	<code>cgdo=0 F/m</code>	Gate-drain overlap capacitance.
178	<code>cgbo=0 F/m</code>	Gate-bulk overlap capacitance.
179	<code>meto=0 m</code>	Metal overlap in fringing field.

Charge model selection parameters

180	<code>capmod=bsim</code>	Intrinsic charge model. Possible values are <code>none</code> , <code>meyer</code> , <code>yang</code> , or <code>bsim</code> .
181	<code>x_{part}=1</code>	Drain/source channel charge partition in saturation for BSIM charge model, use 0.0 for 40/60, 0.5 for 50/50, or 1.0 for 0/100.
182	<code>x_{qc}=0</code>	Drain/source channel charge partition in saturation for charge models, e.g. use 0.4 for 40/60, 0.5 for 50/50, or 0 for 0/100.

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Component Statements Part I

Parasitic resistance parameters

183	$r_s=0 \ \Omega$	Source resistance.
184	$r_d=0 \ \Omega$	Drain resistance.
185	$r_{sh}=0 \ \Omega/\text{sqr}$	Source/drain diffusion sheet resistance.
186	$r_{sc}=0 \ \Omega$	Source contact resistance.
187	$r_{dc}=0 \ \Omega$	Drain contact resistance.
188	$r_{ss}=0 \ \Omega \ \text{m}$	Scalable source resistance.
189	$r_{dd}=0 \ \Omega \ \text{m}$	Scalable drain resistance.
190	$\text{minr}=0.1 \ \Omega$	Minimum source/drain resistance.
191	$\text{hdif}=0 \ \text{m}$	Length of heavily doped diffusion.
192	$\text{ldif}=0 \ \text{m}$	Lateral diffusion beyond the gate.
193	$\text{lgcs}=0 \ \text{m}$	Gate-to-contact length of source side.
194	$\text{lgcd}=0 \ \text{m}$	Gate-to-contact length of drain side.
195	$\text{sc}=\infty \ \text{m}$	Spacing between contacts.

Junction diode parameters

196	$j_s \ (\text{A}/\text{m}^2)$	Bulk junction reverse saturation current density.
197	$i_s=1\text{e-}14 \ \text{A}$	Bulk junction reverse saturation current.
198	$n=1$	Junction emission coefficient.
199	$\text{dskip}=\text{yes}$	Use simple piece-wise linear model for diode currents below $0.1 \cdot i_{\text{abstol}}$. Possible values are <code>no</code> or <code>yes</code> .
200	$\text{imelt}=\text{'imaxA'}$	Explosion current, diode is linearized beyond this current to aid convergence.

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201 $j_{melt} = j_{max} A/m^2$ Explosion current density, diode is linearized beyond this current to aid convergence.

Operating region warning control parameters

202 $alarm = none$ Forbidden operating region.
Possible values are `none`, `off`, `triode`, `sat`, `subth`, or `rev`.

203 $i_{max} = 1 A$ Maximum current, currents above this limit generate a warning.

204 $j_{max} = 1e8 A/m^2$ Maximum current density, currents above this limit generate a warning.

205 $b_{vj} = \infty V$ Junction reverse breakdown voltage.

206 $v_{box} = 1e9 \text{ tox} V$ Oxide breakdown voltage.

Junction capacitance model parameters

207 $c_{bs} = 0 F$ Bulk-source zero-bias junction capacitance.

208 $c_{bd} = 0 F$ Bulk-drain zero-bias junction capacitance.

209 $c_j = 0 F/m^2$ Zero-bias junction bottom capacitance density.

210 $m_j = 1/2$ Bulk junction bottom grading coefficient.

211 $p_b = 0.8 V$ Bulk junction potential.

212 $f_c = 0.5$ Forward-bias depletion capacitance threshold.

213 $c_{jsw} = 0 F/m$ Zero-bias junction sidewall capacitance density.

214 $m_{jsw} = 1/3$ Bulk junction sidewall grading coefficient.

215 $p_{bsw} = 0.8 V$ Side-wall junction potential.

216 $f_{csw} = 0.5$ Side-wall forward-bias depletion capacitance threshold.

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Process and power supply parameters

217	$t_{ox}=4e-8$ m	Gate oxide thickness.
218	$v_{dd}=5$ V	Drain voltage at which parameters are extracted.
219	$v_{gg}=5$ V	Gate voltage at which parameters are extracted.
220	$v_{bb}=-5$ V	Body voltage at which parameters are extracted.

Default device parameters

221	$w=3e-6$ m	Channel width.
222	$l=3e-6$ m	Channel length.
223	$a_s=0$ m ²	Area of source diffusion.
224	$a_d=0$ m ²	Area of drain diffusion.
225	$p_s=0$ m	Perimeter of source diffusion.
226	$p_d=0$ m	Perimeter of drain diffusion.
227	$n_{rd}=0$ m/m	Number of squares of drain diffusion.
228	$n_{rs}=0$ m/m	Number of squares of source diffusion.
229	$l_{dd}=0$ m	Drain diffusion length.
230	$l_{ds}=0$ m	Source diffusion length.

Noise model parameters

231	$noisemod=1$	Noise model selector.
232	$k_f=0$	Flicker (1/f) noise coefficient.
233	$a_f=1$	Flicker (1/f) noise exponent.
234	$e_f=1$	Flicker (1/f) noise frequency exponent.

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235 `wnoi=1e-5 m` Channel width at which noise parameters were extracted.

Auto Model Selector parameters

236 `wmax=1.0 m` Maximum channel width for which the model is valid.

237 `wmin=0.0 m` Minimum channel width for which the model is valid.

238 `lmax=1.0 m` Maximum channel length for which the model is valid.

239 `lmin=0.0 m` Minimum channel length for which the model is valid.

Degradation parameters

240 `degramod=spectre` Degradation model selector.
Possible values are `spectre` or `bert`.

241 `degradation=no` Hot-electron degradation flag.
Possible values are `no` or `yes`.

242 `dvthc=1 V` Degradation coefficient for threshold voltage.

243 `dvthe=1` Degradation exponent for threshold voltage.

244 `duoc=1 S` Degradation coefficient for transconductance.

245 `duoe=1` Degradation exponent for transconductance.

246 `crivth=0.1 V` Maximum allowable threshold voltage shift.

247 `criuo=10%` Maximum allowable normalized mobility change.

248 `crigm=10%` Maximum allowable normalized transconductance change.

249 `criids=10%` Maximum allowable normalized drain current change.

250 `wnom=5e-6 m` Nominal device width in degradation calculation.

251 `lnom=1e-6 m` Nominal device length in degradation calculation.

252 `vbsn=0 V` Substrate voltage in degradation calculation.

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253	$v_{dsni}=0.1$ V	Drain voltage in I_{ds} degradation calculation.
254	$v_{gsni}=5$ V	Gate voltage in I_{ds} degradation calculation.
255	$v_{dsng}=0.1$ V	Drain voltage in G_m degradation calculation.
256	$v_{gsng}=5$ V	Gate voltage in G_m degradation calculation.

Spectre stress parameters

257	$esat=1.1e7$ V/m	Critical field in v_{dsat} calculation.
258	$esatg=2.5e6$ 1/m	Gate voltage dependence of $esat$.
259	$v_{pg}=-0.25$	Gate voltage modifier.
260	$v_{pb}=-0.13$	Gate voltage modifier.
261	$subc1=2.24e-5$	Substrate current coefficient.
262	$subc2=-0.1e-5$ 1/V	Substrate current coefficient.
263	$sube=6.4$	Substrate current exponent.
264	$strc=1$	Stress coefficient.
265	$stre=1$	Stress exponent.

BERT stress parameters

266	$h0=1$	Aging coefficient.
267	$hgd=0$ 1/V	Bias dependence of $h0$.
268	$m0=1$	Aging exponent.
269	$mgd=0$ 1/V	Bias dependence of $m0$.
270	$ecrit0=1.1e5$ V/cm	Critical electric field.
271	$lecrit0=0$ μm V/cm	Length dependence of $ecrit0$.

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272	wecrit0=0	$\mu\text{m V/cm}$	Width dependence of ecrit0.
273	ecritg=0	1/cm	Gate voltage dependence of ecrit0.
274	lecritg=0	$\mu\text{m/cm}$	Length dependence of ecritg.
275	wecritg=0	$\mu\text{m/cm}$	Width dependence of ecritg.
276	ecritb=0	1/cm	Substrate voltage dependence of ecrit0.
277	lecritb=0	$\mu\text{m/cm}$	Length dependence of ecritb.
278	wecritb=0	$\mu\text{m/cm}$	Width dependence of ecritb.
279	lc0=1		Substrate current coefficient.
280	llc0=0	μm	Length dependence of lc0.
281	wlc0=0	μm	Width dependence of lc0.
282	lc1=1		Substrate current coefficient.
283	llc1=0	μm	Length dependence of lc1.
284	wlc1=0	μm	Width dependence of lc1.
285	lc2=1		Substrate current coefficient.
286	llc2=0	μm	Length dependence of lc2.
287	wlc2=0	μm	Width dependence of lc2.
288	lc3=1		Substrate current coefficient.
289	llc3=0	μm	Length dependence of lc3.
290	wlc3=0	μm	Width dependence of lc3.
291	lc4=1		Substrate current coefficient.
292	llc4=0	μm	Length dependence of lc4.
293	wlc4=0	μm	Width dependence of lc4.

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294	$lc5=1$	Substrate current coefficient.
295	$llc5=0$ μm	Length dependence of $lc5$.
296	$wlc5=0$ μm	Width dependence of $lc5$.
297	$lc6=1$	Substrate current coefficient.
298	$llc6=0$ μm	Length dependence of $lc6$.
299	$wlc6=0$ μm	Width dependence of $lc6$.
300	$lc7=1$	Substrate current coefficient.
301	$llc7=0$ μm	Length dependence of $lc7$.
302	$wlc7=0$ μm	Width dependence of $lc7$.

Imax and Imelt

The $imax$ parameter aids convergence and prevents numerical overflow. The junction characteristics of the device are accurately modeled for current up to $imax$. If $imax$ is exceeded during iterations, the linear model is substituted until the current drops below $imax$ or until convergence is achieved. If convergence is achieved with the current exceeding $imax$, the results are inaccurate, and Spectre prints a warning.

A separate model parameter, $imelt$, is used as a limit warning for the junction current. This parameter can be set to the maximum current rating of the device. When any component of the junction current exceeds $imelt$, note that base and collector currents are composed of many exponential terms, Spectre issues a warning and the results become inaccurate. The junction current is linearized above the value of $imelt$ to prevent arithmetic exception, with the exponential term replaced by a linear equation at $imelt$.

Both of these parameters have current density counterparts, $jmax$ and $jmelt$, that you can specify if you want the absolute current values to depend on the device area.

Auto Model Selection

Many models need to be characterized for different geometries in order to obtain accurate results for model development. The model selector program automatically searches for a model with the length and width range specified in the instance statement and uses this model in the simulations.

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For the auto model selector program to find a specific model, the models to be searched should be grouped together within braces. Such a group is called a model group. An opening brace is required at the end of the line defining each model group. Every model in the group is given a name followed by a colon and the list of parameters. Also, the four geometric parameters `lmax`, `lmin`, `wmax`, and `wmin` should be given. The selection criteria to choose a model is as follows:

$$lmin \leq inst_length < lmax \quad \text{and} \quad wmin \leq inst_width < wmax$$

Example:

```
model ModelName ModelType {  
    1: <model parameters> lmin=2 lmax=4 wmin=1 wmax=2  
    2: <model parameters> lmin=1 lmax=2 wmin=2 wmax=4  
    3: <model parameters> lmin=2 lmax=4 wmin=4 wmax=6  
}
```

Then for a given instance

```
M1 1 2 3 4 ModelName w=3 l=1.5
```

the program would search all the models in the model group with the name `ModelName` and then pick the first model whose geometric range satisfies the selection criteria. In the preceding example, the auto model selector program would choose `ModelName.2`.

You must specify both length (`l`) and width (`w`) on the device instance line to enable automatic model selection.

Output Parameters

1	<code>w_{eff}</code> (m)	Effective channel width.
2	<code>l_{eff}</code> (m)	Effective channel length.
3	<code>r_{seff}</code> (Ω)	Effective source resistance.
4	<code>r_{deff}</code> (Ω)	Effective drain resistance.
5	<code>a_{seff}</code> (m ²)	Effective area of source diffusion.

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6	<code>adef</code> (m ²)	Effective area of drain diffusion.
7	<code>pseff</code> (m)	Effective perimeter of source diffusion.
8	<code>pdeff</code> (m)	Effective perimeter of source diffusion.
9	<code>isseff</code> (A)	Effective source-bulk junction reverse saturation current.
10	<code>isdeff</code> (A)	Effective drain-bulk junction reverse saturation current.
11	<code>cbseff</code> (F)	Effective zero-bias source-bulk junction capacitance.
12	<code>cbdeff</code> (F)	Effective zero-bias drain-bulk junction capacitance.
13	<code>vt0</code> (V)	Effective zero-bias threshold voltage.
14	<code>vfb</code> (V)	Effective flat-band voltage.
15	<code>phi</code> (V)	Effective surface potential.
16	<code>k1</code> (\sqrt{V})	Effective body-effect coefficient.
17	<code>k2</code>	Effective charge-sharing parameter.
18	<code>eta</code>	Effective DIBL coefficient.

Operating-Point Parameters

1	<code>type=n</code>	Transistor type. Possible values are <code>n</code> or <code>p</code> .
2	<code>region=triode</code>	Estimated operating region. Spectre outputs number (0-4) in a <code>rawfile</code> . Possible values are <code>off</code> , <code>triode</code> , <code>sat</code> , <code>subth</code> , or <code>breakdown</code> .
3	<code>degradation=no</code>	Hot-electron degradation flag. Possible values are <code>no</code> or <code>yes</code> .
4	<code>reversed</code>	Reverse mode indicator. Possible values are <code>no</code> or <code>yes</code> .
5	<code>ids</code> (A)	Resistive drain-to-source current.

Spectre Circuit Simulator Reference

Component Statements Part I

6	vgs (V)	Gate-source voltage.
7	vds (V)	Drain-source voltage.
8	vbs (V)	Bulk-source voltage.
9	vth (V)	Threshold voltage.
10	vdsat (V)	Drain-source saturation voltage.
11	betaeff (A/V ²)	Effective beta.
12	gm (S)	Common-source transconductance.
13	gds (S)	Common-source output conductance.
14	gmbs (S)	Body-transconductance.
15	cbd (F)	Drain-bulk junction capacitance.
16	cbs (F)	Source-bulk junction capacitance.
17	cgs (F)	Gate-source capacitance.
18	cgd (F)	Gate-drain capacitance.
19	cgb (F)	Gate-bulk capacitance.
20	ron (Ω)	ON-resistance.
21	id (A)	Resistive drain current.
22	ibulk (A)	Resistive bulk current.
23	pwr (W)	Power at op point.
24	gmoverid (1/V)	Gm/Ids.
25	isub (A)	Substrate current.
26	stress	Hot-electron stress.
27	age (s)	Device age.

Spectre Circuit Simulator Reference Component Statements Part I

28 he_vdsat (V) hot electron vdsat.

Parameter Index

In the following index, I refers to instance parameters, M refers to the model parameters section, O refers to the output parameters section, and OP refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of M-35 means the 35th model parameter.

ad	M-224	lc1	M-282	nrs	I-8	ute	M-166
ad	I-4	lc2	M-285	nrs	M-228	uto	M-165
adefeff	O-6	lc3	M-288	pai0	M-134	vbb	M-220
af	M-233	lc4	M-291	paib	M-138	vbox	M-206
age	OP-27	lc5	M-294	pb	M-211	vbs	OP-8
ai0	M-131	lc6	M-297	pbi0	M-142	vbsn	M-252
aib	M-135	lc7	M-300	pbib	M-146	vdd	M-218
alarm	M-202	ld	I-9	pbsw	M-215	vds	OP-7
as	I-3	ldd	M-229	pd	I-6	vdsat	OP-10
as	M-223	ldif	M-192	pd	M-226	vdsng	M-255
aseff	O-5	lds	M-230	pdeff	O-8	vdsni	M-253
betaeff	OP-11	lecrit0	M-271	peta0	M-21	version	M-164
bi0	M-139	lecritb	M-277	petab	M-25	vfb	O-14
bib	M-143	lecritg	M-274	phi	O-15	vfb0	M-2
bvj	M-205	leff	O-2	phi0	M-6	vgg	M-219
capmod	M-180	leta0	M-19	pk1	M-13	vghigh	M-147
cbd	OP-15	letab	M-23	pk2	M-17	vglow	M-151
cbd	M-208	lgcd	M-194	pmu0	M-29	vgs	OP-6
cbdeff	O-12	lgcs	M-193	pmu0b	M-33	vgsng	M-256
cbs	OP-16	lk1	M-11	pmu20	M-45	vgsni	M-254
cbs	M-207	lk2	M-15	pmu2b	M-49	vof0	M-118
cbseff	O-11	llc0	M-280	pmu2g	M-53	vofb	M-122
cgb	OP-19	llc1	M-283	pmu30	M-57	vofd	M-126

Spectre Circuit Simulator Reference Component Statements Part I

cgbo	M-178	llc2	M-286	pmu3b	M-61	vpb	M-260
cgd	OP-18	llc3	M-289	pmu3g	M-65	vpg	M-259
cgdo	M-177	llc4	M-292	pmu40	M-69	vth	OP-9
cgs	OP-17	llc5	M-295	pmu4b	M-73	vto	O-13
cgso	M-176	llc6	M-298	pmu4g	M-77	w	M-221
cj	M-209	llc7	M-301	pmus0	M-37	w	I-1
cjsw	M-213	lmax	M-238	pmusb	M-41	wai0	M-133
crigm	M-248	lmin	M-239	pn0	M-109	waib	M-137
criids	M-249	lmu0	M-27	pnb	M-113	wbi0	M-141
criuo	M-247	lmu0b	M-31	pnd	M-117	wbib	M-145
crivth	M-246	lmu20	M-43	pphi	M-9	wecrit0	M-272
degradation	OP-3	lmu2b	M-47	ps	I-5	wecritb	M-278
degradation	M-241	lmu2g	M-51	ps	M-225	wecritg	M-275
degradation	I-14	lmu30	M-55	pseff	O-7	weff	O-1
degramod	M-240	lmu3b	M-59	ptc	M-169	weta0	M-20
dl0	M-155	lmu3g	M-63	pu10	M-97	wetab	M-24
dskip	M-199	lmu40	M-67	pulb	M-101	wk1	M-12
duoc	M-244	lmu4b	M-71	puld	M-105	wk2	M-16
duoe	M-245	lmu4g	M-75	pua0	M-81	wlc0	M-281
dvthc	M-242	lmus0	M-35	puab	M-85	wlc1	M-284
dvthe	M-243	lmusb	M-39	pub0	M-89	wlc2	M-287
dw0	M-156	ln0	M-107	pubb	M-93	wlc3	M-290
ecrit0	M-270	lnb	M-111	pvfb	M-5	wlc4	M-293
ecritb	M-276	lnd	M-115	pvghigh	M-150	wlc5	M-296
ecritg	M-273	lnom	M-251	pvglow	M-154	wlc6	M-299
ef	M-234	lphi	M-7	pvof0	M-121	wlc7	M-302
eg	M-170	lref	M-157	pvofb	M-125	wmax	M-236
esat	M-257	ls	I-10	pvofd	M-129	wmin	M-237
esatg	M-258	lu10	M-95	pwr	OP-23	wmu0	M-28
eta	O-18	lu1b	M-99	rd	M-184	wmu0b	M-32
eta0	M-18	lu1d	M-103	rdc	M-187	wmu20	M-44
etab	M-22	lua0	M-79	rdd	M-189	wmu2b	M-48

Spectre Circuit Simulator Reference Component Statements Part I

fc	M-212	luab	M-83	rdeff	O-4	wmu2g	M-52
fcs	M-216	lub0	M-87	region	OP-2	wmu30	M-56
gap1	M-171	lubb	M-91	region	I-12	wmu3b	M-60
gap2	M-172	lvfb	M-3	reversed	OP-4	wmu3g	M-64
gds	OP-13	lvghigh	M-148	ron	OP-20	wmu40	M-68
gm	OP-12	lvglow	M-152	rs	M-183	wmu4b	M-72
gmbs	OP-14	lvof0	M-119	rsc	M-186	wmu4g	M-76
gmoverid	OP-24	lvofb	M-123	rseff	O-3	wmus0	M-36
h0	M-266	lvofd	M-127	rsh	M-185	wmusb	M-40
hdif	M-191	m	I-11	rss	M-188	wn0	M-108
he_vdsat	OP-28	m0	M-268	sc	M-195	wnb	M-112
hgd	M-267	meto	M-179	strc	M-264	wnd	M-116
ibulk	OP-22	mgd	M-269	stre	M-265	wnoi	M-235
id	OP-21	minr	M-190	stress	OP-26	wnom	M-250
ids	OP-5	mj	M-210	subc1	M-261	wphi	M-8
imax	M-203	mjsw	M-214	subc2	M-262	wref	M-158
imelt	M-200	mu0	M-26	sube	M-263	wu10	M-96
is	M-197	mu0b	M-30	subthmod	M-130	wulb	M-100
isdeff	O-10	mu20	M-42	temp	M-161	wuld	M-104
isseff	O-9	mu2b	M-46	tempmod	M-163	wua0	M-80
isub	OP-25	mu2g	M-50	tlev	M-167	wuab	M-84
jmax	M-204	mu30	M-54	tlevc	M-168	wub0	M-88
jmelt	M-201	mu3b	M-58	tox	M-217	wubb	M-92
js	M-196	mu3g	M-62	trd	M-174	wvfb	M-4
k1	M-10	mu40	M-66	trise	M-162	wvghigh	M-149
k1	O-16	mu4b	M-70	trise	I-13	wvglow	M-153
k2	M-14	mu4g	M-74	trs	M-173	wvof0	M-120
k2	O-17	mus0	M-34	type	M-1	wvofb	M-124
kf	M-232	musb	M-38	type	OP-1	wvofd	M-128
l	M-222	n	M-198	u10	M-94	x1	M-160
l	I-2	n0	M-106	ulb	M-98	xpart	M-181
lai0	M-132	nb	M-110	uld	M-102	xqc	M-182

Spectre Circuit Simulator Reference Component Statements Part I

laib	M-136	nd	M-114	ua0	M-78	xti	M-175
lbi0	M-140	noisemod	M-231	uab	M-82	xw	M-159
lbib	M-144	nrd	I-7	ub0	M-86		
lc0	M-279	nrd	M-227	ubb	M-90		

BSIM3 MOS Transistor (bsim3)

Description

This is the BSIM3 version-2 (BSIM3v2) model. The BSIM3v2 model is a physically-based, predictive, and computationally efficient model developed at the University of California, Berkeley. It is suitable for both digital and analog applications. In SPICE mode, refer to BSIM3 as MOS level 10. BSIM3 transistors require that you use a model statement.

This device is supported within altergroups.

Sample Instance Statement

```
m3 (1 2 0 0) nchmod l=1.5u w=100u as=450p ad=450p pd=209u ps=209u nrd=207m nrs=207m  
m=1
```

Sample Model Statement

```
model nchmod bsim3 vtho=5.94e-01 phi=0.69 k1=0.72 k2=0 w0=1.3e-07 tox=5.9e-09  
rdsw=80 uo=499 xj=2e-07 vsat=600e+04 at=3.4e+04 a0=0.8 cdsc=1.4e-03 nfactor=1.03
```

Instance Definition

```
Name d g s b ModelName parameter=value ...
```

Instance Parameters

- 1 w (m) Channel width.
- 2 l (m) Channel length.
- 3 as (m²) Area of source diffusion.
- 4 ad (m²) Area of drain diffusion.

Spectre Circuit Simulator Reference

Component Statements Part I

5	<code>ps (m)</code>	Perimeter of source diffusion.
6	<code>pd (m)</code>	Perimeter of drain diffusion.
7	<code>nrd (m/m)</code>	Number of squares of drain diffusion.
8	<code>nrs (m/m)</code>	Number of squares of source diffusion.
9	<code>m=1</code>	Multiplicity factor (number of MOSFETs in parallel).
10	<code>region=triode</code>	Estimated operating region. Spectre outputs number (0-4) in a rawfile. Possible values are <code>off</code> , <code>triode</code> , <code>sat</code> , <code>subth</code> , or <code>breakdown</code> .
11	<code>trise</code>	Temperature rise from ambient.

Model Definition

```
model modelName bsim3 parameter=value ...
```

Model Parameters

Device type parameters

1	<code>type=n</code>	Transistor type. Possible values are <code>n</code> or <code>p</code> .
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Threshold voltage parameters

2	<code>vtho=0 V</code>	Threshold voltage at zero body bias.
3	<code>phi=0.7 V</code>	Surface potential at strong inversion.
4	<code>k1=0.53 \sqrt{V}</code>	Body-effect coefficient.
5	<code>k2=-0.0186</code>	Charge-sharing parameter.
6	<code>k3=80</code>	Narrow width coefficient.
7	<code>k3b=0 1/V</code>	Narrow width coefficient.

Spectre Circuit Simulator Reference

Component Statements Part I

8	$w0=2.5e-6$ m	Narrow width coefficient.
9	$nlx=1.74e-7$ m	Lateral nonuniform doping coefficient.
10	$\gamma1=0$ \sqrt{V}	Body-effect coefficient near the surface.
11	$\gamma2=0$ \sqrt{V}	Body-effect coefficient in the bulk.
12	$\theta=0.02$ 1/V	Drain-induced barrier lowering coefficient.
13	$\eta=0.3$ 1/V	Effective drain voltage coefficient.
14	$lit1$ (m)	Depth of current path.
15	vfb (V)	Flat-band voltage.
16	vbx (V)	Threshold voltage transition body voltage.
17	vbi (V)	Substrate junction built-in potential.
18	$vbm=-5$ V	Maximum applied body voltage.
19	$dvt0=2.2$	First coefficient of short-channel effects.
20	$dvt1=0.53$	Second coefficient of short-channel effects.
21	$dvt2=-0.032$ 1/V	Body-bias coefficient of short-channel effects.
22	$a0=1$ for nmos and 4.4 for pmos	Nonuniform depletion width effect coefficient.
23	$a1=0$ for nmos, 0.23 for pmos	No-saturation coefficient.
24	$a2=1$ for nmos, 0.08 for pmos	No-saturation coefficient.
25	$keta=-0.047$ 1/V	Body-bias coefficient for non-uniform depletion width effect.

Process parameters

26	$n_{sub}=2e15$ cm ⁻³	Substrate doping concentration.
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Spectre Circuit Simulator Reference

Component Statements Part I

27	$n_{\text{peak}}=1.7e17 \text{ cm}^{-3}$	Peak channel doping concentration.
28	$n_{\text{gate}} \text{ (cm}^{-3}\text{)}$	Poly-gate doping concentration.
29	$x_j=0.15e-6 \text{ m}$	Source/drain junction depth.
30	$d_l=0 \text{ m}$	Lateral diffusion for one side.
31	$d_w=0 \text{ m}$	Width reduction for one side.
32	$t_{\text{ox}}=1.5e-8 \text{ m}$	Gate oxide thickness.
33	$v_{\text{dd}}=5 \text{ V}$	Maximum drain voltage.
34	$x_t=1.55e-7 \text{ m}$	Doping depth.
35	$l_{\text{dd}}=0 \text{ m}$	Total length of lightly doped drain region.
36	$r_{\text{ds0}}=0 \text{ } \Omega$	Total drain-source resistance.
37	$r_{\text{dsw}}=0 \text{ } \Omega \text{ } \mu\text{m}$	Width dependence of drain-source resistance.

Mobility parameters

38	$u_0=670 \text{ cm}^2/\text{V s}$	Low-field surface mobility at t_{nom} . Default is 250 for PMOS.
39	$v_{\text{sat}}=9.58e4 \text{ m/s}$	Carrier saturation velocity at t_{nom} .
40	$u_a=2.25e-9 \text{ m/v}$	First-order mobility reduction coefficient.
41	$u_b=5.87e-19 \text{ m}^2/\text{v}^2$	Second-order mobility reduction coefficient.
42	$u_c=0.0465 \text{ 1/V}$	Body-bias dependence of mobility.
43	$u_{c0}=0$	Mobility coefficient.

Output resistance parameters

44	$\text{satmod}=2$	Saturation model selector.
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Spectre Circuit Simulator Reference

Component Statements Part I

45	<code>bulkmod=1</code>	Bulk-charge effect model selector.
46	<code>drout=0.56</code>	DIBL effect on output resistance coefficient.
47	<code>alpha=1.9</code>	Reference voltage multiplication factor.
48	<code>em=4.1e7 V/m</code>	Maximum electric field.
49	<code>pclm=1.3</code>	Channel length modulation coefficient.
50	<code>pdibl1=0.39</code>	First coefficient of drain-induced barrier lowering.
51	<code>pdibl2=8.6e-3</code>	Second coefficient of drain-induced barrier lowering.
52	<code>pscbe1=4.24e8 V/m</code>	First coefficient of substrate current body effect.
53	<code>pscbe2=1e-5 m/v</code>	Second coefficient of substrate current body effect.
54	<code>pvag=0</code>	Gate dependence of Early voltage.

Subthreshold parameters

55	<code>subthmod=2</code>	Subthreshold model selector.
56	<code>vghigh=0.12 V</code>	Upper bound of transition region.
57	<code>vglow=-0.12 V</code>	Lower bound of transition region.
58	<code>cdsc=2.4e-4 F/m²</code>	Source/drain and channel coupling capacitance.
59	<code>cdscb=0 F/m² V</code>	Body-bias dependence of <code>cdsc</code> .
60	<code>nfactor=1</code>	Subthreshold swing coefficient.
61	<code>cit=0 F</code>	Interface trap parameter for subthreshold swing.
62	<code>voff=-0.11 V</code>	Threshold voltage offset.
63	<code>dsub=drout</code>	DIBL effect in subthreshold region.
64	<code>eta0=0.08</code>	DIBL coefficient subthreshold region.

Spectre Circuit Simulator Reference

Component Statements Part I

65 `etab=-0.07 1/V` Body-bias dependence of `et0`.

Parasitic resistance parameters

66 `rsh=0 Ω/sqr` Source/drain diffusion sheet resistance.

67 `rs=0 Ω` Source resistance.

68 `rd=0 Ω` Drain resistance.

69 `lgcs=0 m` Gate-to-contact length of source side.

70 `lgcd=0 m` Gate-to-contact length of drain side.

71 `rsc=0 Ω` Source contact resistance.

72 `rdc=0 Ω` Drain contact resistance.

73 `rss=0 Ω m` Scalable source resistance.

74 `rdd=0 Ω m` Scalable drain resistance.

75 `sc=∞ m` Spacing between contacts.

76 `ldif=0 m` Lateral diffusion beyond the gate.

77 `hdif=0 m` Length of heavily doped diffusion.

78 `minr=0.1 Ω` Minimum source/drain resistance.

Junction diode model parameters

79 `js (A/m2)` Bulk junction reverse saturation current density.

80 `is=1e-14 A` Bulk junction reverse saturation current.

81 `n=1` Junction emission coefficient.

82 `dskip=yes` Use simple piece-wise linear model for diode currents below $0.1 \cdot i_{abstol}$.
Possible values are `no` or `yes`.

Spectre Circuit Simulator Reference

Component Statements Part I

83 `imelt=`imaxA'` Explosion current.

84 `jmelt=`jmaxA/m'2`
Explosion current density.

Overlap capacitance parameters

85 `cgso (F/m)` Gate-source overlap capacitance.

86 `cgdo (F/m)` Gate-drain overlap capacitance.

87 `cgbo (F/m)` Gate-bulk overlap capacitance.

88 `meto=0 m` Metal overlap in fringing field.

Junction capacitance model parameters

89 `cbs=0 F` Bulk-source zero-bias junction capacitance.

90 `cbd=0 F` Bulk-drain zero-bias junction capacitance.

91 `cj=5e-4 F/m2` Zero-bias junction bottom capacitance density.

92 `mj=1/2` Bulk junction bottom grading coefficient.

93 `pb=0.8 V` Bulk junction built-in potential.

94 `fc=0.5` Forward-bias depletion capacitance threshold.

95 `cjsw=5e-10 F/m` Zero-bias junction sidewall capacitance density.

96 `mjsw=1/3` Bulk junction sidewall grading coefficient.

97 `pbsw=0.8 V` Side-wall junction built-in potential.

98 `fcsw=0.5` Side-wall forward-bias depletion capacitance threshold.

Spectre Circuit Simulator Reference

Component Statements Part I

Charge model selection parameters

- 99 `capmod=yang` Intrinsic charge model.
Possible values are `none`, `meyer`, `yang`, or `bsim`.
- 100 `xpart=1` Drain/source channel charge partition in saturation for BSIM charge model, use 0.0 for 40/60, 0.5 for 50/50, or 1.0 for 0/100.
- 101 `xqc=0` Drain/source channel charge partition in saturation for charge models, e.g. use 0.4 for 40/60, 0.5 for 50/50, 0 for 0/100.

Default instance parameters

- 102 `w=5e-6 m` Default channel width.
- 103 `l=5e-6 m` Default channel length.
- 104 `as=0 m2` Default area of source diffusion.
- 105 `ad=0 m2` Default area of drain diffusion.
- 106 `ps=0 m` Default perimeter of source diffusion.
- 107 `pd=0 m` Default perimeter of drain diffusion.
- 108 `nrd=0 m/m` Default number of squares of drain diffusion.
- 109 `nrs=0 m/m` Default number of squares of source diffusion.

Temperature effects parameters

- 110 `tnom (C)` Parameters measurement temperature. Default set by `options`.
- 111 `trise=0 C` Temperature rise from ambient.
- 112 `tlev=0` DC temperature selector.
- 113 `tlevc=0` AC temperature selector.
- 114 `eg=1.12452 V` Energy band gap.

Spectre Circuit Simulator Reference

Component Statements Part I

- 115 $\text{gap1}=7.02\text{e-}4 \text{ V/C}$ Band gap temperature coefficient.
- 116 $\text{gap2}=1108 \text{ C}$ Band gap temperature offset.
- 117 $\text{kt1}=-0.11 \text{ V}$ Temperature coefficient for threshold voltage.
- 118 $\text{kt11}=-1.86\text{e-}7 \text{ v m}$ Temperature coefficient for threshold voltage.
- 119 $\text{kt2}=0.022$ Temperature coefficient for threshold voltage.
- 120 $\text{at}=3.3\text{e}4 \text{ m/s}$ Temperature coefficient for v_{sat} .
- 121 $\text{ua1}=4.31\text{e-}9 \text{ m/v}$ Temperature coefficient for u_a .
- 122 $\text{ub1}=-7.61\text{e-}18 \text{ m}^2/\text{v}^2$
Temperature coefficient for u_b .
- 123 $\text{uc1}=-0.056 \text{ 1/V}$ Temperature coefficient for u_c .
- 124 $\text{trs}=0 \text{ 1/C}$ Temperature parameter for source resistance.
- 125 $\text{trd}=0 \text{ 1/C}$ Temperature parameter for drain resistance.
- 126 $\text{ute}=-1.5$ Mobility temperature exponent.
- 127 $\text{xti}=3$ Saturation current temperature exponent.
- 128 $\text{ptc}=0 \text{ V/C}$ Surface potential temperature coefficient.
- 129 $\text{tcv}=0 \text{ V/C}$ Threshold voltage temperature coefficient.
- 130 $\text{pta}=0 \text{ V/C}$ Junction potential temperature coefficient.
- 131 $\text{ptp}=0 \text{ V/C}$ Sidewall junction potential temperature coefficient.
- 132 $\text{cta}=0 \text{ 1/C}$ Junction capacitance temperature coefficient.
- 133 $\text{ctp}=0 \text{ 1/C}$ Sidewall junction capacitance temperature coefficient.

Noise model parameters

- 134 $\text{noisemod}=1$ Noise model selector.

Spectre Circuit Simulator Reference

Component Statements Part I

- 135 `kf=0` Flicker (1/f) noise coefficient.
- 136 `af=1` Flicker (1/f) noise exponent.
- 137 `ef=1` Flicker (1/f) noise frequency exponent.
- 138 `wnoi=1e-5 m` Channel width at which noise parameters were extracted.
- 139 `a=1e16` for nmos and `9.9e14` for pmos
Oxide trap density coefficient.
- 140 `b=5e4` for nmos and `2.4e3` for pmos
Oxide trap density coefficient.
- 141 `c=-1.4e-8` for nmos and `1.4e-8` for pmos
Oxide trap density coefficient.

Operating region warning control parameters

- 142 `alarm=none` Forbidden operating region.
Possible values are none, off, triode, sat, subth, or rev.
- 143 `imax=1 A` Maximum allowable current.
- 144 `jmax=1e8 A/m2` Maximum allowable current density.
- 145 `bvj=∞ V` Junction reverse breakdown voltage.
- 146 `vbox=1e9 tox V` Oxide breakdown voltage.
- 147 `maxvp=1.12 V` Maximum allowable voltage across the gate poly layer.

Auto Model Selector parameters

- 148 `wmax=1.0 m` Maximum channel width for which the model is valid.
- 149 `wmin=0.0 m` Minimum channel width for which the model is valid.
- 150 `lmax=1.0 m` Maximum channel length for which the model is valid.
- 151 `lmin=0.0 m` Minimum channel length for which the model is valid.

Spectre Circuit Simulator Reference

Component Statements Part I

Imax and Imelt

The `imax` parameter aids convergence and prevents numerical overflow. The junction characteristics of the device are accurately modeled for current up to `imax`. If `imax` is exceeded during iterations, the linear model is substituted until the current drops below `imax` or until convergence is achieved. If convergence is achieved with the current exceeding `imax`, the results are inaccurate, and Spectre prints a warning.

A separate model parameter, `imelt`, is used as a limit warning for the junction current. This parameter can be set to the maximum current rating of the device. When any component of the junction current exceeds `imelt`, note that base and collector currents are composed of many exponential terms, Spectre issues a warning and the results become inaccurate. The junction current is linearized above the value of `imelt` to prevent arithmetic exception, with the exponential term replaced by a linear equation at `imelt`.

Both of these parameters have current density counterparts, `jmax` and `jmelt`, that you can specify if you want the absolute current values to depend on the device area.

Auto Model Selection

Many models need to be characterized for different geometries in order to obtain accurate results for model development. The model selector program automatically searches for a model with the length and width range specified in the instance statement and uses this model in the simulations.

For the auto model selector program to find a specific model, the models to be searched should be grouped together within braces. Such a group is called a model group. An opening brace is required at the end of the line defining each model group. Every model in the group is given a name followed by a colon and the list of parameters. Also, the four geometric parameters `lmax`, `lmin`, `wmax`, and `wmin` should be given. The selection criteria to choose a model is as follows:

$$lmin \leq inst_length < lmax \text{ and } wmin \leq inst_width < wmax$$

Example:

```
model ModelName ModelType {  
  1: <model parameters> lmin=2 lmax=4 wmin=1 wmax=2  
  2: <model parameters> lmin=1 lmax=2 wmin=2 wmax=4  
  3: <model parameters> lmin=2 lmax=4 wmin=4 wmax=6
```


Spectre Circuit Simulator Reference

Component Statements Part I

}

Then for a given instance

```
M1 1 2 3 4 ModelName w=3 l=1.5
```

the program would search all the models in the model group with the name `ModelName` and then pick the first model whose geometric range satisfies the selection criteria. In the preceding example, the auto model selector program would choose `ModelName.2`.

You must specify both length (*l*) and width (*w*) on the device instance line to enable automatic model selection.

Output Parameters

- | | | |
|---|--|------------------------------|
| 1 | <code>w_{eff}</code> (m) | Effective channel width. |
| 2 | <code>l_{eff}</code> (m) | Effective channel length. |
| 3 | <code>r_{seff}</code> (Ω) | Effective source resistance. |
| 4 | <code>r_{deff}</code> (Ω) | Effective drain resistance. |

Operating-Point Parameters

- | | | |
|---|----------------------------|--|
| 1 | <code>type=n</code> | Transistor type.
Possible values are <code>n</code> or <code>p</code> . |
| 2 | <code>region=triode</code> | Estimated operating region. Spectre outputs number (0-4) in a rawfile.
Possible values are <code>off</code> , <code>triode</code> , <code>sat</code> , <code>subth</code> , or <code>breakdown</code> . |
| 3 | <code>reversed</code> | Reverse mode indicator.
Possible values are <code>no</code> or <code>yes</code> . |
| 4 | <code>ids</code> (A) | Resistive drain-to-source current. |
| 5 | <code>vgs</code> (V) | Gate-source voltage. |
| 6 | <code>vds</code> (V) | Drain-source voltage. |
| 7 | <code>vbs</code> (V) | Bulk-source voltage. |

Spectre Circuit Simulator Reference

Component Statements Part I

8	<code>vth</code> (V)	Threshold voltage.
9	<code>vdsat</code> (V)	Drain-source saturation voltage.
10	<code>gm</code> (S)	Common-source transconductance.
11	<code>gds</code> (S)	Common-source output conductance.
12	<code>gmbs</code> (S)	Body-transconductance.
13	<code>betaeff</code> (A/V ²)	Effective beta.
14	<code>cbd</code> (F)	Drain-bulk junction capacitance.
15	<code>cbs</code> (F)	Source-bulk junction capacitance.
16	<code>cgs</code> (F)	Gate-source capacitance.
17	<code>cgd</code> (F)	Gate-drain capacitance.
18	<code>cgb</code> (F)	Gate-bulk capacitance.
19	<code>ron</code> (Ω)	On-resistance.
20	<code>id</code> (A)	Resistive drain current.
21	<code>ibulk</code> (A)	Resistive bulk current.
22	<code>pwr</code> (W)	Power at op point.
23	<code>gmoverid</code> (1/V)	Gm/Ids.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

a M-139 etab M-65 ngate M-28 tnom M-110

Spectre Circuit Simulator Reference Component Statements Part I

a0	M-22	fc	M-94	nlx	M-9	tox	M-32
a1	M-23	fcswh	M-98	noisemod	M-134	trd	M-125
a2	M-24	gamma1	M-10	npeak	M-27	trise	I-11
ad	I-4	gamma2	M-11	nrd	I-7	trise	M-111
ad	M-105	gap1	M-115	nrd	M-108	trs	M-124
af	M-136	gap2	M-116	nrs	M-109	type	M-1
alarm	M-142	gds	OP-11	nrs	I-8	type	OP-1
alpha	M-47	gm	OP-10	nsub	M-26	ua	M-40
as	I-3	gmbs	OP-12	pb	M-93	ua1	M-121
as	M-104	gmoverid	OP-23	pbsw	M-97	ub	M-41
at	M-120	hdif	M-77	pclm	M-49	ubl	M-122
b	M-140	ibulk	OP-21	pd	M-107	uc	M-42
betaeff	OP-13	id	OP-20	pd	I-6	uc0	M-43
bulkmod	M-45	ids	OP-4	pdibl1	M-50	uc1	M-123
bvj	M-145	imax	M-143	pdibl2	M-51	uo	M-38
c	M-141	imelt	M-83	phi	M-3	ute	M-126
capmod	M-99	is	M-80	ps	I-5	vbi	M-17
cbd	OP-14	jmax	M-144	ps	M-106	vbm	M-18
cbd	M-90	jmelt	M-84	pscbe1	M-52	vbox	M-146
cbs	OP-15	js	M-79	pscbe2	M-53	vbs	OP-7
cbs	M-89	k1	M-4	pta	M-130	vbx	M-16
cdsc	M-58	k2	M-5	ptc	M-128	vdd	M-33
cdscb	M-59	k3	M-6	ptp	M-131	vds	OP-6
cgb	OP-18	k3b	M-7	pvag	M-54	vdsat	OP-9
cgbo	M-87	keta	M-25	pwr	OP-22	vfb	M-15
cgd	OP-17	kf	M-135	rd	M-68	vghigh	M-56
cgdo	M-86	kt1	M-117	rdc	M-72	vglow	M-57
cgs	OP-16	kt11	M-118	rdd	M-74	vgs	OP-5
cgso	M-85	kt2	M-119	rdeff	O-4	voff	M-62
cit	M-61	l	M-103	rds0	M-36	vsat	M-39
cj	M-91	l	I-2	rdsw	M-37	vth	OP-8
cjsw	M-95	ldd	M-35	region	I-10	vtho	M-2

Spectre Circuit Simulator Reference Component Statements Part I

cta	M-132	ldif	M-76	region	OP-2	w	I-1
ctp	M-133	leff	O-2	reversed	OP-3	w	M-102
d1	M-30	lgcd	M-70	ron	OP-19	w0	M-8
drou	M-46	lgcs	M-69	rs	M-67	weff	O-1
dskip	M-82	litl	M-14	rsc	M-71	wmax	M-148
dsub	M-63	lmax	M-150	rseff	O-3	wmin	M-149
dvt0	M-19	lmin	M-151	rsh	M-66	wnoi	M-138
dvt1	M-20	m	I-9	rss	M-73	xj	M-29
dvt2	M-21	maxvp	M-147	satmod	M-44	xpart	M-100
dw	M-31	meto	M-88	sc	M-75	xqc	M-101
ef	M-137	minr	M-78	subthmod	M-55	xt	M-34
eg	M-114	mj	M-92	tcv	M-129	xti	M-127
em	M-48	mjsw	M-96	theta	M-12		
eta	M-13	n	M-81	tlev	M-112		
eta0	M-64	nfactor	M-60	tlevc	M-113		

BSIM3v3 MOS Transistor (bsim3v3)

Description

BSIM3v3 is the version-3 of bsim3 model. The versions supported are 3.1, 3.2, 3.21, 3.22, 3.23 and 3.24. It uses single-piece equations for all regions to improve the smoothness of the model characteristics. BSIM3v3 also allows the binning option like the approach used in bsim1 and bsim2. This option is provided for people who want to achieve the highest accuracy of the model. The binning equation is given by

$$P = P_0 + P_I / L_{eff} + P_w / W_{eff} + P_p / (L_{eff} * W_{eff})$$

Only the P₀ parameters are listed. P_I, P_w, and P_p are not shown but can be recognized. The names of P_I, P_w, and P_p are identical to that of P₀ but with a prefix of I, w, and p, respectively. BSIM3v3 transistors require that you use a model statement.

For more information on this model, please consult the University of California at Berkeley BSIM3 home page at

<http://www-device.eecs.berkeley.edu/~bsim3/index.html>

Spectre Circuit Simulator Reference

Component Statements Part I

This device is supported within altergroups.

Sample Instance Statement

```
m4 (0 2 1 1) pchmod w=2u l=0.8u as=250p ad=250p pd=168p ps=168p m=1
```

Sample Model Statement

```
model pchmod bsim3v3 type=p mobmod=1 capmod=2 version=3.1 tox=9e-5 cdsc=1e-3  
cdscb=-4.36889e-4 cdscd=0 cit=0 nfactor=1.79 xj=1.5e-7 vsat=1.5737e5 at=1e5  
a0=1.2522809 ags=0.2912413 a1=1.01222e-4 a2=0.996841 keta=0 nch=4.06263e17  
ngate=7.6e19 kl=0.823562
```

Instance Definition

```
Name d g s b ModelName parameter=value ...
```

Instance Parameters

1	w (m)	Channel width.
2	l (m)	Channel length.
3	as (m ²)	Area of source diffusion.
4	ad (m ²)	Area of drain diffusion.
5	ps (m)	Perimeter of source diffusion.
6	pd (m)	Perimeter of drain diffusion.
7	nrd (m/m)	Number of squares of drain diffusion.
8	nrs (m/m)	Number of squares of source diffusion.
9	m=1	Multiplicity factor (number of MOSFETs in parallel).
10	region=triode	Estimated operating region. Spectre outputs number (0-4) in a rawfile. Possible values are off, triode, sat, subth, or breakdown.
11	nqsmod	NQS flag.
12	trise	Temperature rise from ambient.

Spectre Circuit Simulator Reference

Component Statements Part I

13	<code>aforward=0</code>	Forward gate leakage current coefficient.
14	<code>areverse=0</code>	Reverse gate leakage current coefficient.
15	<code>delvto=0 V</code>	shift in zero-bias threshold voltage <code>vth0</code> .
16	<code>mulmu0=1</code>	mobility multiplier.
17	<code>delk1=0 \sqrt{V}</code>	shift in body bias coefficient <code>k1</code> .
18	<code>delnfcct=0</code>	shift in subthreshold swing factor <code>nfactor</code> .
19	<code>geo=0</code>	Geometry selector.
20	<code>rdc=0</code>	Drain contact resistance.
21	<code>rsc=0</code>	Source contact resistance.
22	<code>sa=0 m</code>	Distance between OD edge to poly of one side.
23	<code>sb=0 m</code>	Distance between OD edge to poly of the other side.

Model Definition

```
model modelName bsim3v3 parameter=value ...
```

Model Parameters

Device type parameters

1	<code>type=n</code>	Transistor type. Possible values are <code>n</code> or <code>p</code> .
---	---------------------	--

Threshold voltage parameters

2	<code>vtho (V)</code>	Threshold voltage at zero body bias for long-channel devices. For enhancement-mode devices, <code>vtho</code> > 0 for n-channel and <code>vth</code> < 0 for p-channel. Default value is calculated from other model parameters.
3	<code>vfb=-1 V</code>	Flat-band voltage.

Spectre Circuit Simulator Reference

Component Statements Part I

4	$k1=0.5 \sqrt{V}$	Body-effect coefficient.
5	$k2=-0.0186$	Charge-sharing parameter.
6	$k3=80$	Narrow width coefficient.
7	$k3b=0 \text{ 1/V}$	Narrow width coefficient.
8	$w0=2.5e-6 \text{ m}$	Narrow width coefficient.
9	$n1x=1.74e-7 \text{ m}$	Lateral nonuniform doping coefficient.
10	$\text{gamma1} (\sqrt{V})$	Body-effect coefficient near the surface.
11	$\text{gamma2} (\sqrt{V})$	Body-effect coefficient in the bulk.
12	$vbx \text{ (V)}$	Threshold voltage transition body voltage.
13	$vbm=-3 \text{ V}$	Maximum applied body voltage.
14	$dvt0=2.2$	First coefficient of short-channel effects.
15	$dvt1=0.53$	Second coefficient of short-channel effects.
16	$dvt2=-0.032 \text{ 1/V}$	Body-bias coefficient of short-channel effects.
17	$dvt0w=0$	First coefficient of narrow-width effects.
18	$dvt1w=5.3e6$	Second coefficient of narrow-width effects.
19	$dvt2w=-0.032 \text{ 1/V}$	Body-bias coefficient of narrow-width effects.
20	$a0=1$	Nonuniform depletion width effect coefficient.
21	$b0=0 \text{ m}$	Bulk charge coefficient due to narrow width effect.
22	$b1=0 \text{ m}$	Bulk charge coefficient due to narrow width effect.
23	$a1=0$	No-saturation coefficient.
24	$a2=1$	No-saturation coefficient.
25	$ags=0 \text{ F/m}^2 \text{ V}$	Gate-bias dependence of A_{bulk} .

Spectre Circuit Simulator Reference

Component Statements Part I

- 26 $keta=-0.047$ $1/V$ Body-bias coefficient for non-uniform depletion width effect.
- 27 $vfbflag=0$ 49 Vfb selector.

Process parameters

- 28 $nsub=6e16$ cm^{-3} Substrate doping concentration.
- 29 $nch=1.7e17$ cm^{-3} Peak channel doping concentration.
- 30 $ngate=0$ cm^{-3} Poly-gate doping concentration.
- 31 $xj=0.15e-6$ m Source/drain junction depth.
- 32 $lint=0$ m Lateral diffusion for one side.
- 33 $wint=0$ m Width reduction for one side.
- 34 $ll=0$ m Length dependence of delta L.
- 35 $lln=1$ Length exponent of delta L.
- 36 $lw=0$ m Width dependence of delta L.
- 37 $lwn=1$ Width exponent of delta L.
- 38 $lwl=0$ m^2 Area dependence of delta L.
- 39 $wl=0$ m Length dependence of delta W.
- 40 $wln=1$ Length exponent of delta W.
- 41 $ww=0$ m Width dependence of delta W.
- 42 $wwn=1$ Width exponent of delta W.
- 43 $wwl=0$ m^2 Area dependence of delta W.
- 44 $dwg=0$ m/v Gate-bias dependence of channel width.
- 45 $dwb=0$ m/\sqrt{v} Body-bias dependence of channel width.

Spectre Circuit Simulator Reference

Component Statements Part I

46	$t_{ox}=1.5e-8$ m	Gate oxide thickness.
47	$dt_{oxcv}=0.0$ m	Delta oxide thickness.
48	$t_{oxm}=t_{ox}$ m	Tox at which parameters were extracted.
49	$x_t=1.55e-7$ m	Doping depth.
50	$r_{dsw}=0$ Ω μm	Width dependence of drain-source resistance.
51	$pr_{wb}=0$ $1/\sqrt{v}$	Body-effect coefficient for Rds.
52	$pr_{wg}=0$ $1/V$	Gate-effect coefficient for Rds.
53	$w_r=1$	Width offset for parasitic resistance.
54	$binunit=1$	Bin parameter unit selector. 1 for microns and 2 for meters.
55	$binflag=0$	49 binning factor.
56	$l_{ref}=1.0e20$	49 binning length factor.
57	$w_{ref}=1.0e20$	49 binning width factor.

Mobility parameters

58	$mobmod=1$	Mobility model selector.
59	$u_0=670$ $\text{cm}^2/\text{V s}$	Low-field surface mobility at t_{nom} . Default is 250 for PMOS Mobility can also be specified in M^2/Vs .
60	$vsat=8e4$ m/s	Carrier saturation velocity at t_{nom} .
61	$ua=2.25e-9$ m/v	First-order mobility reduction coefficient.
62	$ub=5.87e-19$ m^2/v^2	Second-order mobility reduction coefficient.
63	$uc=-4.65e-11$ m/v ²	Body-bias dependence of mobility. Default is -0.046 and unit is 1/ V for $mobmod=3$.

Spectre Circuit Simulator Reference

Component Statements Part I

Output resistance parameters

64	<code>drout=0.56</code>	DIBL effect on output resistance coefficient.
65	<code>pclm=1.3</code>	Channel length modulation coefficient.
66	<code>pdiblc1=0.39</code>	First coefficient of drain-induced barrier lowering.
67	<code>pdiblc2=8.6e-3</code>	Second coefficient of drain-induced barrier lowering.
68	<code>pdiblc1b=0 1/V</code>	Body-effect coefficient for DIBL.
69	<code>pscbe1=4.24e8 V/m</code>	First coefficient of substrate current body effect.
70	<code>pscbe2=1e-5 m/v</code>	Second coefficient of substrate current body effect.
71	<code>pvag=0</code>	Gate dependence of Early voltage.
72	<code>delta=0.01 V</code>	Effective drain voltage smoothing parameter.

Subthreshold parameters

73	<code>cdsc=2.4e-4 F/m²</code>	Source/drain and channel coupling capacitance.
74	<code>cdscb=0 F/m² V</code>	Body-bias dependence of <code>cdsc</code> .
75	<code>cdscd=0 F/m² V</code>	Drain-bias dependence of <code>cdsc</code> .
76	<code>nfactor=1</code>	Subthreshold swing coefficient.
77	<code>cit=0 F</code>	Interface trap parameter for subthreshold swing.
78	<code>voff=-0.08 V</code>	Threshold voltage offset.
79	<code>dsub=drout</code>	DIBL effect in subthreshold region.
80	<code>eta0=0.08</code>	DIBL coefficient subthreshold region.
81	<code>etab=-0.07 1/V</code>	Body-bias dependence of <code>et0</code> .

Spectre Circuit Simulator Reference

Component Statements Part I

Substrate current parameters

- 82 $\alpha_0=0$ m/v Substrate current impact ionization coefficient.
- 83 $\alpha_1=0$ 1/V Substrate current impact ionization coefficient.
- 84 $\beta_0=30$ 1/V Substrate current impact ionization exponent.

Parasitic resistance parameters

- 85 $r_{sh}=0$ Ω/sqr Source/drain diffusion sheet resistance.
- 86 $r_s=0$ Ω Source resistance.
- 87 $r_d=0$ Ω Drain resistance.
- 88 $lgcs=0$ m Gate-to-contact length of source side.
- 89 $lgcd=0$ m Gate-to-contact length of drain side.
- 90 $r_{sc}=0$ Ω Source contact resistance.
- 91 $r_{dc}=0$ Ω Drain contact resistance.
- 92 $r_{ss}=0$ Ω m Scalable source resistance.
- 93 $r_{dd}=0$ Ω m Scalable drain resistance.
- 94 $s_c=\infty$ m Spacing between contacts.
- 95 $ldif=0$ m Lateral diffusion beyond the gate.
- 96 $hdif=0$ m Length of heavily doped diffusion.
- 97 $minr=0.1$ Ω Minimum source/drain resistance.

Junction diode model parameters

- 98 j_s (A/m^2) Bulk junction reverse saturation current density.
- 99 $j_{sw}=0$ A/m Sidewall junction reverse saturation current density.

Spectre Circuit Simulator Reference

Component Statements Part I

100	<code>is=1e-14 A</code>	Bulk junction reverse saturation current.
101	<code>n=1</code>	Junction emission coefficient.
102	<code>dskip=yes</code>	Use simple piece-wise linear model for diode currents below $0.1 \cdot i_{abstol}$. Possible values are <code>no</code> or <code>yes</code> .
103	<code>imelt=`imaxA'</code>	Explosion current.
104	<code>ijth (A)</code>	Alias to <code>imelt</code> .
105	<code>jmelt=`jmaxA/m'²</code>	Explosion current density.
106	<code>vnds=-1</code>	Reverse diode current transition point.
107	<code>nds=1</code>	Reverse bias slope coefficient.
108	<code>tt=0.0 s</code>	Transit time.

Overlap capacitance parameters

109	<code>cgso (F/m)</code>	Gate-source overlap capacitance.
110	<code>cgdo (F/m)</code>	Gate-drain overlap capacitance.
111	<code>cgbo=2 Dwc Cox F/m</code>	Gate-bulk overlap capacitance. The default value is 0 if <code>version=3.0</code> .
112	<code>meto=0 m</code>	Metal overlap in fringing field.
113	<code>cgsl=0 F/m</code>	Gate-source overlap capacitance in LDD region.
114	<code>cgdl=0 F/m</code>	Gate-drain overlap capacitance in LDD region.
115	<code>ckappa=0.6</code>	Overlap capacitance fitting parameter.

Spectre Circuit Simulator Reference

Component Statements Part I

Junction capacitance model parameters

116	<code>cbs=0</code>	F	Bulk-source zero-bias junction capacitance.
117	<code>cbd=0</code>	F	Bulk-drain zero-bias junction capacitance.
118	<code>cj=5e-4</code>	F/m ²	Zero-bias junction bottom capacitance density.
119	<code>mj=1/2</code>		Bulk junction bottom grading coefficient.
120	<code>pb=1</code>	V	Bulk junction built-in potential.
121	<code>fc=0.5</code>		Forward-bias depletion capacitance threshold.
122	<code>cjsw=5e-10</code>	F/m	Zero-bias junction sidewall capacitance density.
123	<code>mjsw=0.33</code>		Bulk junction sidewall grading coefficient.
124	<code>pbsw=1</code>	V	Side-wall junction built-in potential.
125	<code>cjswg=cjsw</code>	F/m	Zero-bias gate-side junction capacitance density.
126	<code>mjswg=mjsw</code>		Gate-side junction grading coefficient.
127	<code>pbswg=pbsw</code>	V	Gate-side junction built-in potential.
128	<code>fcsw=0.5</code>		Side-wall forward-bias depletion capacitance threshold.

Charge model selection parameters

129	<code>capmod=2</code>		Intrinsic charge model.
130	<code>nqsmod=0</code>		Non-quasi static model selector. Set to 1 to turn on nqs.
131	<code>dwc=wint</code>	m	Delta W for capacitance model.
132	<code>dlc=lint</code>	m	Delta L for capacitance model.
133	<code>clc=1e-7</code>	m	Intrinsic capacitance fitting parameter.
134	<code>cle=0.6</code>		Intrinsic capacitance fitting parameter.

Spectre Circuit Simulator Reference

Component Statements Part I

135	c_f (F/m)	Fringe capacitance parameter.
136	$e_{lm}=5$	Elmore constant of the channel.
137	$v_{fbcv}=-1$	Flat-band voltage for $capmod=0$.
138	$acde=1$ 1/V	CV parameter.
139	$mo_{in}=15$ 1/V	CV parameter.
140	$n_{off}=1$	Transition parameter.
141	$v_{offcv}=0$	Transition parameter.
142	$x_{part}=0$	Drain/source channel charge partition in saturation for BSIM charge model, use 0.0 for 40/60, 0.5 for 50/50, or 1.0 for 0/100.
143	$llc=11$ m	Length dependence of delta L for CV.
144	$lwc=1w$ m	Width dependence of delta L for CV.
145	$lwl_c=lwl$ m ²	Area dependence of delta L for CV.
146	$wlc=w1$ m	Length dependence of delta W for CV.
147	$wwc=ww$ m	Width dependence of delta W for CV.
148	$wwl_c=wwl$ m ²	Area dependence of delta W for CV.
149	$wmlt=1.0$	Width shrink reduction factor.
150	$lm1t=1.0$	Length shrink reduction factor.

Default for instance parameters

151	$w=5e-6$ m	Default channel width.
152	$l=5e-6$ m	Default channel length.
153	$a_s=0$ m ²	Default area of source diffusion.
154	$a_d=0$ m ²	Default area of drain diffusion.

Spectre Circuit Simulator Reference

Component Statements Part I

155	<code>ps=0 m</code>	Default perimeter of source diffusion.
156	<code>pd=0 m</code>	Default perimeter of drain diffusion.
157	<code>nrd=0 m/m</code>	Default number of squares of drain diffusion.
158	<code>nrs=0 m/m</code>	Default number of squares of source diffusion.
159	<code>version=3.1</code>	Model version selector. The available versions are 3.1, 3.2, 3.21, 3.22, 3.23 and 3.24.
160	<code>paramchk=1</code>	Model parameter checking selector.
161	<code>fullreinit=0</code>	Model parameter full reinit selector.
162	<code>level=11</code>	BSIM3v3 model selector. The available level are 11, 49 and 53.
163	<code>acm=0</code>	BSIM3v3 area calculation method selector.
164	<code>geo=0</code>	geometry selector.
165	<code>calcacm=0</code>	49 geometry factor.

Temperature effects parameters

166	<code>tnom (C)</code>	Parameters measurement temperature. Default set by <code>options</code> .
167	<code>trise=0 C</code>	Temperature rise from ambient.
168	<code>tlev=0</code>	DC temperature selector.
169	<code>tlevc=0</code>	AC temperature selector.
170	<code>eg=1.12452 V</code>	Energy band gap.
171	<code>gap1=7.02e-4 V/C</code>	Band gap temperature coefficient.
172	<code>gap2=1108 C</code>	Band gap temperature offset.
173	<code>diomod=1</code>	a flag to select junction model. <code>diomod=1</code> junction model described in Common MOSFET Equations section will be used. <code>diomod=0</code> Berkeley junction model is used.

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174	$kt1=-0.11$	V	Temperature coefficient for threshold voltage.
175	$kt11=0$	v m	Temperature coefficient for threshold voltage.
176	$kt2=0.022$		Temperature coefficient for threshold voltage.
177	$at=3.3e4$	m/s	Temperature coefficient for v_{sat} .
178	$ua1=4.31e-9$	m/v	Temperature coefficient for u_a .
179	$ub1=-7.61e-18$	m^2/v^2	Temperature coefficient for u_b .
180	$uc1=-5.5e-11$	m/v^2	Temperature coefficient for u_c . Default is -0.056 for $mobmod=3$.
181	$prt=0$	Ω	Temperature coefficient for R_{ds} .
182	$trs=0$	1/C	Temperature parameter for source resistance.
183	$trd=0$	1/C	Temperature parameter for drain resistance.
184	$ute=-1.5$		Mobility temperature exponent.
185	$x_{ti}=3$		Saturation current temperature exponent.
186	$pta=0$	V/C	Junction potential temperature coefficient.
187	$tpb=0$	V/C	Temperature coefficient for p_b .
188	$ptp=0$	V/C	Sidewall junction potential temperature coefficient.
189	$tpbsw=0$	V/C	Temperature coefficient for p_{bsw} .
190	$tpbswg=0$	V/C	Temperature coefficient for p_{bswg} .
191	$cta=0$	1/C	Junction capacitance temperature coefficient.
192	$tcj=0$	1/C	Temperature coefficient for c_j .
193	$ctp=0$	1/C	Sidewall junction capacitance temperature coefficient.
194	$tcjsw=0$	1/C	Temperature coefficient for c_{jsw} .

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195 `tcjswg=0 1/C` Temperature coefficient for `cjswg`.

Noise model parameters

196 `noimod=1` Noise model selector.

197 `kf=0` Flicker (1/f) noise coefficient.

198 `af=1` Flicker (1/f) noise exponent.

199 `ef=1` Flicker (1/f) noise frequency exponent.

200 `noia=1e20` Oxide trap density coefficient. Default is 9.9e18 for pmos.

201 `noib=5e4` Oxide trap density coefficient. Default is 2.4e3 for pmos.

202 `noic=-1.4e-12` Oxide trap density coefficient. Default is 1.4e-8 for pmos.

203 `noid=2e14` flicker noise subthreshold-above threshold transition coefficient.

204 `wnoi=1e-5 m` Channel width at which noise parameters were extracted.

205 `em=4.1e7 V/m` Maximum electric field.

206 `flkmod=0` Flicker noise model (0 for `Ids` based model, 1 for gm based model).

207 `gamma=2.0/3.0` Thermal noise coefficient.

208 `nlev=2.0` 49 noise selector.

Gate leak current parameters

209 `bforward=0` Forward gate leakage current coefficient in `pow()`.

210 `breverse=0` Reverse gate leakage current coefficient in `pow()`.

211 `cforward=0` Forward gate leakage current coefficient in `exp()`.

212 `creverse=0` Reverse gate leakage current coefficient in `exp()`.

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213 `tcc=0` Gate leakage current temperature coefficient.

Auto Model Selector parameters

214 `wmax=1 m` Maximum channel width for which the model is valid.

215 `wmin=0 m` Minimum channel width for which the model is valid.

216 `lmax=1 m` Maximum channel length for which the model is valid.

217 `lmin=0 m` Minimum channel length for which the model is valid.

Operating region warning control parameters

218 `alarm=none` Forbidden operating region.
Possible values are `none`, `off`, `triode`, `sat`, `subth`, or `rev`.

219 `imax=1 A` Maximum allowable current.

220 `jmax=1e8 A/m2` Maximum allowable current density.

221 `bvj= ∞ V` Junction reverse breakdown voltage.

222 `vbox=1e9 tox V` Oxide breakdown voltage.

223 `warn=on` Parameter to turn warnings on and off.
Possible values are `off` or `on`.

224 `apwarn=0` Warning message flag.

Length dependent parameters

225 `x1=0 m` Length variation due to masking and etching.

Width dependent parameters

226 `xw=0 m` Width variation due to masking and etching.

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Cross-term dependent parameters (Not listed)

DC-mismatch dependent parameters

- 227 `mvtwl=0.0` v m Threshold mismatch area dependence.
- 228 `mvtwl2=0.0` v m^{1.5} Threshold mismatch area square dependence.
- 229 `mvto=0.0` V Threshold mismatch intercept.
- 230 `mbewl=0.0` m Beta mismatch area dependence.
- 231 `mbeo=0.0` Beta mismatch intercept.

Mos Table Model parameters

- 232 `mos_method=a` Table model enable.
Possible values are `s` or `a`.

LOD model parameters

- 233 `sa0=1e-6` m reference distance between od edge to poly of one side.
- 234 `sb0=1e-6` m reference distance between od edge to poly of the other side.
- 235 `wlod=0` m length parameter for stress effect.
- 236 `ku0=0` m mobility degradation/enhancement coefficient for stress effect.
- 237 `kvsat=0` saturation velocity degradation/enhancement parameter for stress effect.
- 238 `kvth0=0` v m threshold shift parameter for stress effect.
- 239 `tku0=0` temperature coefficient of `ku0`.
- 240 `llodku0=0` length parameter for `u0` stress effect.
- 241 `wlodku0=0` width parameter for `u0` stress effect.

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- 242 `llodvth=0` length parameter for vth stress effect.
- 243 `wlodvth=0` width parameter for vth stress effect.
- 244 `lku0=0 m^llodku0` length dependence of ku0.
- 245 `wku0=0 m^wloдку0` width dependence of ku0.
- 246 `pku0=0 m^(llodku0+wloдку0)`
cross-term dependence of ku0.
- 247 `lkvth0=0 v m^llodku0`
length dependence of kvth0.
- 248 `wkvth0=0 v m^wloдку0`
width dependence of kvth0.
- 249 `pkvth0=0 v m^(llodku0+wloдку0)`
cross-term dependence of kvth0.
- 250 `stk2=0 m` k2 shift factor related to vth0 change.
- 251 `lodk2=0` k2 shift modification factor for stress effect.
- 252 `steta0=0 m` eta0 shift factor related to vth0 change.
- 253 `lodeta0=0` eta0 shift modification factor for stress effect.

Imax and Imelt

The `imax` parameter aids convergence and prevents numerical overflow. The junction characteristics of the device are accurately modeled for current up to `imax`. If `imax` is exceeded during iterations, the linear model is substituted until the current drops below `imax` or until convergence is achieved. If convergence is achieved with the current exceeding `imax`, the results are inaccurate, and Spectre prints a warning.

A separate model parameter, `imelt`, is used as a limit warning for the junction current. This parameter can be set to the maximum current rating of the device. When any component of the junction current exceeds `imelt`, note that base and collector currents are composed of many exponential terms, Spectre issues a warning and the results become inaccurate. The junction current is linearized above the value of `imelt` to prevent arithmetic exception, with the exponential term replaced by a linear equation at `imelt`.

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Both of these parameters have current density counterparts, j_{max} and j_{melt} , that you can specify if you want the absolute current values to depend on the device area.

Auto Model Selection

Many models need to be characterized for different geometries in order to obtain accurate results for model development. The model selector program automatically searches for a model with the length and width range specified in the instance statement and uses this model in the simulations.

For the auto model selector program to find a specific model, the models to be searched should be grouped together within braces. Such a group is called a model group. An opening brace is required at the end of the line defining each model group. Every model in the group is given a name followed by a colon and the list of parameters. Also, the four geometric parameters l_{max} , l_{min} , w_{max} , and w_{min} should be given. The selection criteria to choose a model is as follows:

$$l_{min} \leq inst_length < l_{max} \text{ and } w_{min} \leq inst_width < w_{max}$$

Example:

```
model ModelName ModelType {  
    1: <model parameters> lmin=2 lmax=4 wmin=1 wmax=2  
    2: <model parameters> lmin=1 lmax=2 wmin=2 wmax=4  
    3: <model parameters> lmin=2 lmax=4 wmin=4 wmax=6  
}
```

Then for a given instance

```
M1 1 2 3 4 ModelName w=3 l=1.5
```

the program would search all the models in the model group with the name `ModelName` and then pick the first model whose geometric range satisfies the selection criteria. In the preceding example, the auto model selector program would choose `ModelName.2`.

You must specify both length (l) and width (w) on the device instance line to enable automatic model selection.

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Output Parameters

- | | | |
|---|--|------------------------------|
| 1 | <code>w_{eff}</code> (m) | Effective channel width. |
| 2 | <code>l_{eff}</code> (m) | Effective channel length. |
| 3 | <code>r_{seff}</code> (Ω) | Effective source resistance. |
| 4 | <code>r_{deff}</code> (Ω) | Effective drain resistance. |

Operating-Point Parameters

- | | | |
|----|--|--|
| 1 | <code>type=n</code> | Transistor type.
Possible values are <code>n</code> or <code>p</code> . |
| 2 | <code>region=triode</code> | Estimated operating region. Spectre outputs number (0-4) in a rawfile.
Possible values are <code>off</code> , <code>triode</code> , <code>sat</code> , <code>subth</code> , or <code>breakdown</code> . |
| 3 | <code>reversed</code> | Reverse mode indicator.
Possible values are <code>no</code> or <code>yes</code> . |
| 4 | <code>ids</code> (A) | Resistive drain-to-source current. |
| 5 | <code>vgs</code> (V) | Gate-source voltage. |
| 6 | <code>vds</code> (V) | Drain-source voltage. |
| 7 | <code>vbs</code> (V) | Bulk-source voltage. |
| 8 | <code>vth</code> (V) | Threshold voltage. |
| 9 | <code>vdsat</code> (V) | Drain-source saturation voltage. |
| 10 | <code>gm</code> (S) | Common-source transconductance. |
| 11 | <code>gds</code> (S) | Common-source output conductance. |
| 12 | <code>gmbs</code> (S) | Body-transconductance. |
| 13 | <code>betaeff</code> (A/V ²) | Effective beta. |
| 14 | <code>cjd</code> (F) | Drain-bulk junction capacitance. |

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15	<code>cjs</code> (F)	Source-bulk junction capacitance.
16	<code>cgg</code> (F)	dQg_dVg .
17	<code>cgd</code> (F)	dQg_dVd .
18	<code>cgs</code> (F)	dQg_dVs .
19	<code>cgb</code> (F)	dQg_dVbk .
20	<code>cdg</code> (F)	dQd_dVg .
21	<code>cdd</code> (F)	dQd_dVd .
22	<code>cds</code> (F)	dQd_dVs .
23	<code>cdb</code> (F)	dQd_dVb .
24	<code>csg</code> (F)	dQs_dVg .
25	<code>csd</code> (F)	dQs_dVd .
26	<code>css</code> (F)	dQs_dVs .
27	<code>csb</code> (F)	dQs_dVb .
28	<code>cbg</code> (F)	dQb_dVg .
29	<code>cbd</code> (F)	dQb_dVd .
30	<code>cbs</code> (F)	dQb_dVs .
31	<code>cbb</code> (F)	dQb_dVb .
32	<code>ron</code> (Ω)	On-resistance.
33	<code>id</code> (A)	Resistive drain current.
34	<code>is</code> (A)	Resistive source current.
35	<code>ibulk</code> (A)	Resistive bulk current.
36	<code>pwr</code> (W)	Power at op point.

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37 gmoverid (1/V) Gm/lds.

Parameter Index

In the following index, I refers to instance parameters, M refers to the model parameters section, O refers to the output parameters section, and OP refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of M-35 means the 35th model parameter.

a0	M-20	dvt0w	M-17	m	I-9	stk2	M-250
a1	M-23	dvt1	M-15	mbe0	M-231	tcc	M-213
a2	M-24	dvt1w	M-18	mbewl	M-230	tcj	M-192
acde	M-138	dvt2	M-16	meto	M-112	tcjsw	M-194
acm	M-163	dvt2w	M-19	minr	M-97	tcjswg	M-195
ad	I-4	dwb	M-45	mj	M-119	tku0	M-239
ad	M-154	dwc	M-131	mjsw	M-123	tlev	M-168
af	M-198	dwg	M-44	mjswg	M-126	tlevc	M-169
aforward	I-13	ef	M-199	mobmod	M-58	tnom	M-166
ags	M-25	eg	M-170	moin	M-139	tox	M-46
alarm	M-218	elm	M-136	mos_method	M-232	toxm	M-48
alpha0	M-82	em	M-205	mulmu0	I-16	tpb	M-187
alpha1	M-83	eta0	M-80	mvt0	M-229	tpbsw	M-189
apwarn	M-224	etab	M-81	mvtw1	M-227	tpbswg	M-190
areverse	I-14	fc	M-121	mvtw12	M-228	trd	M-183
as	M-153	fcsw	M-128	n	M-101	trise	M-167
as	I-3	flkmod	M-206	nch	M-29	trise	I-12
at	M-177	fullreinit	M-161	nds	M-107	trs	M-182
b0	M-21	gamma	M-207	nfactor	M-76	tt	M-108
b1	M-22	gamma1	M-10	ngate	M-30	type	OP-1
beta0	M-84	gamma2	M-11	nlev	M-208	type	M-1
betaeff	OP-13	gap1	M-171	nlx	M-9	u0	M-59
bforward	M-209	gap2	M-172	noff	M-140	ua	M-61

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binflag	M-55	gds	OP-11	noia	M-200	ual	M-178
binunit	M-54	geo	I-19	noib	M-201	ub	M-62
breverse	M-210	geo	M-164	noic	M-202	ubl	M-179
bvj	M-221	gm	OP-10	noid	M-203	uc	M-63
calcacm	M-165	gmbs	OP-12	noimod	M-196	uc1	M-180
capmod	M-129	gmoverid	OP-37	nqsmod	M-130	ute	M-184
cbb	OP-31	hdif	M-96	nqsmod	I-11	vbm	M-13
cbd	M-117	ibulk	OP-35	nrd	I-7	vbox	M-222
cbd	OP-29	id	OP-33	nrd	M-157	vbs	OP-7
cbg	OP-28	ids	OP-4	nrs	I-8	vbx	M-12
cbs	OP-30	ijth	M-104	nrs	M-158	vds	OP-6
cbs	M-116	imax	M-219	nsub	M-28	vdsat	OP-9
cdb	OP-23	imelt	M-103	paramchk	M-160	version	M-159
cdd	OP-21	is	OP-34	pb	M-120	vfb	M-3
cdg	OP-20	is	M-100	pbsw	M-124	vfbcv	M-137
cds	OP-22	jmax	M-220	pbswg	M-127	vfbflag	M-27
cdsc	M-73	jmelt	M-105	pclm	M-65	vgs	OP-5
cdscb	M-74	js	M-98	pd	M-156	vnds	M-106
cdscd	M-75	jsw	M-99	pd	I-6	voff	M-78
cf	M-135	k1	M-4	pdiblc1	M-66	voffcv	M-141
cforward	M-211	k2	M-5	pdiblc2	M-67	vsat	M-60
cgb	OP-19	k3	M-6	pdiblc3	M-68	vth	OP-8
cgbo	M-111	k3b	M-7	pku0	M-246	vtho	M-2
cgd	OP-17	keta	M-26	pkvth0	M-249	w	M-151
cgdl	M-114	kf	M-197	prt	M-181	w	I-1
cgdo	M-110	kt1	M-174	prwb	M-51	w0	M-8
cgg	OP-16	kt11	M-175	prwg	M-52	warn	M-223
cgs	OP-18	kt2	M-176	ps	I-5	weff	O-1
cgs1	M-113	ku0	M-236	ps	M-155	wint	M-33
cgso	M-109	kvsat	M-237	pscbe1	M-69	wku0	M-245
cit	M-77	kvth0	M-238	pscbe2	M-70	wkvth0	M-248
cj	M-118	l	I-2	pta	M-186	wl	M-39

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cjd	OP-14	l	M-152	ptp	M-188	wlc	M-146
cjs	OP-15	ldif	M-95	pvag	M-71	wln	M-40
cjsw	M-122	leff	O-2	pwr	OP-36	wlod	M-235
cjswg	M-125	level	M-162	rd	M-87	wlodku0	M-241
ckappa	M-115	lgcd	M-89	rdc	M-91	wlodvth	M-243
clc	M-133	lgcs	M-88	rdc	I-20	wmax	M-214
cle	M-134	lint	M-32	rdd	M-93	wmin	M-215
creverse	M-212	lku0	M-244	rdeff	O-4	wmlt	M-149
csb	OP-27	lkvth0	M-247	rdsw	M-50	wnoi	M-204
csd	OP-25	ll	M-34	region	I-10	wr	M-53
csg	OP-24	llc	M-143	region	OP-2	wref	M-57
css	OP-26	lln	M-35	reversed	OP-3	ww	M-41
cta	M-191	llodku0	M-240	ron	OP-32	wwc	M-147
ctp	M-193	llodvth	M-242	rs	M-86	wwl	M-43
delk1	I-17	lmax	M-216	rsc	M-90	wwlc	M-148
delnfct	I-18	lmin	M-217	rsc	I-21	wwn	M-42
delta	M-72	lmlt	M-150	rseff	O-3	xj	M-31
delvto	I-15	lodeta0	M-253	rsh	M-85	xl	M-225
diomod	M-173	lodk2	M-251	rss	M-92	xpart	M-142
dlc	M-132	lref	M-56	sa	I-22	xt	M-49
drout	M-64	lw	M-36	sa0	M-233	xti	M-185
dskip	M-102	lwc	M-144	sb	I-23	xw	M-226
dsub	M-79	lwl	M-38	sb0	M-234		
dtoxcv	M-47	lwlc	M-145	sc	M-94		
dvt0	M-14	lwn	M-37	steta0	M-252		

BSIM4 MOS Transistor (bsim4)

Description

BSIM4 is the version-4.21 of bsim model. It uses single-piece equations for all regions to improve the smoothness of the model characteristics. BSIM4 also allows the binning option

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like the approach used in `bsim3`. This option is provided for people who want to achieve the highest accuracy of the model. The binning equation is given by

$$P = P_0 + P_I / L_{eff} + P_w / W_{eff} + P_p / (L_{eff} * W_{eff})$$

Only the `P0` parameters are listed. `PI`, `Pw`, and `Pp` are not shown but can be recognized. The names of `PI`, `Pw`, and `Pp` are identical to that of `P0` but with a prefix of `I`, `w`, and `p`, respectively. BSIM4 transistors require that you use a model statement.

For more information on this model, please consult the University of California at Berkeley BSIM4 home page at

<http://www-device.eecs.berkeley.edu/~bsim3/bsim4.html>

This device is supported within `altergroups`.

Sample Instance Statement

```
m4 (0 2 1 1) pchmod w=2u l=0.8u as=250p ad=250p pd=168p ps=168p m=1
```

Sample Model Statement

```
model pchmod bsim4 type=p mobmod=0 capmod=2 version=4.21 tox=3e-9 cdsc=2.58e-4
cdscb=0 cdscd=6.1e-8 cit=0 nfactor=1.1 xj=9e-8 vfb=0.76vsat=9.2e4 at=3.3e4 a0=1.1
ags=1.0e-20 al=0 ngate=9e19 vth0=-0.42a1=0 a2=1 delta=0.014 pvag=1e-20 pclm=6.28e-
4 pdits=0.2 pditsl=2.3e6pditsd=0.23 fprout=0.2 pdiblcb=3.4e-8 pdiblc1=0.81
drout=0.56pdiblc2=9.84e-6 psobel=8.14e8 psobe2=9.58e-07 lint=5e-9 wint=5e-
9dmcg=5e-6 dmci=5e-6 dmdg=5e-6 dmcgt=6e-7 dwj=4.5e-8 rsh=6cgso=7.43e-10
cgdo=7.43e-10 cgbo=2.56e-11 cgsl=1e-14 cgdl=1e-14ckappas=0.5 ckappad=0.5 noff=0.9
voffcv=0.02 acde=1 moin=15 xpart=0ktl1=0 kt2=2.2e-2 lpe0=5.75e-8 lpeb=2.3e-10
dvt0=2.89 dvt1=0.53dvt2=-3.2e-2 dvt0w=0 dvt1w=0 dvt2w=0 dvtp0=7.32e-7
dvtp1=0.12dsub=0.058 eta0=0.001 u0=4.19e-2 ua=8.7e-16 ub=3.06e-18 k1=0.33uc=4.6e-
13 ute=-1.5 ual=4.31e-9 ub1=7.61e-18 uc1=-5.6e-11 k2=-1.87e-2rds=369.4 rdw=184.7
rsw=184.7 prwg=3.22e-8 prwb=6.8e-11 wr=1rdsmin=0 rdwmin=0 rswmin=0 prt=0 b0=-1e-
20 k3=80 k3b=0 w0=2.5e-6b1=0 keta=-0.047 alpha0=7.4e-2 alpha1=0.005 beta0=30
```

Instance Definition

```
Name d g s b ModelName parameter=value ...
```

Instance Parameters

- | | | |
|---|----------------------|---------------------------|
| 1 | w (m) | Channel width. |
| 2 | l (m) | Channel length. |
| 3 | as (m ²) | Area of source diffusion. |

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4	<code>ad</code> (m^2)	Area of drain diffusion.
5	<code>ps</code> (m)	Perimeter of source diffusion.
6	<code>pd</code> (m)	Perimeter of drain diffusion.
7	<code>nrd</code> (m/m)	Number of squares of drain diffusion.
8	<code>nrs</code> (m/m)	Number of squares of source diffusion.
9	<code>m=1</code>	Multiplicity factor (number of MOSFETs in parallel).
10	<code>region=triode</code>	Estimated operating region. Spectre outputs number (0-4) in a rawfile. Possible values are <code>off</code> , <code>triode</code> , <code>sat</code> , <code>subth</code> , or <code>breakdown</code> .
11	<code>trnqsmo</code>	Transient NQS flag.
12	<code>acnqsmo</code>	AC NQS flag.
13	<code>trise</code>	Temperature rise from ambient.
14	<code>rgatemod</code>	Rgate flag.
15	<code>rbodymod</code>	Rbody flag.
16	<code>geomod</code>	Geometry flag.
17	<code>rgeomod</code>	Diffusion resistance and contact model flag.
18	<code>rbpb</code> (Ω)	Resistance connected between bNode and bNode .
19	<code>rbpd</code> (Ω)	Resistance connected between bNode and dbNode.
20	<code>rbps</code> (Ω)	Resistance connected between bNode and sbNode.
21	<code>rbdb</code> (Ω)	Resistance connected between dbNode and bNode.
22	<code>rbbs</code> (Ω)	Resistance connected between sbNode and bNode.
23	<code>nf</code>	Number of device fingers.
24	<code>min</code>	Minimum number of device fingers.

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- 25 `delvto=0 V` shift in zero-bias threshold voltage `vth0`.
- 26 `mulmu0=1` mobility multiplier.
- 27 `delk1=0 \sqrt{V}` shift in body bias coefficient `k1`.
- 28 `delnfct=0` shift in subthreshold swing factor `nfactor`.

LOD model parameters

- 29 `sa=0` Distance between OD edge to poly of one side.
- 30 `sb=0` Distance between OD edge to poly of the other side.
- 31 `sd=0` Distance between neighbour fingers.

Model Definition

```
model modelName bsim4 parameter=value ...
```

Model Parameters

Device type parameters

- 1 `type=n` Transistor type.
Possible values are `n` or `p`.

Threshold voltage parameters

- 2 `vtho (V)` Threshold voltage at zero body bias for long-channel devices.
For enhancement-mode devices, `vtho > 0` for n-channel and `vth < 0` for p-channel. Default value is calculated from other model parameters.
- 3 `vfb=-1 V` Flat-band voltage.
- 4 `phin=0 V` Non-uniform vertical doping effect on surface potential.
- 5 `k1=0.53 \sqrt{V}` Body-effect coefficient.

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6	$k2 = -0.0186$	Charge-sharing parameter.
7	$k3 = 80$	Narrow width coefficient.
8	$k3b = 0 \text{ 1/V}$	Narrow width coefficient.
9	$w0 = 2.5e-6 \text{ m}$	Narrow width coefficient.
10	$lpe0 = 1.74e-7 \text{ m}$	Lateral non-uniform doping at $V_{bs} = 0$.
11	$lpeb = 0$	Lateral non-uniform doping effect on K1.
12	$\gamma1 \text{ (}\sqrt{V}\text{)}$	Body-effect coefficient near the surface.
13	$\gamma2 \text{ (}\sqrt{V}\text{)}$	Body-effect coefficient in the bulk.
14	$vbx \text{ (V)}$	Threshold voltage transition body voltage.
15	$vbm = -3 \text{ V}$	Maximum applied body voltage.
16	$dvt0 = 2.2$	First coefficient of short-channel effects.
17	$dvt1 = 0.53$	Second coefficient of short-channel effects.
18	$dvt2 = -0.032 \text{ 1/V}$	Body-bias coefficient of short-channel effects.
19	$dvtp0 = 0 \text{ m}$	First Coef. of drain-induced V_{th} shift for long-channel pocket devices.
20	$dvtp1 = 0 \text{ 1/V}$	Second Coef. of drain-induced V_{th} shift for long-channel pocket devices.
21	$dvt0w = 0$	First coefficient of narrow-width effects.
22	$dvt1w = 5.3e6 \text{ 1/m}$	Second coefficient of narrow-width effects.
23	$dvt2w = -0.032 \text{ 1/V}$	Body-bias coefficient of narrow-width effects.
24	$a0 = 1$	Nonuniform depletion width effect coefficient.
25	$b0 = 0 \text{ m}$	Bulk charge coefficient due to narrow width effect.
26	$b1 = 0 \text{ m}$	Bulk charge coefficient due to narrow width effect.

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27	a1=0	No-saturation coefficient.
28	a2=1	No-saturation coefficient.
29	ags=0 F/m ² V	Gate-bias dependence of Abulk.
30	keta=-0.047 1/V	Body-bias coefficient for non-uniform depletion width effect.

Process parameters

31	epsrox=3.9	Gate dielectric constant.
32	toxe=3.0e-9 m	Electrical gate oxide thickness.
33	toxp=toxe m	Electrical gate oxide thickness.
34	dtox=0.0 m	Difference between electrical and physical gate oxide thickness.
35	ndep=1.7e17 cm ⁻³	Channel doping concentration.
36	nsd=1.0e20 cm ⁻³	Source-drain doping concentration.
37	nsub=6e16 cm ⁻³	Substrate doping concentration.
38	ngate=0 cm ⁻³	Poly-gate doping concentration.
39	xj=0.15e-6 m	Source/drain junction depth.
40	lint=0 m	Lateral diffusion for one side.
41	wint=0 m	Width reduction for one side.
42	ll=0	Length dependence of delta L.
43	lln=1	Length exponent of delta L.
44	lw=0	Width dependence of delta L.
45	lwn=1	Width exponent of delta L.
46	lw1=0	Area dependence of delta L.

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47	<code>wl=0</code>	Length dependence of delta W.
48	<code>wln=1</code>	Length exponent of delta W.
49	<code>ww=0</code>	Width dependence of delta W.
50	<code>wwn=1</code>	Width exponent of delta W.
51	<code>wwl=0</code>	Area dependence of delta W.
52	<code>dwg=0 m/v</code>	Gate-bias dependence of channel width.
53	<code>dwb=0 m/\sqrt{v}</code>	Body-bias dependence of channel width.
54	<code>toxm=toxe m</code>	Toxe at which parameters were extracted.
55	<code>xt=1.55e-7 m</code>	Doping depth.
56	<code>binunit=1</code>	Bin parameter unit selector. 1 for microns and 2 for meters.

Bias-dependent Rds parameters

57	<code>rdsmode=1</code>	bias-dependent D/S model selector.
58	<code>rdsw=200</code>	Zero bias LDD resistance per unit width for RDSMOD=0..
59	<code>rdswmin=0</code>	LDD resistance per unit width at high Vgs and zero Vbs for RDSMOD=0.
60	<code>rdw=100</code>	Zero bias LDD resistance per unit width for RDSMOD=1..
61	<code>rdwmin=0</code>	LDD resistance per unit width at high Vgs and zero Vbs for RDSMOD=1.
62	<code>rsw=100</code>	Zero bias LDD resistance per unit width for RDSMOD=1..
63	<code>rswmin=0</code>	LDD resistance per unit width at high Vgs and zero Vbs for RDSMOD=1.
64	<code>prwb=0 1/\sqrt{v}</code>	Body-effect coefficient for Rds.
65	<code>prwg=1 1/V</code>	Gate-effect coefficient for Rds.

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66 `wr=1` Width offset for parasitic resistance.

Mobility parameters

67 `mobmod=0` Mobility model selector.

68 `u0=670 cm2/V s` Low-field surface mobility at t_{nom} . Default is 250 for PMOS.

69 `vsat=8e4 m/s` Carrier saturation velocity at t_{nom} .

70 `ua=1.0e-9 m/v` First-order mobility reduction coefficient. Default is 1.0e-15 if Mobmod = 2.

71 `ub=1.0e-19 m2/v2` Second-order mobility reduction coefficient.

72 `uc=-4.65e-11 m/v2` Body-bias dependence of mobility. Default is -0.0465 and unit is 1/V for mobmod=1.

73 `eu=1.67` Exponent for mobility degradation of mobmod=2. Default is 1.0 for Pmos.

Output resistance parameters

74 `drout=0.56` DIBL effect on output resistance coefficient.

75 `fprout=0.0 V/ \sqrt{m}` Effect of pocket implant on R_{out} degradation.

76 `pclm=1.3` Channel length modulation coefficient.

77 `pdiblc1=0.39` First coefficient of drain-induced barrier lowering.

78 `pdiblc2=8.6e-3` Second coefficient of drain-induced barrier lowering.

79 `pdiblcb=0 1/V` Body-effect coefficient for DIBL.

80 `pscbe1=4.24e8 V/m` First coefficient of substrate current body effect.

81 `pscbe2=1e-5 m/v` Second coefficient of substrate current body effect.

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82	<code>pvag=0</code>	Gate dependence of Early voltage.
83	<code>delta=0.01 V</code>	Effective drain voltage smoothing parameter.
84	<code>pdits=0.0 1/V</code>	Effect of pocket implant on Rout degradation.
85	<code>pditsl=0.0 1/m</code>	Channel-length of drain-induced Vth shift on Rout.
86	<code>pditsd=0.0 1/V</code>	Channel-length of drain-induced Vth shift on Rout.

Subthreshold parameters

87	<code>cdsc=2.4e-4 F/m²</code>	Source/drain and channel coupling capacitance.
88	<code>cdscb=0 F/m² V</code>	Body-bias dependence of <code>cdsc</code> .
89	<code>cdscd=0 F/m² V</code>	Drain-bias dependence of <code>cdsc</code> .
90	<code>nfactor=1</code>	Subthreshold swing coefficient.
91	<code>cit=0 F/m²</code>	Interface trap parameter for subthreshold swing.
92	<code>voff=-0.08 V</code>	Threshold voltage offset.
93	<code>voffl=0.0 V</code>	Channel-length dependence of Voff..
94	<code>minv=0</code>	Vgsteff fitting parameter for moderate inversion condition..
95	<code>dsub=drout</code>	DIBL effect in subthreshold region.
96	<code>eta0=0.08</code>	DIBL coefficient subthreshold region.
97	<code>etab=-0.07 1/V</code>	Body-bias dependence of <code>et0</code> .

Substrate current parameters

98	<code>alpha0=0 m/v</code>	Substrate current impact ionization coefficient.
99	<code>alpha1=0 1/V</code>	Substrate current impact ionization coefficient.
100	<code>beta0=30 1/V</code>	Substrate current impact ionization exponent.

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Parasitic resistance parameters

101	<code>rgatemod</code>	Rgate flag.
102	<code>rsh=0 Ω/sqr</code>	Source/drain diffusion sheet resistance.
103	<code>rshg=0.1 Ω/sqr</code>	Gate electrode diffusion sheet resistance.
104	<code>dmcg=0 m</code>	Distance from S/D contact center to the gate edge.
105	<code>dmci=dmcg m</code>	Distance from S/D contact center to the isolation edge in the channel-length direction.
106	<code>dmdg=0 m</code>	Distance from S/D contact center to the gate edge.
107	<code>dmcgt=0 m</code>	DMCG of test structures.
108	<code>dwj=Dwc</code>	Offset of the S/D junction width.
109	<code>xgw=0 m</code>	Distance from the gate contact to the channel edge.
110	<code>xgl=0 m</code>	Offset of the gate length due to variations in patterning.
111	<code>ngcon=1</code>	Number of gate contacts.
112	<code>nf=1</code>	Number of device fingers.
113	<code>min</code>	Minimum number of device fingers.
114	<code>permod=1</code>	Perimeter model selector.
115	<code>geomod</code>	Geometry flag.
116	<code>rgeomod</code>	Diffusion resistance and contact model flag.
117	<code>xw=0 m</code>	Width variation due to masking and etching.
118	<code>xl=0 m</code>	Length variation due to masking and etching.
119	<code>minr=0.001 Ω</code>	Minimum source/drain resistance.

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Gate-Induced drain leakage parameters

- 120 `agidl=0` $1/\Omega$ Pre-exponential coefficient for GIDL.
- 121 `bgidl=2.3e9` V Exponential coefficient for GIDL.
- 122 `cgidl=0.5` V^3 Exponential coefficient for GIDL.
- 123 `egidl=0.8` V Fitting parameter for band bending for GIDL.

Gate Tunneling parameters

- 124 `igcmod=0` Gate-to-channel tunneling model selector.
- 125 `igbmod=1` Gate-to-substrate tunneling model selector.
- 126 `aigbacc=0.43` $\sqrt{F/g}$ s/m
Parameter for I_{gb} in accumulation.
- 127 `bigbacc=0.054` $\sqrt{F/g}$ /sm
Parameter for I_{gb} in accumulation.
- 128 `cigbacc=0.075` $1/V$ Parameter for I_{gb} in accumulation.
- 129 `nigbacc=1` Parameter for I_{gb} in accumulation.
- 130 `aigbinv=0.35` $\sqrt{F/g}$ s/m
Parameter for I_{gb} in inversion.
- 131 `bigbinv=0.03` $\sqrt{F/g}$ /sm
Parameter for I_{gb} in inversion.
- 132 `cigbinv=0.006` $1/V$ Parameter for I_{gb} in inversion.
- 133 `eigbinv=1.1` V Parameter for I_{gb} in inversion.
- 134 `nigbinv=3` Parameter for I_{gb} in inversion.
- 135 `aigc=0.054` $\sqrt{F/g}$ s/m
Parameter for I_{gcs} and I_{gcd} . Default value for Pmos is 0.31.

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- 136 $\text{bigc}=0.054 \sqrt{F/g} / \text{sm}$ Parameter for Igcs and Igcd. Default value for Pmos is 0.024.
- 137 $\text{cigc}=0.075 1/V$ Parameter for Igcs and Igcd. Default value for Pmos is 0.03.
- 138 $\text{aigsd}=0.43 \sqrt{F/g} \text{ s/m}$ Parameter for Igs and Igd. Default value for Pmos is 0.31.
- 139 $\text{bigsd}=0.054 \sqrt{F/g} / \text{sm}$ Parameter for Igs and Igd. Default value for Pmos is 0.024.
- 140 $\text{cigsd}=0.075 1/V$ Parameter for Igs and Igd. Default value for Pmos is 0.03.
- 141 $\text{dlcig}=\text{Lint m}$ Source/drain overlap length for Igs and Igd.
- 142 $\text{nigc}=1.0$ Source/drain overlap length for Igs and Igd.
- 143 $\text{poxedge}=1.0$ Factor for the gate oxide thickness in source/drain overlap regions.
- 144 $\text{pigcd}=1.0$ Vds dependence of Igcs and Igcd.
- 145 $\text{ntox}=1.0$ Exponent for the gate oxide ratio.
- 146 $\text{toxref}=3.0\text{e-}9 \text{ m}$ Nominal gate oxide thickness for gate dielectric tunneling current model only.

Junction diode model parameters

- 147 $\text{diomod}=1$ Diode model selector.
- 148 $\text{js}=1.0\text{e-}4 \text{ A/m}^2$ Bottom junction reverse saturation current density..
- 149 $\text{jss}=1.0\text{e-}4 \text{ A/m}^2$ Bottom junction reverse saturation current density..
- 150 $\text{jsd}=\text{Jss} \text{ A/m}^2$ Bottom junction reverse saturation current density..
- 151 $\text{jsws}=0 \text{ A/m}$ Isolation-edge sidewall reverse saturation current density..
- 152 $\text{jswd}=\text{Jsws} \text{ A/m}$ Isolation-edge sidewall reverse saturation current density..
- 153 $\text{jswgs}=0 \text{ A/m}$ Gate-edge sidewall reverse saturation current density..

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154	<code>jswgd=Jswgs</code>	A/m	Gate-edge sidewall reverse saturation current density.
155	<code>is=1e-14</code>	A	Bulk junction reverse saturation current.
156	<code>n=1</code>		Junction emission coefficient.
157	<code>njs=1</code>		Bulk-Source junction emission coefficient.
158	<code>njd=Njs</code>		Bulk-Source junction emission coefficient.
159	<code>dskip=yes</code>		Use simple piece-wise linear model for diode currents below $0.1 \cdot i_{abstol}$. Possible values are <code>no</code> or <code>yes</code> .
160	<code>imelt='imaxA'</code>		Explosion current.
161	<code>jmelt='jmaxA/m'²</code>		Explosion current density.
162	<code>ijthsrev=0.1</code>	A	Limiting current in reverse bias region.
163	<code>ijthdrev=Ijthsrev</code>	A	Limiting current in reverse bias region.
164	<code>ijthsfwd=0.1</code>	A	Limiting current in forward bias region.
165	<code>ijthdfwd=Ijthsfwd</code>	A	Limiting current in forward bias region.
166	<code>xjbvs=1.0</code>		Limiting current in forward bias region.
167	<code>xjbvd=xjbvs</code>		Limiting current in forward bias region.

Overlap capacitance parameters

168	<code>cgso</code>	(F/m)	Non LDD region source-gate overlap capacitance per unit channel width.
169	<code>cgdo</code>	(F/m)	Non LDD region drain-gate overlap capacitance per unit channel width.

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170	$cgbo=2$	Dwc	Coxe	F/m	Non LDD region drain-gate overlap capacitance per unit channel width..
171	$meto=0$	m			Metal overlap in fringing field.
172	$cgs1=0$	F/m			Overlap capacitance between gate and lightly-doped source region.
173	$cgd1=Cgd1$	F/m			Overlap capacitance between gate and lightly-doped drain region.
174	$ckappas=0.6$	V			Coefficient of bias-dependent overlap capacitance for the source side.
175	$ckappad=Ckappas$	V			Coefficient of bias-dependent overlap capacitance for the source side.

Junction capacitance model parameters

176	$cj=5e-4$	F/m ²			Zero-bias junction bottom capacitance density.
177	$cjs=5e-4$	F/m ²			Zero bias bottom junction capacitance per unit area..
178	$cjd=5e-4$	F/m ²			Zero bias bottom junction capacitance per unit area..
179	$mj=1/2$				Bulk junction bottom grading coefficient.
180	$mjs=1/2$				Bulk junction bottom grading coefficient.
181	$mjd=Mjs$				Bulk junction bottom grading coefficient.
182	$pb=1$	V			Bottom junction built-in potential.
183	$pbs=1$	V			Bottom junction built-in potential.
184	$pbd=pbs$	V			Bottom junction built-in potential.
185	$fc=0.5$				Forward-bias depletion capacitance threshold.
186	$cjsw=5e-10$	F/m			Zero-bias junction sidewall capacitance density.

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187	$c_{jsws}=5e-10$ F/m	Zero-bias junction sidewall capacitance density.
188	$c_{jswd}=C_{jsws}$ F/m	Zero-bias junction sidewall capacitance density.
189	$m_{jsw}=0.33$	Isolation-edge sidewall junction capacitance grading coefficient..
190	$m_{jsws}=0.33$	Isolation-edge sidewall junction capacitance grading coefficient..
191	$m_{jswd}=M_{jsws}$	Isolation-edge sidewall junction capacitance grading coefficient..
192	$p_{bsw}=1$ V	Isolation-edge sidewall junction built-in potential.
193	$p_{bsws}=1$ V	Isolation-edge sidewall junction built-in potential.
194	$p_{bswd}=p_{bsws}$ V	Isolation-edge sidewall junction built-in potential.
195	c_{jswg} (F/m)	Zero-bias gate-side junction capacitance density.
196	$c_{jswgs}=c_{jsws}$ F/m	Zero-bias gate-side junction capacitance density.
197	$c_{jswgd}=c_{jswgs}$ F/m	Zero-bias gate-side junction capacitance density.
198	$m_{jswg}=m_{jsw}$	Gate-edge sidewall junction grading coefficient.
199	$m_{jswgs}=m_{jsws}$	Gate-edge sidewall junction grading coefficient.
200	$m_{jswgd}=m_{jsws}$	Gate-edge sidewall junction grading coefficient.
201	$p_{bswg}=p_{bsw}$ V	Gate-edge sidewall junction built-in potential.
202	$p_{bswgs}=p_{bsws}$ V	Gate-edge sidewall junction built-in potential.
203	$p_{bswgd}=p_{bsws}$ V	Gate-edge sidewall junction built-in potential.
204	$f_{csw}=0.5$	Side-wall forward-bias depletion capacitance threshold.
205	$bvs=10.0$ V	Breakdown voltage.
206	$bvd=Bvs$ V	Breakdown voltage.

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Charge model selection parameters

207	capmod=2	Intrinsic charge model.
208	trnqsmod=0	Transient Non-quasi static model selector. Set to 1 to turn on nqs.
209	acnqsmod=0	Ac Non-quasi static model selector. Set to 1 to turn on nqs.
210	dwc=wint m	Delta W for capacitance model.
211	dlc=Lint m	Delta L for capacitance model.
212	clc=1e-7 m	Constant term for the short channel model..
213	cle=0.6	Intrinsic capacitance fitting parameter.
214	cf (F/m)	Coefficient of bias-dependent overlap capacitance for the source side.
215	vfbcv=-1	Flat-band voltage for capmod=0.
216	acde=1 m/v	Exponential coefficient for charge thickness in CAPMOD=2 for accumulation and depletion regions.
217	moim=15 1/V	Exponential coefficient for charge thickness for accumulation and depletion regions.
218	noff=1	Transition parameter.
219	voffcv=0 V	CV parameter in VgsteffCV for weak to strong inversion..
220	xpart=0	Charge partition number. Use 0.0 for 40/60, 0.5 for 50/50, or 1.0 for 0/100.
221	llc=ll	Length dependence of delta L for CV.
222	lwc=lw	Width dependence of delta L for CV.
223	lwlc=lwl	Area dependence of delta L for CV.
224	wlc=wl	Length dependence of delta W for CV.
225	wwc=ww	Width dependence of delta W for CV.

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226 `wwlc=wwl` Area dependence of delta W for CV.

Default for instance parameters

227 `w=5e-6 m` Default channel width.

228 `l=5e-6 m` Default channel length.

229 `as=0 m2` Default area of source diffusion.

230 `ad=0 m2` Default area of drain diffusion.

231 `ps=0 m` Default perimeter of source diffusion.

232 `pd=0 m` Default perimeter of drain diffusion.

233 `nrd=0 m/m` Default number of squares of drain diffusion.

234 `nrs=0 m/m` Default number of squares of source diffusion.

235 `version=4.21` Model version selector.

236 `level=14` Model level selector for spice compatibility.

237 `paramchk=1` Model parameter checking selector.

238 `fullreinit=0` Model parameter full reinit selector.

Temperature effects parameters

239 `tnom (C)` Parameters measurement temperature. Default set by `options`.

240 `trise=0 C` Temperature rise from ambient.

241 `tlev=0` DC temperature selector.

242 `tlevc=0` AC temperature selector.

243 `eg=1.12452 V` Energy band gap.

244 `gap1=7.02e-4 V/C` Band gap temperature coefficient.

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245	gap2=1108 C	Band gap temperature offset.
246	kt1=-0.11 V	Temperature coefficient for threshold voltage.
247	kt11=0 v m	Temperature coefficient for threshold voltage.
248	kt2=0.022	Temperature coefficient for threshold voltage.
249	at=3.3e4 m/s	Temperature coefficient for v_{sat} .
250	ua1=4.31e-9 m/v	Temperature coefficient for u_a .
251	ub1=-7.61e-18 m ² /v ²	Temperature coefficient for u_b .
252	uc1=-5.5e-11 m/v ²	Temperature coefficient for u_c . Default is -0.056 for mobmod=3.
253	prt=0 Ω m	Temperature coefficient for R_{ds} .
254	ute=-1.5	Mobility temperature exponent.
255	xti=3	Saturation current temperature exponent.
256	xtis=3	Bulk-Source junction saturation current temperature exponent.
257	xtid=3	Bulk-Source junction saturation current temperature exponent.
258	pta=0 V/C	Temperature coefficient for p_b .
259	tpb=0 V/C	Temperature coefficient for p_b .
260	ptp=0 V/C	Temperature coefficient for $p_{b_{sw}}$.
261	tpbsw=0 V/C	Temperature coefficient for $p_{b_{sw}}$.
262	tpbswg=0 V/C	Temperature coefficient for $p_{b_{swg}}$.
263	cta=0 1/C	Temperature coefficient for c_j .
264	tcj=0 1/C	Temperature coefficient for c_j .
265	ctp=0 1/C	Temperature coefficient for $c_{j_{sw}}$.

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266 $t_{cjsw}=0$ 1/C Temperature coefficient for c_{jsw} .

267 $t_{cjswg}=0$ 1/C Temperature coefficient for c_{jswg} .

LOD model parameters

268 $s_{aref}=1e-6$ m reference distance between od edge to poly of one side.

269 $s_{bref}=1e-6$ m reference distance between od edge to poly of the other side.

270 $w_{lod}=0$ m length parameter for stress effect.

271 $k_{u0}=0$ m mobility degradation/enhancement coefficient for stress effect.

272 $k_{vsat}=0$ saturation velocity degradation/enhancement parameter for stress effect.

273 $k_{vth0}=0$ v m threshold shift parameter for stress effect.

274 $t_{ku0}=0$ temperature coefficient of k_{u0} .

275 $l_{lodku0}=0$ length parameter for u_0 stress effect.

276 $w_{lodku0}=0$ width parameter for u_0 stress effect.

277 $l_{lodvth0}=0$ length parameter for v_{th} stress effect.

278 $w_{lodvth0}=0$ width parameter for v_{th} stress effect.

279 $l_{ku0}=0$ m ^{l_{lodku0}} length dependence of k_{u0} .

280 $w_{ku0}=0$ m ^{w_{lodku0}} width dependence of k_{u0} .

281 $p_{ku0}=0$ m ^{$(l_{lodku0}+w_{lodku0})$} cross-term dependence of k_{u0} .

282 $l_{kvth0}=0$ v m ^{l_{lodku0}} length dependence of k_{vth0} .

283 $w_{kvth0}=0$ v m ^{w_{lodku0}} width dependence of k_{vth0} .

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284	<code>pkvth0=0</code>	<code>v m^(1lodku0+w1odku0)</code> cross-term dependence of kvth0.
285	<code>stk2=0</code>	<code>m</code> k2 shift factor related to vth0 change.
286	<code>lodk2=0</code>	k2 shift modification factor for stress effect.
287	<code>steta0=0</code>	<code>m</code> eta0 shift factor related to vth0 change.
288	<code>lodeta0=0</code>	eta0 shift modification factor for stress effect.

Noise model parameters

289	<code>fnoimod=1</code>	Flicker noise model selector.
290	<code>tnoimod=0</code>	Thermal noise model selector.
291	<code>kf=0</code>	Flicker noise exponent.
292	<code>af=1</code>	Flicker noise exponent.
293	<code>ef=1</code>	Flicker noise frequency exponent.
294	<code>noia=6.25e41</code>	Flicker noise parameter B. Default is 6.188e40 for pmos.
295	<code>noib=3.125e26</code>	Flicker noise parameter C. Default is 1.5e25 for pmos.
296	<code>noic=8.75e9</code>	Flicker noise parameter C.
297	<code>wnoi=1e-5</code>	<code>m</code> Channel width at which noise parameters were extracted.
298	<code>em=4.1e7</code>	<code>V/m</code> Saturation field.
299	<code>flkmod</code>	Flicker Noise Model.
300	<code>ntnoi=1</code>	Noise factor for short-channel devices for TNOIMOD=0 only.
301	<code>tnoia=1.5</code>	Coefficient of channel-length dependence of total channel thermal noise.
302	<code>tnoib=3.5</code>	Coefficient of channel-length dependence of total channel thermal noise.

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Substrate Network parameters

303	<code>rbodymod</code>	Rbody flag.
304	<code>xrcrg1=12</code>	Parameter for distributed channel-resistance effect for both intrinsic-input resistance and charge-deficit NQS models.
305	<code>xrcrg2=1</code>	Parameter to account for the excess channel diffusion resistance for both intrinsic-input resistance and charge-deficit NQS models.
306	<code>rbpb (Ω)</code>	Resistance connected between bNode and bNode .
307	<code>rbpd (Ω)</code>	Resistance connected between bNode and dbNode.
308	<code>rbps (Ω)</code>	Resistance connected between bNode and sbNode.
309	<code>rbdb (Ω)</code>	Resistance connected between dbNode and bNode.
310	<code>rbsb (Ω)</code>	Resistance connected between sbNode and bNode.
311	<code>gbmin=1.0e-12 1/Ω</code>	Conductance in parallel with each of the five substrate resistances to avoid potential numerical instability due to unreasonably too large a substrate resistance.

Auto Model Selector parameters

312	<code>wmax=1 m</code>	Maximum channel width for which the model is valid.
313	<code>wmin=0 m</code>	Minimum channel width for which the model is valid.
314	<code>lmax=1 m</code>	Maximum channel length for which the model is valid.
315	<code>lmin=0 m</code>	Minimum channel length for which the model is valid.

Operating region warning control parameters

316	<code>alarm=none</code>	Forbidden operating region. Possible values are <code>none</code> , <code>off</code> , <code>triode</code> , <code>sat</code> , <code>subth</code> , or <code>rev</code> .
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317	<code>imax=1</code>	A	Maximum allowable current.
318	<code>jmax=1e8</code>	A/m ²	Maximum allowable current density.
319	<code>bvj=∞</code>	V	Junction reverse breakdown voltage.
320	<code>vbox=1e9</code>	<code>toxe</code> V	Oxide breakdown voltage.
321	<code>warn=on</code>		Parameter to turn warnings on and off. Possible values are <code>off</code> or <code>on</code> .

Length dependent parameters (Not listed)

Width dependent parameters (Not listed)

Cross-term dependent parameters (Not listed)

DC-mismatch dependent parameters

322	<code>mvtwl=0.0</code>	v m	Threshold mismatch area dependence.
323	<code>mvtwl2=0.0</code>	v m ^{1.5}	Threshold mismatch area square dependence.
324	<code>mvt0=0.0</code>	V	Threshold mismatch intercept.
325	<code>mbewl=0.0</code>	m	Beta mismatch area dependence.
326	<code>mbe0=0.0</code>		Beta mismatch intercept.

Mos Table Model parameters

327	<code>mos_method=a</code>		Table model enable. Possible values are <code>s</code> or <code>a</code>
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Spectre Circuit Simulator Reference

Component Statements Part I

Imax and Imelt

The `imax` parameter aids convergence and prevents numerical overflow. The junction characteristics of the device are accurately modeled for current up to `imax`. If `imax` is exceeded during iterations, the linear model is substituted until the current drops below `imax` or until convergence is achieved. If convergence is achieved with the current exceeding `imax`, the results are inaccurate, and Spectre prints a warning.

A separate model parameter, `imelt`, is used as a limit warning for the junction current. This parameter can be set to the maximum current rating of the device. When any component of the junction current exceeds `imelt`, note that base and collector currents are composed of many exponential terms, Spectre issues a warning and the results become inaccurate. The junction current is linearized above the value of `imelt` to prevent arithmetic exception, with the exponential term replaced by a linear equation at `imelt`.

Both of these parameters have current density counterparts, `jmax` and `jmelt`, that you can specify if you want the absolute current values to depend on the device area.

Auto Model Selection

Many models need to be characterized for different geometries in order to obtain accurate results for model development. The model selector program automatically searches for a model with the length and width range specified in the instance statement and uses this model in the simulations.

For the auto model selector program to find a specific model, the models to be searched should be grouped together within braces. Such a group is called a model group. An opening brace is required at the end of the line defining each model group. Every model in the group is given a name followed by a colon and the list of parameters. Also, the four geometric parameters `lmax`, `lmin`, `wmax`, and `wmin` should be given. The selection criteria to choose a model is as follows:

$$lmin \leq inst_length < lmax \text{ and } wmin \leq inst_width < wmax$$

Example:

```
model ModelName ModelType {  
  1: <model parameters> lmin=2 lmax=4 wmin=1 wmax=2  
  2: <model parameters> lmin=1 lmax=2 wmin=2 wmax=4  
  3: <model parameters> lmin=2 lmax=4 wmin=4 wmax=6
```


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}

Then for a given instance

```
M1 1 2 3 4 ModelName w=3 l=1.5
```

the program would search all the models in the model group with the name `ModelName` and then pick the first model whose geometric range satisfies the selection criteria. In the preceding example, the auto model selector program would choose `ModelName.2`.

You must specify both length (`l`) and width (`w`) on the device instance line to enable automatic model selection.

Output Parameters

1	<code>w_{eff}</code> (m)	Effective channel width.
2	<code>l_{eff}</code> (m)	Effective channel length.
3	<code>rg_{bi}</code> (Ω)	Gate bias-independent resistance.
4	<code>v_{tho}</code> (V)	V _{tho} model defined or calculated.
5	<code>k₁</code> (\sqrt{V})	K1 model defined or calculated.
6	<code>k₂</code>	K2 model defined or calculated.

Operating-Point Parameters

1	<code>type=n</code>	Transistor type. Possible values are <code>n</code> or <code>p</code> .
2	<code>region=triode</code>	Estimated operating region. Spectre outputs number (0-4) in a rawfile. Possible values are <code>off</code> , <code>triode</code> , <code>sat</code> , <code>subth</code> , or <code>breakdown</code> .
3	<code>reversed</code>	Reverse mode indicator. Possible values are <code>no</code> or <code>yes</code> .
4	<code>ids</code> (A)	Resistive drain-to-source current.
5	<code>vgs</code> (V)	Gate-source voltage.

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6	vds (V)	Drain-source voltage.
7	vbs (V)	Bulk-source voltage.
8	vth (V)	Threshold voltage.
9	vdsat (V)	Drain-source saturation voltage.
10	gm (S)	Common-source transconductance.
11	gds (S)	Common-source output conductance.
12	gmbs (S)	Body-transconductance.
13	betaeff (A/V ²)	Effective beta.
14	cjd (F)	Drain-bulk junction capacitance.
15	cjs (F)	Source-bulk junction capacitance.
16	cgg (F)	dQg_dVg.
17	cgd (F)	dQg_dVd.
18	cgs (F)	dQg_dVs.
19	cgb (F)	dQg_dVbk.
20	cdg (F)	dQd_dVg.
21	cdd (F)	dQd_dVd.
22	cds (F)	dQd_dVs.
23	cdb (F)	dQd_dVb.
24	csg (F)	dQs_dVg.
25	csd (F)	dQs_dVd.
26	css (F)	dQs_dVs.
27	csb (F)	dQs_dVb.

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28	cbg (F)	dQb_dVg.
29	cbd (F)	dQb_dVd.
30	cbs (F)	dQb_dVs.
31	cbb (F)	dQb_dVb.
32	ron (Ω)	On-resistance.
33	id (A)	Resistive drain current.
34	ibulk (A)	Resistive bulk current.
35	pwr (W)	Power at op point.
36	gmoverid (1/V)	Gm/Ids.
37	rdeff (Ω)	Effective drain resistance.
38	rseff (Ω)	Effective source resistance.
39	rgbd (Ω)	Gate bias-dependent resistance.
40	igidl (A)	Gate-induced drain leakage current.
41	igd (A)	Gate-to-drain tunneling current.
42	igs (A)	Gate-to-source tunneling current.
43	igb (A)	Gate-to-bulk tunneling current.
44	igcs (A)	Gate-to-channel (source side) tunneling current.
45	igcd (A)	Gate-to-channel (drain side) tunneling current.
46	gbs (S)	bulk-source diode conductance.
47	gbd (S)	bulk-drain diode conductance.

Spectre Circuit Simulator Reference

Component Statements Part I

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

a0	M-24	dvt2w	M-23	min	I-24	rgbd	OP-39
a1	M-27	dvtp0	M-19	min	M-113	rgbi	O-3
a2	M-28	dvtp1	M-20	minr	M-119	rgeomod	I-17
acde	M-216	dwb	M-53	minv	M-94	rgeomod	M-116
acnqsmod	M-209	dwc	M-210	mj	M-179	ron	OP-32
acnqsmod	I-12	dwg	M-52	mjd	M-181	rseff	OP-38
ad	I-4	dwj	M-108	mjs	M-180	rsh	M-102
ad	M-230	ef	M-293	mjsw	M-189	rshg	M-103
af	M-292	eg	M-243	mjswd	M-191	rsw	M-62
agidl	M-120	egidl	M-123	mjswg	M-198	rswmin	M-63
ags	M-29	eigbinv	M-133	mjswgd	M-200	sa	I-29
aigbacc	M-126	em	M-298	mjswgs	M-199	saref	M-268
aigbinv	M-130	epsrox	M-31	mjsws	M-190	sb	I-30
aigc	M-135	eta0	M-96	mobmod	M-67	sbref	M-269
aigsd	M-138	etab	M-97	moin	M-217	sd	I-31
alarm	M-316	eu	M-73	mos_method	M-327	steta0	M-287
alpha0	M-98	fc	M-185	mulmu0	I-26	stk2	M-285
alpha1	M-99	fcsw	M-204	mvt0	M-324	tcj	M-264
as	M-229	flkmod	M-299	mvtw1	M-322	tcjsw	M-266
as	I-3	fnoimod	M-289	mvtw12	M-323	tcjswg	M-267
at	M-249	fprout	M-75	n	M-156	tku0	M-274
b0	M-25	fullreinit	M-238	ndep	M-35	tlev	M-241
b1	M-26	gamma1	M-12	nf	I-23	tlevc	M-242
beta0	M-100	gamma2	M-13	nf	M-112	tnoia	M-301
betaeff	OP-13	gap1	M-244	nfactor	M-90	tnoib	M-302

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bigbacc M-127	gbd OP-47	ngcon M-111	tnom M-239
bigbinv M-131	gbmin M-311	nigbacc M-129	toxe M-32
bigc M-136	gbs OP-46	nigbinv M-134	toxm M-54
bigsd M-139	gds OP-11	nigc M-142	toxp M-33
binunit M-56	geomod M-115	njd M-158	toxref M-146
bvd M-206	geomod I-16	njs M-157	tpb M-259
bvj M-319	gm OP-10	noff M-218	tpbsw M-261
bvs M-205	gmbs OP-12	noia M-294	tpbswg M-262
capmod M-207	gmoverid OP-36	noib M-295	trise M-240
cbb OP-31	ibulk OP-34	noic M-296	trise I-13
cbd OP-29	id OP-33	nrd M-233	trnqsmod I-11
cbg OP-28	ids OP-4	nrd I-7	trnqsmod M-208
cbs OP-30	igb OP-43	nrs I-8	type OP-1
cdb OP-23	igbmod M-125	nrs M-234	type M-1
cdd OP-21	igcd OP-45	nsd M-36	u0 M-68
cdg OP-20	igcmmod M-124	nsub M-37	ua M-70
cds OP-22	igcs OP-44	ntnoi M-300	ual M-250
cdsc M-87	igd OP-41	ntox M-145	ub M-71
cdscb M-88	igidl OP-40	paramchk M-237	ubl M-251
cdscd M-89	igs OP-42	pb M-182	uc M-72
cf M-214	ijthdfwd M-165	pbd M-184	uc1 M-252
cgb OP-19	ijthdrev M-163	pbs M-183	ute M-254
cgbo M-170	ijthsfwd M-164	pbsw M-192	vbm M-15
cgd OP-17	ijthsrev M-162	pbswd M-194	vbox M-320
cgdl M-173	imax M-317	pbswg M-201	vbs OP-7
cgdo M-169	imelt M-160	pbswgd M-203	vbx M-14
cgg OP-16	is M-155	pbswgs M-202	vds OP-6
cgidl M-122	jmax M-318	pbsws M-193	vdsat OP-9
cgs OP-18	jmelt M-161	pclm M-76	version M-235
cgs1 M-172	js M-148	pd M-232	vfb M-3
cgso M-168	jsd M-150	pd I-6	vfbcv M-215

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cigbacc M-128	jss M-149	pdiblc1 M-77	vgs OP-5
cigbinv M-132	jswd M-152	pdiblc2 M-78	voff M-92
cigc M-137	jswgd M-154	pdiblc3 M-79	voffcv M-219
cigsd M-140	jswgs M-153	pdits M-84	voffl M-93
cit M-91	jsws M-151	pditsd M-86	vsat M-69
cj M-176	k1 M-5	pditsl M-85	vth OP-8
cjd M-178	k1 O-5	permod M-114	vtho O-4
cjd OP-14	k2 M-6	phin M-4	vtho M-2
cjs M-177	k2 O-6	pigcd M-144	w I-1
cjs OP-15	k3 M-7	pku0 M-281	w M-227
cjsw M-186	k3b M-8	pkvth0 M-284	w0 M-9
cjswd M-188	keta M-30	poxedge M-143	warn M-321
cjswg M-195	kf M-291	prt M-253	weff O-1
cjswgd M-197	kt1 M-246	prwb M-64	wint M-41
cjswgs M-196	kt1l M-247	prwg M-65	wku0 M-280
cjsws M-187	kt2 M-248	ps M-231	wkvth0 M-283
ckappad M-175	ku0 M-271	ps I-5	wl M-47
ckappas M-174	kvsat M-272	pscbe1 M-80	wlc M-224
clc M-212	kvth0 M-273	pscbe2 M-81	wln M-48
cle M-213	l I-2	pta M-258	wlod M-270
csb OP-27	l M-228	ptp M-260	wlodku0 M-276
csd OP-25	leff O-2	pvag M-82	wlodvth M-278
csg OP-24	level M-236	pwr OP-35	wmax M-312
css OP-26	lint M-40	rbdb I-21	wmin M-313
cta M-263	lku0 M-279	rbdb M-309	wnoi M-297
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delk1 I-27	ll M-42	rbodymod I-15	ww M-49
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delta M-83	lln M-43	rbpb I-18	wwl M-51
delvto I-25	llodku0 M-275	rbpd I-19	wwlc M-226
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dlc M-211	lmax M-314	rbps I-20	xgl M-110

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dmcgt M-107	lodk2 M-286	rbsb M-310	xjbvd M-167
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dmdg M-106	lpeb M-11	rdsmod M-57	xl M-118
droul M-74	lw M-44	rdsb M-58	xpart M-220
dskip M-159	lwc M-222	rdsbmin M-59	xrcrg1 M-304
dsub M-95	lwl M-46	rdw M-60	xrcrg2 M-305
dtox M-34	lwlc M-223	rdwmin M-61	xt M-55
dvt0 M-16	lwn M-45	region OP-2	xti M-255
dvt0w M-21	m I-9	region I-10	xtid M-257
dvt1 M-17	mbe0 M-326	reversed OP-3	xtis M-256
dvt1w M-22	mbewl M-325	rgatemod I-14	xw M-117
dvt2 M-18	meto M-171	rgatemod M-101	

BSIMSOI-PD/FD Transistor (bsimsoi)

Description

B3SOI is an SOI model developed by U.C. Berkeley based on bsim3v3. B3SOI devices require that you use a model statement. This is the B3SOI version-2.0 model.

This device is supported within altergroups.

Instance Definition

```
Name d g s e [p] [b] [t] ModelName parameter=value ...
```

Instance Parameters

- 1 w (m) Channel width.
- 2 l (m) Channel length.
- 3 as (m²) Area of source diffusion.

Spectre Circuit Simulator Reference

Component Statements Part I

4	<code>ad</code> (m^2)	Area of drain diffusion.
5	<code>ps</code> (m)	Perimeter of source diffusion.
6	<code>pd</code> (m)	Perimeter of drain diffusion.
7	<code>nrd</code> (m/m)	Number of squares of drain diffusion.
8	<code>nrs</code> (m/m)	Number of squares of source diffusion.
9	<code>nrb</code> (m/m)	Number of body squares.
10	<code>m=1</code>	Multiplicity factor (number of MOSFETs in parallel).
11	<code>region=triode</code>	Estimated operating region. Possible values are <code>off</code> , <code>triode</code> , <code>sat</code> , or <code>subth</code> .
12	<code>rth0</code> (Ω)	Thermal resistance.
13	<code>cth0</code> (F)	Thermal capacitance.
14	<code>bjtoff=0</code>	BJT off flag.
15	<code>nbc=0</code> m/m	Number of body contact isolation edge.
16	<code>nseg=1</code> m/m	Number of segments for channel width partitioning.
17	<code>pdbcpr=0</code> m	Perimeter length for body contact parasitic at drain.
18	<code>psbcpr=0</code> m	Perimeter length for body contact parasitic at source.
19	<code>agbcpr=0</code> m	Gate to body overlap for body contact parasitic.
20	<code>aebcpr=0</code> m	Gate to body overlap for body contact parasitic.
21	<code>vbsusr=0.0</code> V	Optional initial value of <code>Vbs</code> for transient.
22	<code>tnodeout=0</code>	Temperature node flag associated with T node.

Model Definition

```
model modelName bsimsoi parameter=value ...
```


Spectre Circuit Simulator Reference

Component Statements Part I

Model Parameters

Device type parameters

- | | | |
|---|--------------------------|--|
| 1 | <code>type=n</code> | Transistor type.
Possible values are n or p. |
| 2 | <code>version=2.0</code> | Model version selector. The available version is 2.0. |
| 3 | <code>soimod=0</code> | SOI model selector. SoiMod=0:PD module. SoiMod=1: FD module. |

Threshold voltage parameters

- | | | |
|----|---|--|
| 4 | <code>vtho (V)</code> | Threshold voltage at zero body bias for long-channel devices. For enhancement-mode devices, $v_{tho} > 0$ for n-channel and $v_{th} < 0$ for p-channel. Default value is calculated from other model parameters. |
| 5 | <code>k1=0.5 \sqrt{V}</code> | Body-effect coefficient. |
| 6 | <code>k1w1=0.0 m</code> | First body effect width dependent parameter. |
| 7 | <code>k1w2=0.0 m</code> | Second body effect width dependent parameter. |
| 8 | <code>k2=-0.0186</code> | Charge-sharing parameter. |
| 9 | <code>k3=0</code> | Narrow width coefficient. |
| 10 | <code>k3b=0 1/V</code> | Narrow width coefficient. |
| 11 | <code>w0=2.5e-6 m</code> | Narrow width coefficient. |
| 12 | <code>n1x=1.74e-7 m</code> | Lateral nonuniform doping coefficient. |
| 13 | <code>gamma1 (\sqrt{V})</code> | Body-effect coefficient near the surface. |
| 14 | <code>gamma2 (\sqrt{V})</code> | Body-effect coefficient in the bulk. |
| 15 | <code>vbx (V)</code> | Threshold voltage transition body voltage. |

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Component Statements Part I

16	$v_{bm} = -3 \text{ V}$	Maximum applied body voltage.
17	$dvt0 = 2.2$	First coefficient of short-channel effects.
18	$dvt1 = 0.53$	Second coefficient of short-channel effects.
19	$dvt2 = -0.032 \text{ 1/V}$	Body-bias coefficient of short-channel effects.
20	$dvt0w = 0$	First coefficient of narrow-width effects.
21	$dvt1w = 5.3e6$	Second coefficient of narrow-width effects.
22	$dvt2w = -0.032 \text{ 1/V}$	Body-bias coefficient of narrow-width effects.
23	$a0 = 1$	Nonuniform depletion width effect coefficient.
24	$b0 = 0 \text{ m}$	Bulk charge coefficient due to narrow width effect.
25	$b1 = 0 \text{ m}$	Bulk charge coefficient due to narrow width effect.
26	$a1 = 0$	No-saturation coefficient.
27	$a2 = 1$	No-saturation coefficient.
28	$ags = 0 \text{ F/m}^2 \text{ V}$	Gate-bias dependence of a_{bulk} .
29	$keta = -0.6 \text{ 1/V}$	Body-bias coefficient for non-uniform depletion width effect.
30	$ketas = 0.0 \text{ V}$	Surface Potential adjustment for bulk charge effect.

Process parameters

31	$n_{sub} = 6e16 \text{ cm}^{-3}$	Substrate doping concentration.
32	$n_{ch} = 1.7e17 \text{ cm}^{-3}$	Peak channel doping concentration.
33	$n_{gate} \text{ (cm}^{-3}\text{)}$	Poly-gate doping concentration.
34	$x_j = 0.15e-6 \text{ m}$	Source/drain junction depth.
35	$l_{int} = 0 \text{ m}$	Lateral diffusion for one side.

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Component Statements Part I

36	wint=0 m	Width reduction for one side.
37	ll=0 m	Length dependence of delta L.
38	lln=1	Length exponent of delta L.
39	lw=0 m	Width dependence of delta L.
40	lwn=1	Width exponent of delta L.
41	lwl=0 m ²	Area dependence of delta L.
42	wl=0 m	Length dependence of delta W.
43	wln=1	Length exponent of delta W.
44	ww=0 m	Width dependence of delta W.
45	wwn=1	Width exponent of delta W.
46	wwl=0 m ²	Area dependence of delta W.
47	dwg=0 m/v	Gate-bias dependence of channel width.
48	dwb=0 m/ \sqrt{v}	Body-bias dependence of channel width.
49	dwbc=0.0 m	Width offset for body contact isolation edge.
50	tox=1e-8 m	Gate oxide thickness.
51	tbox=3e-7 m	Buried oxide thickness.
52	tsi=1e-7 m	Silicon film thickness.
53	xt=1.55e-7 m	Doping depth.
54	rdsw=100 Ω μm	Width dependence of drain-source resistance.
55	prwb=0 1/ \sqrt{v}	Body-effect coefficient for Rds.
56	prwg=0 1/V	Gate-effect coefficient for Rds.
57	wr=1	Width offset for parasitic resistance.

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Component Statements Part I

- 58 `xl=0 m` Length variation due to masking and etching.
- 59 `xw=0 m` Width variation due to masking and etching.
- 60 `binunit=1` Bin parameter unit selector. 1 for microns and 2 for meters.

Mobility parameters

- 61 `mobmod=1` Mobility model selector.
- 62 `u0=670 cm2/V s` Low-field surface mobility at t_{nom} . Default is 250 for PMOS.
- 63 `vsat=8e4 m/s` Carrier saturation velocity at t_{nom} .
- 64 `ua=2.25e-9 m/v` First-order mobility reduction coefficient.
- 65 `ub=5.87e-19 m2/v2` Second-order mobility reduction coefficient.
- 66 `uc=-4.65e-11 m/v2` Body-bias dependence of mobility. Default is -0.046 and unit is 1/V for `mobmod=3`.

Output resistance parameters

- 67 `drout=0.56` DIBL effect on output resistance coefficient.
- 68 `pclm=1.3` Channel length modulation coefficient.
- 69 `pdiblc1=0.39` First coefficient of drain-induced barrier lowering.
- 70 `pdiblc2=8.6e-3` Second coefficient of drain-induced barrier lowering.
- 71 `pdiblcb=0 1/V` Body-effect coefficient for DIBL.
- 72 `pvag=0` Gate dependence of Early voltage.
- 73 `delta=0.01 V` Effective drain voltage smoothing parameter.

Spectre Circuit Simulator Reference

Component Statements Part I

Subthreshold parameters

- 74 $cdsc=2.4e-4 \text{ F/m}^2$ Source/drain and channel coupling capacitance.
- 75 $cdscb=0 \text{ F/m}^2 \text{ V}$ Body-bias dependence of $cdsc$.
- 76 $cdscd=0 \text{ F/m}^2 \text{ V}$ Drain-bias dependence of $cdsc$.
- 77 $nfactor=1$ Subthreshold swing coefficient.
- 78 $cit=0 \text{ F}$ Interface trap parameter for subthreshold swing.
- 79 $voff=-0.08 \text{ V}$ Threshold voltage offset.
- 80 $dsub=drout$ DIBL effect in subthreshold region.
- 81 $eta0=0.08$ DIBL coefficient subthreshold region.
- 82 $etab=-0.07 \text{ 1/V}$ Body-bias dependence of $et0$.

Substrate current parameters

- 83 $alpha0=0 \text{ m/v}$ Substrate current impact ionization coefficient.
- 84 $beta0=0 \text{ 1/V}$ First V_{ds} dependent parameter of impact ionization current.
- 85 $fbjtii=0.0$ Fraction of bipolar current affecting the impact ionization.
- 86 $beta1=0$ Second V_{ds} dependent parameter of impact ionization current.
- 87 $beta2=0 \text{ V}$ Third V_{ds} dependent parameter of impact ionization current.
- 88 $vdsatii0=0.9 \text{ V}$ Nominal drain saturation voltage at threshold for impact ionization current.
- 89 $tii=0$ Temperature dependent parameter for impact ionization current.
- 90 $lii=0$ Channel length dependent parameter at threshold for impact ionization current.
- 91 $esatii=1e7 \text{ V/m}$ Saturation channel electric field for impact ionization current.

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92	<code>sii0=0.5 1/V</code>	First V_{gs} dependent parameter for impact ionization current.
93	<code>sii1=0.1 1/V</code>	Second V_{gs} dependent parameter for impact ionization current.
94	<code>sii2=0.0 1/V</code>	Third V_{gs} dependent parameter for impact ionization current.
95	<code>siid=0 1/V</code>	V_{ds} dependent parameter of drain saturation voltage for impact ionization current.

Parasitic resistance parameters

96	<code>rbsh=0 Ω</code>	Extrinsic body contact sheet resistance.
97	<code>rsh=0 Ω/sqr</code>	Source/drain diffusion sheet resistance.
98	<code>rs=0 Ω</code>	Source resistance.
99	<code>rd=0 Ω</code>	Drain resistance.
100	<code>rbody=0 F</code>	Body resistance.
101	<code>rsc=0 Ω</code>	Source contact resistance.
102	<code>rdc=0 Ω</code>	Drain contact resistance.
103	<code>rss=0 Ω m</code>	Scalable source resistance.
104	<code>rdd=0 Ω m</code>	Scalable drain resistance.
105	<code>hdif=0 m</code>	Length of heavily doped diffusion.
106	<code>ldif=0 m</code>	Lateral diffusion beyond the gate.
107	<code>minr=0.1 Ω</code>	Minimum source/drain resistance.

Junction diode model parameters

108	<code>dskip=yes</code>	Use simple piece-wise linear model for diode currents below $0.1 \cdot i_{abstol}$. Possible values are <code>no</code> or <code>yes</code> .
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Component Statements Part I

Overlap capacitance parameters

- 109 c_{gso} (F/m) Gate-source overlap capacitance.
- 110 c_{gdo} (F/m) Gate-drain overlap capacitance.
- 111 $c_{geo}=0.0$ F/m Gate-substrate overlap capacitance.
- 112 $c_{gbo}=2 D_{wc} C_{ox}$ F/m Gate-bulk overlap capacitance..
- 113 $c_{gsl}=0$ F/m Gate-source overlap capacitance in LDD region.
- 114 $c_{gdl}=0$ F/m Gate-drain overlap capacitance in LDD region.
- 115 $c_{kappa}=0.6$ Overlap capacitance fitting parameter.

Junction capacitance model parameters

- 116 $c_{jswg}=c_{jsw}$ F/m Zero-bias gate-side junction capacitance density.
- 117 $m_{jswg}=0.5$ Gate-side junction grading coefficient.
- 118 $p_{bswg}=0.7$ V Gate-side junction built-in potential.
- 119 $t_t=1e-12$ s Transit time.
- 120 $n_{dif}=1$ Power coefficient of channel length dependency for diffusion capacitance.
- 121 $l_{dif0}=1$ Power coefficient of channel length dependency for diffusion capacitance.

Charge model selection parameters

- 122 $capmod=2$ Intrinsic charge model.
- 123 $d_{wc}=w_{int}$ m Delta W for capacitance model.
- 124 $delvt=0.0$ V Threshold voltage adjustment for C-V.

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125	<code>fbody=1.0</code>	Scaling factor for body charge.
126	<code>dlc=lint m</code>	Delta L for capacitance model.
127	<code>dlcb=lint m</code>	Length offset fitting parameter for body charge.
128	<code>dlbg=0.0 m</code>	Length offset fitting parameter for backgate charge.
129	<code>clc=1e-8 m</code>	Intrinsic capacitance fitting parameter.
130	<code>cle=0.0</code>	Intrinsic capacitance fitting parameter.
131	<code>cf (F/m)</code>	Fringe capacitance parameter.
132	<code>xpart=0</code>	Drain/source channel charge partition in saturation for BSIM charge model, use 0.0 for 40/60, 0.5 for 50/50, or 1.0 for 0/100.

Default instance parameters

133	<code>w=5e-6 m</code>	Default channel width.
134	<code>l=5e-6 m</code>	Default channel length.
135	<code>as=0 m²</code>	Default area of source diffusion.
136	<code>ad=0 m²</code>	Default area of drain diffusion.
137	<code>ps=0 m</code>	Default perimeter of source diffusion.
138	<code>pd=0 m</code>	Default perimeter of drain diffusion.
139	<code>nrd=0 m/m</code>	Default number of squares of drain diffusion.
140	<code>nrs=0 m/m</code>	Default number of squares of source diffusion.
141	<code>nrb=0 m/m</code>	Default body squares.
142	<code>paramchk=1</code>	Model parameter checking selector.

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Temperature effects parameters

143	t_{nom} (C)	Parameters measurement temperature. Default set by <code>options</code> .
144	$t_{max}=500$ C	Maximum device temperature above ambient.
145	$shmod=0$	Self-heating selector.
146	$t_{lev}=0$	DC temperature selector.
147	$t_{levc}=0$	AC temperature selector.
148	$kt1=-0.11$ V	Temperature coefficient for threshold voltage.
149	$kt1l=0$ v m	Temperature coefficient for threshold voltage.
150	$kt2=0.022$	Temperature coefficient for threshold voltage.
151	$at=3.3e4$ m/s	Temperature coefficient for v_{sat} .
152	$tcjswg=0$ 1/K	Temperature coefficient of C_{jswg} .
153	$tpbswg=0$ V/K	Temperature coefficient of P_{bswg} .
154	$ua1=4.31e-9$ m/v	Temperature coefficient for u_a .
155	$ub1=-7.61e-18$ m^2/v^2	Temperature coefficient for u_b .
156	$uc1=-5.5e-11$ m/v^2	Temperature coefficient for u_c . Default is -0.056 for <code>mobmod=3</code> .
157	$prt=0$ Ω	Temperature coefficient for R_{ds} .
158	$trs=0$ 1/C	Temperature parameter for source resistance.
159	$trd=0$ 1/C	Temperature parameter for drain resistance.
160	$ute=-1.5$	Mobility temperature exponent.
161	$cth0=1e-5$ F	Self-heating thermal capacitance.
162	$rth0=0$ Ω	Self-heating thermal resistance.

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163	<code>ntrecf=0</code>	Temperature coefficient of Ntrecf.
164	<code>ntreocr=0</code>	Temperature coefficient of Ntrecr.
165	<code>xbjt=2</code>	BJT current temperature exponent.
166	<code>xdif=2</code>	Diffusion current temperature exponent.
167	<code>xrec=1</code>	Recombination current temperature exponent.
168	<code>xtun=0</code>	Tunneling current temperature exponent.

Noise model parameters

169	<code>noimod=1</code>	Noise model selector.
170	<code>kf=0</code>	Flicker (1/f) noise coefficient.
171	<code>af=1</code>	Flicker (1/f) noise exponent.
172	<code>ef=1</code>	Flicker (1/f) noise frequency exponent.
173	<code>noia=1e20</code>	Oxide trap density coefficient. Default is 9.9e18 for pmos.
174	<code>noib=5e4</code>	Oxide trap density coefficient. Default is 2.4e3 for pmos.
175	<code>noic=-1.4e-12</code>	Oxide trap density coefficient. Default is 1.4e-8 for pmos.
176	<code>em=4.1e7 V/m</code>	Maximum electric field.
177	<code>noif=1</code>	Floating body excess noise ideality factor.
178	<code>w0flk=0 m</code>	Width constant for IBM flicker noise equation.

Auto Model Selector parameters

179	<code>wmax=1 m</code>	Maximum channel width for which the model is valid.
180	<code>wmin=0 m</code>	Minimum channel width for which the model is valid.
181	<code>lmax=1 m</code>	Maximum channel length for which the model is valid.

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182 `lmin=0 m` Minimum channel length for which the model is valid.

Operating region warning control parameters

183 `alarm=none` Forbidden operating region.
Possible values are `none`, `off`, `triode`, `sat`, `subth`, or `rev`.

184 `imax=1 A` Maximum allowable current.

185 `bvj= ∞ V` Junction reverse breakdown voltage.

186 `vbox=1e9 tox V` Oxide breakdown voltage.

187 `warn=on` Parameter to turn warnings on and off.
Possible values are `off` or `on`.

SOI specific parameters

188 `kb1=1` Scaling factor for backgate charge.

189 `kb3=1` Backgate coupling coefficient at subthreshold.

190 `agidl=0` GIDL constant.

191 `bgidl=0 V/m` GIDL exponential coefficient.

192 `ngidl=1.2 V` GIDL Vds enhancement coefficient.

193 `ntun=10` Reverse tunneling non-ideality factor.

194 `nrecf0=2.0` Recombination non-ideality factor at forward bias.

195 `nrecr0=10` Recombination non-ideality factor at reversed bias.

196 `vsdfb (F/m)` Source/Drain diffusion flatband voltage.

197 `vsdth` Source/Drain diffusion threshold voltage.

198 `csdmin (F)` Source/Drain diffusion bottom minimum capacitance.

199 `csdesw=0` Source/drain sidewall fringing constant.

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200	<code>ndiode=1</code>	Diode non-ideality factor.
201	<code>asd=0.3</code>	Source/Drain diffusion smoothing parameter.
202	<code>isbjt=1e-6 A</code>	BJT saturation current.
203	<code>isdif=0 A</code>	Diffusion saturation current.
204	<code>isrec=1e-5 A</code>	Recombination saturation current.
205	<code>istun=0 A</code>	Tunneling saturation current.
206	<code>ln=2e-6 m</code>	Electron diffusion length.
207	<code>vrec0=0 V</code>	Voltage dependent parameter for recombination current.
208	<code>vtun0=0 V</code>	Voltage dependent parameter for tunneling current.
209	<code>nbjt=1</code>	Power coefficient of channel length dependency for bipolar current.
210	<code>lbjt0=0.20e-6 m</code>	Reference channel length for bipolar current.
211	<code>vabjt=10 V</code>	Early voltage for bipolar current.
212	<code>aely=0 V</code>	Channel length dependency of early voltage for bipolar current.
213	<code>ahli=0</code>	High level injection parameter for bipolar current.
214	<code>frbody=1</code>	Layout dependent body-resistance coefficient.
215	<code>vevb=0.075</code>	Vaux parameter for valence-band electron tunneling.
216	<code>vecb=0.026</code>	Vaux parameters for conduction-band electron tunneling.
217	<code>dtoxcv=0.0 m</code>	Delta oxide thickness in Capmod3.
218	<code>aigc=0.43</code>	Parameter for I_{gc} .
219	<code>bigc=0.054</code>	Parameter for I_{gc} .
220	<code>cigc=0.075 1/V</code>	Parameter for I_{gc} .

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221	<code>aigsd=0.43</code>	Parameter for I_{gs}/I_{gd} .
222	<code>bigsd=0.054</code>	Parameter for I_{gs}/I_{gd} .
223	<code>cigsd=0.075</code> 1/V	parameter for I_{gs}/I_{gd} .
224	<code>nigc=1.0</code>	Parameter for I_{gc} slope.
225	<code>pigcd=1.0</code>	Parameter for I_{gc} partition.
226	<code>poxedge=1.0</code>	Factor for the gate edge T_{ox} .
227	<code>dlcig=0.0</code> m	Delta Length for I_g model.
228	<code>vbsa=0.0</code> V	Non-ideal offset voltage.
229	<code>nofffd=1.0</code>	Smooth parameter in FD module.
230	<code>vofffd=0.0</code>	Smooth parameter in FD module.
231	<code>k1b=1.0</code>	First backgate body effect parameter.
232	<code>k2b=0.0</code>	Second backgate body effect parameter for short channel effect.
233	<code>dk2b=0.0</code>	Third backgate body effect parameter for short channel effect.
234	<code>dvbd0=0.0</code>	First short-channel effect parameter in FD module.
235	<code>dvbd1=0.0</code>	Second short-channel effect parameter in FD module.
236	<code>moinfd=1.0e3</code>	Gate bias dependence coefficient of surface potential.

Gate tunneling parameters

237	<code>wth0=0.0</code> μm	Minimum width for thermal resistance calculation..
238	<code>rhalo=1.0e15</code> Ω/sqr	Body halo sheet resistance.
239	<code>ntox=1.0</code>	Power term of gate current.
240	<code>toxref=2.5e-9</code> m	Target oxide thickness.

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241	<code>ebg=1.2 V</code>	Effective bandgap in gate current calculation.
242	<code>nevb=3.0</code>	Valence-band electron non-ideality factor.
243	<code>alphagb1=0.35</code>	First V_{ox} dependent parameter for gate current in inversion..
244	<code>betagb1=0.03</code>	Second V_{ox} dependent parameter for gate current in inversion..
245	<code>vgb1=300</code>	Third V_{ox} dependent parameter for gate current in inversion..
246	<code>alphagb2=0.43</code>	First V_{ox} dependent parameter for gate current in accumulation..
247	<code>betagb2=0.05</code>	Second V_{ox} dependent parameter for gate current in accumulation..
248	<code>necb=1.0</code>	Conduction-band electron non-ideality factor.
249	<code>vgb2=17</code>	Third V_{ox} dependent parameter for gate current in accumulation..
250	<code>toxqm=Tox m</code>	Effective oxide thickness considering quantum effects..
251	<code>voxh=5.0 V</code>	Limit of V_{ox} in gate current calculation..
252	<code>deltavox=0.005 V</code>	Smoothing parameter in the V_{ox} smoothing function..
253	<code>igbmod=0</code>	Gate-body tunneling current model selector.
254	<code>igcmod=0</code>	Gate-channel tunneling current model selector.

Length dependent parameters

255	<code>llc=0 m</code>	Length dependence of delta LC.
256	<code>lwc=0 m</code>	Width dependence of delta LC.
257	<code>lwlc=0 m²</code>	Area dependence of delta LC.

Width dependent parameters

258	<code>wlc=0 m</code>	Length dependence of delta WC.
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259 $w_{WC}=0$ m Width dependence of delta WC.

260 $w_{wlc}=0$ m² Area dependence of delta WC.

Cross-term dependent parameters

Auto Model Selection

Many models need to be characterized for different geometries in order to obtain accurate results for model development. The model selector program automatically searches for a model with the length and width range specified in the instance statement and uses this model in the simulations.

For the auto model selector program to find a specific model, the models to be searched should be grouped together within braces. Such a group is called a model group. An opening brace is required at the end of the line defining each model group. Every model in the group is given a name followed by a colon and the list of parameters. Also, the four geometric parameters `lmax`, `lmin`, `wmax`, and `wmin` should be given. The selection criteria to choose a model is as follows:

$$l_{min} \leq inst_length < l_{max} \text{ and } w_{min} \leq inst_width < w_{max}$$

Example:

```
model ModelName ModelType {  
  1: <model parameters> lmin=2 lmax=4 wmin=1 wmax=2  
  2: <model parameters> lmin=1 lmax=2 wmin=2 wmax=4  
  3: <model parameters> lmin=2 lmax=4 wmin=4 wmax=6  
}
```

Then for a given instance

```
M1 1 2 3 4 ModelName w=3 l=1.5
```

the program would search all the models in the model group with the name `ModelName` and then pick the first model whose geometric range satisfies the selection criteria. In the preceding example, the auto model selector program would choose `ModelName.2`.

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You must specify both length (l) and width (w) on the device instance line to enable automatic model selection.

Output Parameters

1	<code>w_{eff}</code> (m)	Effective channel width.
2	<code>l_{eff}</code> (m)	Effective channel length.
3	<code>r_{th_{eff}}</code> (Ω)	Effective thermal resistance.
4	<code>C_{th_{eff}}</code> (F)	Effective thermal capacitance.
5	<code>r_{seff}</code> (Ω)	Effective source resistance.
6	<code>r_{deff}</code> (Ω)	Effective drain resistance.

Operating-Point Parameters

1	<code>type=n</code>	Transistor type. Possible values are <code>n</code> or <code>p</code> .
2	<code>region=triode</code>	Estimated operating region. Possible values are <code>off</code> , <code>triode</code> , <code>sat</code> , or <code>subth</code> .
3	<code>reversed</code>	Reverse mode indicator. Possible values are <code>no</code> or <code>yes</code> .
4	<code>v_{gs}</code> (V)	Gate-source voltage.
5	<code>v_{ds}</code> (V)	Drain-source voltage.
6	<code>v_{bs}</code> (V)	Bulk-source voltage.
7	<code>v_{bgs}</code> (V)	Back-Gate-source voltage.
8	<code>i_{ds}</code> (A)	Resistive drain-to-source current.
9	<code>i_c</code> (A)	BJT collector current.
10	<code>i_{sgidl}</code> (A)	Source GIDL current.

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11	idgidl (A)	Drain GIDL current.
12	iii (A)	Impact ionization current.
13	ibd (A)	Resistive bulk-to-drain junction current.
14	igbt (A)	Gate-to-body tunneling current.
15	ibs (A)	Resistive bulk-to-source junction current.
16	vth (V)	Threshold voltage.
17	vdsat (V)	Drain-source saturation voltage.
18	gm (S)	Common-source transconductance.
19	gds (S)	Common-source output conductance.
20	gmb (S)	Body-transconductance.
21	gmbg (S)	Back-gate-transconductance.
22	ueff (cm ² /V s)	Effective mobility.
23	betaeff (A/V ²)	Effective beta.
24	qg (Coul)	Gate charge.
25	qd (Coul)	Drain charge.
26	qs (Coul)	Source charge.
27	qb (Coul)	Body charge.
28	qbg (Coul)	Back-Gate charge.
29	cgg (F)	dQg_dVg.
30	cgd (F)	dQg_dVd.
31	cgs (F)	dQg_dVs.
32	cgb (F)	dQg_dVbk .

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33	<code>cdg</code> (F)	<code>dQd_dVg</code> .
34	<code>cdd</code> (F)	<code>dQd_dVd</code> .
35	<code>cds</code> (F)	<code>dQd_dVs</code> .
36	<code>cdb</code> (F)	<code>dQd_dVb</code> .
37	<code>csq</code> (F)	<code>dQs_dVg</code> .
38	<code>csd</code> (F)	<code>dQs_dVd</code> .
39	<code>css</code> (F)	<code>dQs_dVs</code> .
40	<code>csb</code> (F)	<code>dQs_dVb</code> .
41	<code>cbg</code> (F)	<code>dQb_dVg</code> .
42	<code>cbd</code> (F)	<code>dQb_dVd</code> .
43	<code>cbs</code> (F)	<code>dQb_dVs</code> .
44	<code>cbb</code> (F)	<code>dQb_dVb</code> .
45	<code>id</code> (A)	Total resistive drain current.
46	<code>is</code> (A)	Total resistive source current.
47	<code>ib</code> (A)	Total resistive bulk current.
48	<code>pwr</code> (W)	Power at op point.
49	<code>gmoverid</code> (1/V)	<code>Gm/Ids</code> .
50	<code>tdev</code> (C)	Temperature rise from ambient.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the

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description for that parameter. For example, a reference of M-35 means the 35th model parameter.

Ctheff	O-4	dsub	M-80	minr	M-107	tbox	M-51
a0	M-23	dtoxcv	M-217	mjswg	M-117	tcjswg	M-152
a1	M-26	dvbd0	M-234	mobmod	M-61	tdev	OP-50
a2	M-27	dvbd1	M-235	moinfd	M-236	tii	M-89
ad	I-4	dvt0	M-17	nbcb	I-15	tlev	M-146
ad	M-136	dvt0w	M-20	nbjt	M-209	tlevc	M-147
aebcp	I-20	dvt1	M-18	nch	M-32	tmax	M-144
aely	M-212	dvt1w	M-21	ndif	M-120	tnodeout	I-22
af	M-171	dvt2	M-19	ndiode	M-200	tnom	M-143
agbcp	I-19	dvt2w	M-22	necb	M-248	tox	M-50
agidl	M-190	dwb	M-48	nevb	M-242	toxqm	M-250
ags	M-28	dwbc	M-49	nfactor	M-77	toxref	M-240
ahli	M-213	dwc	M-123	ngate	M-33	tpbswg	M-153
aigc	M-218	dwg	M-47	ngidl	M-192	trd	M-159
aigsd	M-221	ebg	M-241	nigc	M-224	trs	M-158
alarm	M-183	ef	M-172	nlx	M-12	tsi	M-52
alpha0	M-83	em	M-176	nofffd	M-229	tt	M-119
alphagb1	M-243	esatii	M-91	noia	M-173	type	M-1
alphagb2	M-246	eta0	M-81	noib	M-174	type	OP-1
as	M-135	etab	M-82	noic	M-175	u0	M-62
as	I-3	fbjtii	M-85	noif	M-177	ua	M-64
asd	M-201	fbody	M-125	noimod	M-169	ua1	M-154
at	M-151	frbody	M-214	nrb	M-141	ub	M-65
b0	M-24	gamma1	M-13	nrb	I-9	ub1	M-155
b1	M-25	gamma2	M-14	nrd	M-139	uc	M-66
beta0	M-84	gds	OP-19	nrd	I-7	uc1	M-156
beta1	M-86	gm	OP-18	nrecf0	M-194	ueff	OP-22
beta2	M-87	gmb	OP-20	nrecr0	M-195	ute	M-160
betaeff	OP-23	gmbg	OP-21	nrs	M-140	vabjt	M-211

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betagb1 M-244	gmoverid OP-49	nrs I-8	vbgs OP-7
betagb2 M-247	hdif M-105	nseg I-16	vbm M-16
bgid1 M-191	ib OP-47	nsub M-31	vbox M-186
bigc M-219	ibd OP-13	ntox M-239	vbs OP-6
bigsd M-222	ibs OP-15	ntrecf M-163	vbsa M-228
binunit M-60	ic OP-9	ntrechr M-164	vbsusr I-21
bjtoff I-14	id OP-45	ntun M-193	vbv M-15
bvj M-185	idgid1 OP-11	paramchk M-142	vds OP-5
capmod M-122	ids OP-8	pbswg M-118	vdsat OP-17
cbb OP-44	igbmod M-253	pclm M-68	vdsatii0 M-88
cbd OP-42	igbt OP-14	pd I-6	vecb M-216
cbg OP-41	igcmmod M-254	pd M-138	version M-2
cbs OP-43	iii OP-12	pdbcpl I-17	vevb M-215
cdb OP-36	imax M-184	pdiblc1 M-69	vgb1 M-245
cdd OP-34	is OP-46	pdiblc2 M-70	vgb2 M-249
cdg OP-33	isbjt M-202	pdiblc3 M-71	vgs OP-4
cds OP-35	isdif M-203	pigcd M-225	voff M-79
cdsc M-74	isgid1 OP-10	poxedge M-226	vofffd M-230
cdscb M-75	isrec M-204	pvt M-157	voxh M-251
cdscd M-76	istun M-205	prwb M-55	vrec0 M-207
cf M-131	k1 M-5	prwg M-56	vsat M-63
cgb OP-32	k1b M-231	ps I-5	vsdfb M-196
cgbo M-112	k1w1 M-6	ps M-137	vsdth M-197
cgd OP-30	k1w2 M-7	psbcp I-18	vth OP-16
cgdl M-114	k2 M-8	pvag M-72	vtho M-4
cgdo M-110	k2b M-232	pwr OP-48	vtun0 M-208
cgeo M-111	k3 M-9	qb OP-27	w M-133
cgg OP-29	k3b M-10	qbg OP-28	w I-1
cgs OP-31	kb1 M-188	qd OP-25	w0 M-11
cgs1 M-113	kb3 M-189	qg OP-24	w0flk M-178
cgso M-109	keta M-29	qs OP-26	warn M-187
cigc M-220	ketas M-30	rbody M-100	weff O-1

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cigsd M-223	kf M-170	rbsh M-96	wint M-36
cit M-78	kt1 M-148	rd M-99	wl M-42
cjswg M-116	kt11 M-149	rdc M-102	wlc M-258
ckappa M-115	kt2 M-150	rdd M-104	wln M-43
clc M-129	l I-2	rdeff O-6	wmax M-179
cle M-130	l M-134	rdsw M-54	wmin M-180
csb OP-40	lbjt0 M-210	region OP-2	wr M-57
csd OP-38	ldif M-106	region I-11	wth0 M-237
csdesw M-199	ldif0 M-121	reversed OP-3	ww M-44
csdmin M-198	leff O-2	rhalo M-238	wwc M-259
csg OP-37	lii M-90	rs M-98	wwl M-46
css OP-39	lint M-35	rsc M-101	wwlc M-260
cth0 I-13	ll M-37	rseff O-5	wwn M-45
cth0 M-161	llc M-255	rsh M-97	xbjt M-165
delta M-73	lln M-38	rss M-103	xdif M-166
deltavox M-252	lmax M-181	rth0 I-12	xj M-34
delvt M-124	lmin M-182	rth0 M-162	xl M-58
dk2b M-233	ln M-206	rtheff O-3	xpart M-132
dlbg M-128	lw M-39	shmod M-145	xrec M-167
dlc M-126	lwc M-256	sii0 M-92	xt M-53
dlcb M-127	lwl M-41	sii1 M-93	xtun M-168
dlcig M-227	lwlc M-257	sii2 M-94	xw M-59
drout M-67	lwn M-40	siid M-95	
dskip M-108	m I-10	soimod M-3	

BTA SOI Transistor (btasoi)

Description

BTASOI is an SOI model developed by BTA Technology based on bsim3v3. It is a new, simple and compact SOI model that can accommodate both the fully-depleted, FD and partially-depleted, PD modes, adopting the transition voltage, V_{tr} , for the definition of body condition. It can also simulate the special characteristics of SOI devices such as kink effect and

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reduction of saturation current due to self-heating. Simulation results with this model are in excellent agreement with the experimental data for 0.25um SIMOX technology. BTASOI devices require that you use a model statement.

If you want to get more information about this model, please contact BTA Technology at <http://www.btat.com>

This device is supported within altergroups.

Sample Instance Statement

```
m5 (1 2 0 0) nchmod l=1.5u w=100u as=450p ad=450p pd=209u ps=209u m=1
```

Sample Model Statement

```
model nchmod btasoi type=n b3v3mod=no version=3.1 vtho=0.62 k1=0.672 k2=0.038  
nlx=1.14e-7 dvt0=4.1 a0=1.08 nch=2.65e17 u0=4.01e-2 a1=0 a2=1 ags=9.8e-4  
vsat=1.77e5
```

Instance Definition

Name d g s [bg] [b] ModelName parameter=value ...

Instance Parameters

1	w (m)	Channel width.
2	l (m)	Channel length.
3	as (m ²)	Area of source diffusion.
4	ad (m ²)	Area of drain diffusion.
5	ps (m)	Perimeter of source diffusion.
6	pd (m)	Perimeter of drain diffusion.
7	nrd (m/m)	Number of squares of drain diffusion.
8	nrs (m/m)	Number of squares of source diffusion.
9	m=1	Multiplicity factor (number of MOSFETs in parallel).

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- 10 `region=triode` Estimated operating region. Spectre outputs number (0-4) in a rawfile.
Possible values are `off`, `triode`, `sat`, or `subth`.
- 11 `trise` Temperature rise from ambient.
- 12 `rbody (Ω)` Body resistance.

Model Definition

```
model modelName btasoi parameter=value ...
```

Model Parameters

Device type parameters

- 1 `type=n` Transistor type.
Possible values are `n` or `p`.
- 2 `b3v3mod=no` B3v3 compatible flag.
Possible values are `no` or `yes`.
- 3 `version=3.1` Model version selector.
- 4 `btasoiver=1.0` BTASOI Model version selector.

Threshold voltage parameters

- 5 `vtho (V)` Threshold voltage at zero body bias for long-channel devices.
For enhancement-mode devices, `vtho` > 0 for n-channel and `vth` < 0 for p-channel. Default value is calculated from other model parameters.
- 6 `k1=0.5 \sqrt{V}` Body-effect coefficient.
- 7 `k2=-0.0186` Charge-sharing parameter.
- 8 `k3=80` Narrow width coefficient.
- 9 `k3b=0 1/V` Narrow width coefficient.

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10	$w0=2.5e-6$ m	Narrow width coefficient.
11	$n1x=1.74e-7$ m	Lateral nonuniform doping coefficient.
12	$\gamma1$ (\sqrt{V})	Body-effect coefficient near the surface.
13	$\gamma2$ (\sqrt{V})	Body-effect coefficient in the bulk.
14	vbx (V)	Threshold voltage transition body voltage.
15	$vbm=-3$ V	Maximum applied body voltage.
16	$dvt0=2.2$	First coefficient of short-channel effects.
17	$dvt1=0.53$	Second coefficient of short-channel effects.
18	$dvt2=-0.032$ 1/V	Body-bias coefficient of short-channel effects.
19	$dvt0w=0$	First coefficient of narrow-width effects.
20	$dvt1w=5.3e6$	Second coefficient of narrow-width effects.
21	$dvt2w=-0.032$ 1/V	Body-bias coefficient of narrow-width effects.
22	$a0=1$	Nonuniform depletion width effect coefficient.
23	$b0=0$ m	Bulk charge coefficient due to narrow width effect.
24	$b1=0$ m	Bulk charge coefficient due to narrow width effect.
25	$a1=0$	No-saturation coefficient.
26	$a2=1$	No-saturation coefficient.
27	$ags=0$ F/m ² V	Gate-bias dependence of A_{bulk} .
28	$keta=-0.047$ 1/V	Body-bias coefficient for non-uniform depletion width effect.

Process parameters

29	$n_{sub}=6e16$ cm ⁻³	Substrate doping concentration.
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30	$n_{ch}=1.7e17 \text{ cm}^{-3}$	Peak channel doping concentration.
31	$n_{gate} \text{ (cm}^{-3}\text{)}$	Poly-gate doping concentration.
32	$x_j=0.15e-6 \text{ m}$	Source/drain junction depth.
33	$l_{int}=0 \text{ m}$	Lateral diffusion for one side.
34	$w_{int}=0 \text{ m}$	Width reduction for one side.
35	$l_l=0 \text{ m}$	Length dependence of delta L.
36	$l_{ln}=1$	Length exponent of delta L.
37	$l_w=0 \text{ m}$	Width dependence of delta L.
38	$l_{wn}=1$	Width exponent of delta L.
39	$l_{wl}=0 \text{ m}^2$	Area dependence of delta L.
40	$w_l=0 \text{ m}$	Length dependence of delta W.
41	$w_{ln}=1$	Length exponent of delta W.
42	$w_w=0 \text{ m}$	Width dependence of delta W.
43	$w_{wn}=1$	Width exponent of delta W.
44	$w_{wl}=0 \text{ m}^2$	Area dependence of delta W.
45	$d_{wg}=0 \text{ m/v}$	Gate-bias dependence of channel width.
46	$d_{wb}=0 \text{ m}/\sqrt{v}$	Body-bias dependence of channel width.
47	$t_{ox}=1.5e-8 \text{ m}$	Gate oxide thickness.
48	$t_{box}=4e-7 \text{ m}$	Buried oxide thickness.
49	$t_{si}=8e-8 \text{ m}$	Silicon film thickness.
50	$x_t=1.55e-7 \text{ m}$	Doping depth.
51	$r_{dsw}=0 \text{ } \Omega \text{ } \mu\text{m}$	Width dependence of drain-source resistance.

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52	<code>prwb=0 1/√v</code>	Body-effect coefficient for Rds.
53	<code>prwg=0 1/V</code>	Gate-effect coefficient for Rds.
54	<code>wr=1</code>	Width offset for parasitic resistance.
55	<code>xl=0 m</code>	Length variation due to masking and etching.
56	<code>xw=0 m</code>	Width variation due to masking and etching.
57	<code>binunit=1</code>	Bin parameter unit selector. 1 for microns and 2 for meters.

Mobility parameters

58	<code>mobmod=1</code>	Mobility model selector.
59	<code>u0=670 cm²/V s</code>	Low-field surface mobility at t_{nom} . Default is 250 for PMOS.
60	<code>vsat=8e4 m/s</code>	Carrier saturation velocity at t_{nom} .
61	<code>ua=2.25e-9 m/v</code>	First-order mobility reduction coefficient.
62	<code>ub=5.87e-19 m²/v²</code>	Second-order mobility reduction coefficient.
63	<code>uc=-4.65e-11 m/v²</code>	Body-bias dependence of mobility. Default is -0.046 and unit is 1/V for <code>mobmod=3</code> .

Output resistance parameters

64	<code>drout=0.56</code>	DIBL effect on output resistance coefficient.
65	<code>pclm=1.3</code>	Channel length modulation coefficient.
66	<code>pdiblc1=0.39</code>	First coefficient of drain-induced barrier lowering.
67	<code>pdiblc2=8.6e-3</code>	Second coefficient of drain-induced barrier lowering.
68	<code>pdiblcb=0 1/V</code>	Body-effect coefficient for DIBL.

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- 69 $\text{pscbe1}=4.24\text{e}8 \text{ V/m}$ First coefficient of substrate current body effect.
- 70 $\text{pscbe2}=1\text{e}-5 \text{ m/v}$ Second coefficient of substrate current body effect.
- 71 $\text{pvag}=0$ Gate dependence of Early voltage.
- 72 $\text{delta}=0.01 \text{ V}$ Effective drain voltage smoothing parameter.

Subthreshold parameters

- 73 $\text{cdsc}=2.4\text{e}-4 \text{ F/m}^2$ Source/drain and channel coupling capacitance.
- 74 $\text{cdscb}=0 \text{ F/m}^2 \text{ V}$ Body-bias dependence of cdsc .
- 75 $\text{cdscd}=0 \text{ F/m}^2 \text{ V}$ Drain-bias dependence of cdsc .
- 76 $\text{nfactor}=1$ Subthreshold swing coefficient.
- 77 $\text{cit}=0 \text{ F}$ Interface trap parameter for subthreshold swing.
- 78 $\text{voff}=-0.08 \text{ V}$ Threshold voltage offset.
- 79 $\text{dsub}=\text{drout}$ DIBL effect in subthreshold region.
- 80 $\text{eta0}=0.08$ DIBL coefficient subthreshold region.
- 81 $\text{etab}=-0.07 \text{ 1/V}$ Body-bias dependence of et0 .

Substrate current parameters

- 82 $\text{alpha0}=0 \text{ m/v}$ Substrate current impact ionization coefficient.
- 83 $\text{beta0}=30 \text{ 1/V}$ Substrate current impact ionization exponent.

Parasitic resistance parameters

- 84 $\text{rsh}=0 \text{ }\Omega/\text{sqr}$ Source/drain diffusion sheet resistance.
- 85 $\text{rs}=0 \text{ }\Omega$ Source resistance.

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Component Statements Part I

86	<code>rd=0</code>	Ω	Drain resistance.
87	<code>rsc=0</code>	Ω	Source contact resistance.
88	<code>rdc=0</code>	Ω	Drain contact resistance.
89	<code>rss=0</code>	Ω m	Scalable source resistance.
90	<code>rdd=0</code>	Ω m	Scalable drain resistance.
91	<code>hdif=0</code>	m	Length of heavily doped diffusion.
92	<code>ldif=0</code>	m	Lateral diffusion beyond the gate.
93	<code>minr=0.1</code>	Ω	Minimum source/drain resistance.

Junction diode model parameters

94	<code>js</code>	(A/m ²)	Bulk junction reverse saturation current density.
95	<code>jsw=0</code>	A/m	Sidewall junction reverse saturation current density.
96	<code>is=1e-14</code>	A	Bulk junction reverse saturation current.
97	<code>n=1</code>		Junction emission coefficient.
98	<code>dskip=yes</code>		Use simple piecewise linear model for diode currents below $0.1 \cdot i_{abstol}$. Possible values are <code>no</code> or <code>yes</code> .
99	<code>imelt=`imaxA`</code>		Explosion current.
100	<code>imelt1=imax</code>	A/m	Explosion current density for <code>is1</code> .
101	<code>imelt2=imax</code>	A/m	Explosion current density for <code>is2</code> .
102	<code>imelt3=imax</code>	A/m	Explosion current density for <code>is3</code> .

Overlap capacitance parameters

103	<code>cgso</code>	(F/m)	Gate-source overlap capacitance.
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104	c_{gdo} (F/m)	Gate-drain overlap capacitance.
105	$c_{gbo}=2$ D_{wc} C_{ox} F/m	Gate-bulk overlap capacitance. The default value is 0 if version=3.0.
106	$m_{eto}=0$ m	Metal overlap in fringing field.
107	$c_{gsl}=0$ F/m	Gate-source overlap capacitance in LDD region.
108	$c_{gdl}=0$ F/m	Gate-drain overlap capacitance in LDD region.
109	$c_{kappa}=0.6$	Overlap capacitance fitting parameter.

Junction capacitance model parameters

110	$c_{bs}=0$ F	Bulk-source zero-bias junction capacitance.
111	$c_{bd}=0$ F	Bulk-drain zero-bias junction capacitance.
112	$c_j=5e-4$ F/m ²	Zero-bias junction bottom capacitance density.
113	$m_j=1/2$	Bulk junction bottom grading coefficient.
114	$p_b=1$ V	Bulk junction built-in potential.
115	$f_c=0.5$	Forward-bias depletion capacitance threshold.
116	$c_{jsw}=5e-10$ F/m	Zero-bias junction sidewall capacitance density.
117	$m_{jsw}=0.33$	Bulk junction sidewall grading coefficient.
118	$p_{bsw}=1$ V	Side-wall junction built-in potential.
119	$c_{jswg}=c_{jsw}$ F/m	Zero-bias gate-side junction capacitance density.
120	$m_{jswg}=m_{jsw}$	Gate-side junction grading coefficient.
121	$p_{bswg}=p_{bsw}$ V	Gate-side junction built-in potential.
122	$f_{csw}=f_c$	Side-wall forward-bias depletion capacitance threshold.

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Component Statements Part I

123 `tau=0 s` Transit time.

Charge model selection parameters

124 `capmod=2` Intrinsic charge model.

125 `dwc=wint m` Delta W for capacitance model.

126 `dlc=lint m` Delta L for capacitance model.

127 `clc=1e-7 m` Intrinsic capacitance fitting parameter.

128 `cle=0.6` Intrinsic capacitance fitting parameter.

129 `cf (F/m)` Fringe capacitance parameter.

130 `vfbcv=-1` Flat-band voltage for `capmod=0`.

131 `xpart=0` Drain/source channel charge partition in saturation for BSIM charge model, use 0.0 for 40/60, 0.5 for 50/50, or 1.0 for 0/100.

Default instance parameters

132 `w=5e-6 m` Default channel width.

133 `l=5e-6 m` Default channel length.

134 `as=0 m2` Default area of source diffusion.

135 `ad=0 m2` Default area of drain diffusion.

136 `ps=0 m` Default perimeter of source diffusion.

137 `pd=0 m` Default perimeter of drain diffusion.

138 `nrd=0 m/m` Default number of squares of drain diffusion.

139 `nrs=0 m/m` Default number of squares of source diffusion.

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Temperature effects parameters

140	<code>tnom (C)</code>	Parameters measurement temperature. Default set by <code>options</code> .
141	<code>tmax=500 C</code>	Maximum device temperature above ambient.
142	<code>trise=0 C</code>	Temperature rise from ambient.
143	<code>selft=0</code>	Self heating option.
144	<code>tlev=0</code>	DC temperature selector.
145	<code>tlevc=0</code>	AC temperature selector.
146	<code>eg=1.12452 V</code>	Energy band gap.
147	<code>gap1=7.02e-4 V/C</code>	Band gap temperature coefficient.
148	<code>gap2=1108 C</code>	Band gap temperature offset.
149	<code>kt1=-0.11 V</code>	Temperature coefficient for threshold voltage.
150	<code>kt11=0 v m</code>	Temperature coefficient for threshold voltage.
151	<code>kt2=0.022</code>	Temperature coefficient for threshold voltage.
152	<code>at=3.3e4 m/s</code>	Temperature coefficient for <code>vsat</code> .
153	<code>ua1=4.31e-9 m/v</code>	Temperature coefficient for <code>ua</code> .
154	<code>ub1=-7.61e-18 m²/v²</code>	Temperature coefficient for <code>ub</code> .
155	<code>uc1=-5.5e-11 m/v²</code>	Temperature coefficient for <code>uc</code> . Default is -0.056 for <code>mobmod=3</code> .
156	<code>prr=0 Ω</code>	Temperature coefficient for <code>Rds</code> .
157	<code>trs=0 1/C</code>	Temperature parameter for source resistance.
158	<code>trd=0 1/C</code>	Temperature parameter for drain resistance.
159	<code>ute=-1.5</code>	Mobility temperature exponent.

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160	$dt1=0$	First temperature coefficient for tau.
161	$dt2=0$	Second temperature coefficient for tau.
162	$x_{ti}=3$	Saturation current temperature exponent.
163	$x_{ti1}=3$	Saturation current temperature exponent.
164	$x_{ti2}=x_{ti1}$	Saturation current temperature exponent.
165	$x_{ti3}=x_{ti1}$	Saturation current temperature exponent.
166	$ptc=0$ V/C	Surface potential temperature coefficient.
167	$pta=0$ V/C	Junction potential temperature coefficient.
168	$ptp=0$ V/C	Sidewall junction potential temperature coefficient.
169	$cta=0$ 1/C	Junction capacitance temperature coefficient.
170	$ctp=0$ 1/C	Sidewall junction capacitance temperature coefficient.

Noise model parameters

171	$noimod=1$	Noise model selector.
172	$kf=0$	Flicker (1/f) noise coefficient.
173	$af=1$	Flicker (1/f) noise exponent.
174	$ef=1$	Flicker (1/f) noise frequency exponent.
175	$noia=1e20$	Oxide trap density coefficient. Default is $9.9e18$ for pmos.
176	$noib=5e4$	Oxide trap density coefficient. Default is $2.4e3$ for pmos.
177	$noic=-1.4e-12$	Oxide trap density coefficient. Default is $1.4e-8$ for pmos.
178	$em=4.1e7$ V/m	Maximum electric field.

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Auto Model Selector parameters

179	<code>wmax=1 m</code>	Maximum channel width for which the model is valid.
180	<code>wmin=0 m</code>	Minimum channel width for which the model is valid.
181	<code>lmax=1 m</code>	Maximum channel length for which the model is valid.
182	<code>lmin=0 m</code>	Minimum channel length for which the model is valid.

Operating region warning control parameters

183	<code>alarm=none</code>	Forbidden operating region. Possible values are <code>none</code> , <code>off</code> , <code>triode</code> , <code>sat</code> , <code>subth</code> , or <code>rev</code> .
184	<code>imax=1 A</code>	Maximum allowable current.
185	<code>bvj=∞ V</code>	Junction reverse breakdown voltage.
186	<code>vbox=1e9 tox V</code>	Oxide breakdown voltage.
187	<code>warn=on</code>	Parameter to turn warnings on and off. Possible values are <code>off</code> or <code>on</code> .

SOI specific parameters

188	<code>vbtho=10 V</code>	Back-gate threshold voltage..
189	<code>vtr0=0.3 V</code>	Long-channel transition body voltage at $V_{ds}=0$.
190	<code>knk=0.01</code>	V_{tr} smoothing factor.
191	<code>dice=0</code>	Drain-induced charge-sharing parameter.
192	<code>dvtrd=0 V</code>	V_{tr} dependence on V_{ds} .
193	<code>dvtrg=1 V</code>	V_{tr} dependence on V_{gs} .
194	<code>dvtrbg=1.0</code>	Smoothing factor for back-gate bias.
195	<code>a0bg=0</code>	Back-gate saturation region coefficient.

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196	dbg=1	Diode fully depletion adjustment factor.
197	dvtr=1.0	Diode back-gate dependence factor.
198	vbgf=0.0	Flat-band voltage for back-gate.
199	rth0=0 Ω	Self-heating thermal resistance.
200	cth0=1 F	Self-heating thermal capacitance.
201	ll=0	Vgs dependence of characteristic length.
202	a _{ii} =0	First parameter for critical field.
203	b _{ii} =0	Second parameter for critical field.
204	c _{ii} =0	Gate dependence of critical field.
205	d _{ii} =0	Body dependence of critical field.
206	ndiode=1	Diode non-ideality factor.
207	nt=1	Reverse tunneling non-ideality factor.
208	is1=1e-16 A	First diode parameter.
209	is2=0 A	Second diode parameter.
210	is3=0 A	Tunneling diode parameter.
211	edl=2e-6 m	Electron diffusion length.
212	kb=0 m	Parasitic bipolar base width.
213	delacc=0.02 V	Capacitance smoothing parameter in accumulation region.
214	delr=0.01 V	V _{bs} smoothing parameter for C-V.
215	dqsq=8e-3 V	V _{tr} smoothing parameter for C-V.
216	a0cv=0.1	A0 for C-V calculation.
217	qgvd0=1	Cgd fitting parameter.

Length dependent parameters (Not listed)

Width dependent parameters (Not listed)

Cross-term dependent parameters (Not listed)

The `jmelt` parameter is used to aid convergence and prevent numerical overflow. The junction characteristics of the FET are accurately modeled for current (density) up to `jmelt`. For current density above `jmelt`, the junction is modeled as a linear resistor and a warning is printed.

Auto Model Selection

Many models need to be characterized for different geometries in order to obtain accurate results for model development. The model selector program automatically searches for a model with the length and width range specified in the instance statement and uses this model in the simulations.

For the auto model selector program to find a specific model, the models to be searched should be grouped together within braces. Such a group is called a model group. An opening brace is required at the end of the line defining each model group. Every model in the group is given a name followed by a colon and the list of parameters. Also, the four geometric parameters `lmax`, `lmin`, `wmax`, and `wmin` should be given. The selection criteria to choose a model is as follows:

$$l_{min} \leq inst_length < l_{max} \quad \text{and} \quad w_{min} \leq inst_width < w_{max}$$

Example:

```
model ModelName ModelType {  
  1: <model parameters> lmin=2 lmax=4 wmin=1 wmax=2  
  2: <model parameters> lmin=1 lmax=2 wmin=2 wmax=4  
  3: <model parameters> lmin=2 lmax=4 wmin=4 wmax=6  
}
```

Then for a given instance

```
M1 1 2 3 4 ModelName w=3 l=1.5
```

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the program would search all the models in the model group with the name `ModelName` and then pick the first model whose geometric range satisfies the selection criteria. In the preceding example, the auto model selector program would choose `ModelName.2`.

You must specify both length (`l`) and width (`w`) on the device instance line to enable automatic model selection.

Output Parameters

1	<code>w_{eff}</code> (m)	Effective channel width.
2	<code>l_{eff}</code> (m)	Effective channel length.
3	<code>r_{th_{eff}}</code> (Ω)	Effective thermal resistance.
4	<code>c_{th_{eff}}</code> (F)	Effective thermal capacitance.
5	<code>r_{seff}</code> (Ω)	Effective source resistance.
6	<code>r_{deff}</code> (Ω)	Effective drain resistance.

Operating-Point Parameters

1	<code>type=n</code>	Transistor type. Possible values are <code>n</code> or <code>p</code> .
2	<code>region=triode</code>	Estimated operating region. Spectre outputs number (0-4) in a rawfile. Possible values are <code>off</code> , <code>triode</code> , <code>sat</code> , or <code>subth</code> .
3	<code>reversed</code>	Reverse mode indicator. Possible values are <code>no</code> or <code>yes</code> .
4	<code>v_{gs}</code> (V)	Gate-source voltage.
5	<code>v_{ds}</code> (V)	Drain-source voltage.
6	<code>v_{bs}</code> (V)	Bulk-source voltage.
7	<code>i_{ds}</code> (A)	Resistive drain-to-source current.
8	<code>i_{sub}</code> (A)	Resistive substrate current.

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Component Statements Part I

9	<code>ibd</code> (A)	Resistive bulk-to-drain junction current.
10	<code>ibs</code> (A)	Resistive bulk-to-source junction current.
11	<code>vth</code> (V)	Threshold voltage.
12	<code>vdsat</code> (V)	Drain-source saturation voltage.
13	<code>gm</code> (S)	Common-source transconductance.
14	<code>gds</code> (S)	Common-source output conductance.
15	<code>gmb</code> (S)	Body-transconductance.
16	<code>ueff</code> ($\text{cm}^2/\text{V s}$)	Effective mobility.
17	<code>betaeff</code> (A/V^2)	Effective beta.
18	<code>cjd</code> (F)	Drain-bulk junction capacitance.
19	<code>cjs</code> (F)	Source-bulk junction capacitance.
20	<code>cgg</code> (F)	dQ_g/dV_g .
21	<code>cgd</code> (F)	dQ_g/dV_d .
22	<code>cgs</code> (F)	dQ_g/dV_s .
23	<code>cgb</code> (F)	dQ_g/dV_{bk} .
24	<code>cdg</code> (F)	dQ_d/dV_g .
25	<code>cdd</code> (F)	dQ_d/dV_d .
26	<code>cds</code> (F)	dQ_d/dV_s .
27	<code>cdb</code> (F)	dQ_d/dV_b .
28	<code>csg</code> (F)	dQ_s/dV_g .
29	<code>csd</code> (F)	dQ_s/dV_d .
30	<code>css</code> (F)	dQ_s/dV_s .

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31	csb (F)	dQs_dVb.
32	cbg (F)	dQb_dVg.
33	cbd (F)	dQb_dVd.
34	cbs (F)	dQb_dVs.
35	cbb (F)	dQb_dVb.
36	ron (Ω)	On-resistance.
37	id (A)	Total resistive drain current.
38	is (A)	Total resistive source current.
39	ib (A)	Total resistive bulk current.
40	pwr (W)	Power at op point.
41	gmoverid (1/V)	Gm/Ids.
42	tdev (C)	Temperature rise from ambient.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the

Spectre Circuit Simulator Reference Component Statements Part I

description for that parameter. For example, a reference of M-35 means the 35th model parameter.

Ctheff	O-4	dice	M-191	ll	M-201	rss	M-89
a0	M-22	dii	M-205	ldif	M-92	rth0	M-199
a0bg	M-195	dlc	M-126	leff	O-2	rtheff	O-3
a0cv	M-216	dqsq	M-215	lint	M-33	selft	M-143
a1	M-25	drout	M-64	ll	M-35	tau	M-123
a2	M-26	dskip	M-98	lln	M-36	tbox	M-48
ad	I-4	dsub	M-79	lmax	M-181	tdev	OP-42
ad	M-135	dt1	M-160	lmin	M-182	tlev	M-144
af	M-173	dt2	M-161	lw	M-37	tlevc	M-145
ags	M-27	dvt0	M-16	lwl	M-39	tmax	M-141
aia	M-202	dvt0w	M-19	lwn	M-38	tnom	M-140
alarm	M-183	dvt1	M-17	m	I-9	tox	M-47
alpha0	M-82	dvt1w	M-20	meto	M-106	trd	M-158
as	I-3	dvt2	M-18	minr	M-93	trise	M-142
as	M-134	dvt2w	M-21	mj	M-113	trise	I-11
at	M-152	dvtr	M-197	mjsw	M-117	trs	M-157
b0	M-23	dvtrbg	M-194	mjswg	M-120	tsi	M-49
b1	M-24	dvtrd	M-192	mobmod	M-58	type	OP-1
b3v3mod	M-2	dvtrg	M-193	n	M-97	type	M-1
beta0	M-83	dwb	M-46	nch	M-30	u0	M-59
betaeff	OP-17	dwc	M-125	ndiode	M-206	ua	M-61
bii	M-203	dwg	M-45	nfactor	M-76	ua1	M-153
binunit	M-57	edl	M-211	ngate	M-31	ub	M-62
btasoiver	M-4	ef	M-174	nlx	M-11	ub1	M-154
bvj	M-185	eg	M-146	noia	M-175	uc	M-63
capmod	M-124	em	M-178	noib	M-176	uc1	M-155
cbb	OP-35	eta0	M-80	noic	M-177	ueff	OP-16
cbd	M-111	etab	M-81	noimod	M-171	ute	M-159
cbd	OP-33	fc	M-115	nrd	I-7	vbgf	M-198

Spectre Circuit Simulator Reference Component Statements Part I

cbg	OP-32	fcsw	M-122	nrd	M-138	vbm	M-15
cbs	M-110	gamma1	M-12	nrs	M-139	vbox	M-186
cbs	OP-34	gamma2	M-13	nrs	I-8	vbs	OP-6
cdb	OP-27	gap1	M-147	nsub	M-29	vbtho	M-188
cdd	OP-25	gap2	M-148	nt	M-207	vbx	M-14
cdg	OP-24	gds	OP-14	pb	M-114	vds	OP-5
cds	OP-26	gm	OP-13	pbsw	M-118	vdsat	OP-12
cdsc	M-73	gmbs	OP-15	pbswg	M-121	version	M-3
cdscb	M-74	gmoverid	OP-41	pclm	M-65	vfbcv	M-130
cdscd	M-75	hdif	M-91	pd	M-137	vgs	OP-4
cf	M-129	ib	OP-39	pd	I-6	voff	M-78
cgb	OP-23	ibd	OP-9	pdiblc1	M-66	vsat	M-60
cgbo	M-105	ibs	OP-10	pdiblc2	M-67	vth	OP-11
cgd	OP-21	id	OP-37	pdiblc3	M-68	vtho	M-5
cgdl	M-108	ids	OP-7	prt	M-156	vtr0	M-189
cgdo	M-104	imax	M-184	prwb	M-52	w	M-132
cgg	OP-20	imelt	M-99	prwg	M-53	w	I-1
cgs	OP-22	imelt1	M-100	ps	M-136	w0	M-10
cgs1	M-107	imelt2	M-101	ps	I-5	warn	M-187
cgso	M-103	imelt3	M-102	pscbe1	M-69	weff	O-1
cii	M-204	is	M-96	pscbe2	M-70	wint	M-34
cit	M-77	is	OP-38	pta	M-167	wl	M-40
cj	M-112	is1	M-208	ptc	M-166	wln	M-41
cjd	OP-18	is2	M-209	ptp	M-168	wmax	M-179
cjs	OP-19	is3	M-210	pvag	M-71	wmin	M-180
cjsw	M-116	isub	OP-8	pwr	OP-40	wr	M-54
cjswg	M-119	js	M-94	qgvd0	M-217	ww	M-42
ckappa	M-109	jsw	M-95	rbody	I-12	ww1	M-44
clc	M-127	k1	M-6	rd	M-86	wwn	M-43
cle	M-128	k2	M-7	rdc	M-88	xj	M-32
csb	OP-31	k3	M-8	rdd	M-90	x1	M-55
csd	OP-29	k3b	M-9	rdef	O-6	xpart	M-131

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csg	OP-28	kb	M-212	rds	M-51	xt	M-50
css	OP-30	keta	M-28	region	OP-2	xti	M-162
cta	M-169	kf	M-172	region	I-10	xti1	M-163
cth0	M-200	knk	M-190	reversed	OP-3	xti2	M-164
ctp	M-170	kt1	M-149	ron	OP-36	xti3	M-165
dbg	M-196	kt11	M-150	rs	M-85	xw	M-56
delacc	M-213	kt2	M-151	rsc	M-87		
delr	M-214	l	M-133	rseff	O-5		
delta	M-72	l	I-2	rsh	M-84		

Component Statements Part II

This chapter discusses the following topics:

- [Two Terminal Capacitor \(capacitor\)](#) on page 282
- [Linear Current Controlled Current Source \(cccs\)](#) on page 286
- [Linear Current Controlled Voltage Source \(ccvs\)](#) on page 288
- [Circuit Reduced Order Model \(cktrom\)](#) on page 290
- [Magnetic Core with Hysteresis \(core\)](#) on page 292
- [Logic-to-Analog Converter \(d2a\)](#) on page 296
- [Delay Line \(delay\)](#) on page 298
- [Diode Level 500 \(dio500\)](#) on page 298
- [Junction Diode \(diode\)](#) on page 302
- [EKV MOSFET Transistor \(ekv\)](#) on page 309
- [Ratiometric Fourier Analyzer \(fourier\)](#) on page 323
- [GaAs MESFET \(gaas\)](#) on page 326
- [Hetero-Junction Bipolar Transistor \(hbt\)](#) on page 331
- [HiSIM1 Field Effect Transistor \(hisim\)](#) on page 341
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- [Two Terminal Inductor \(inductor\)](#) on page 372
- [Interconnect Capacitance \(intcap\)](#) on page 375
- [Current Probe \(iprobe\)](#) on page 379
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Component Statements Part II

- Junction Capacitor (juncap) on page 392
- MISN Field Effect Transistor (misnan) on page 397
- MOS Level-0 Transistor (mos0) on page 405
- MOS Level-1 Transistor (mos1) on page 408
- Compact MOS-Transistor Distortion Model (mos1000) on page 424
- Compact MOS-Transistor Distortion Model (mos1100) on page 435
- Compact MOS-Transistor Distortion Model (mos11010) on page 447
- Compact MOS-Transistor Distortion Model (mos11011) on page 459
- MOS Level-15 Transistor (mos15) on page 474

Two Terminal Capacitor (capacitor)

Description

You can assign the capacitance or let Spectre compute it from the physical length and width of the capacitor. In either case, the capacitance can be a function of temperature or applied voltage.

This device is supported within altergroups.

If the C(inst) is not given,

$$C(\text{inst}) = C(\text{model})$$

if C(model) is given and ,

if Area(inst) or Perim(inst) is given

$$\text{Area_eff} = \text{Area} - (\text{Perim}) \cdot \text{etch} + 4 \cdot \text{etch}^2$$

$$\text{Perim_eff} = \text{Perim} - 8 \cdot \text{etch}$$

else

$$\text{Area_eff} = (\text{L} - 2 \cdot \text{etch}) \cdot (\text{W} - 2 \cdot \text{etch})$$

$$\text{Perim_eff} = 2 \cdot (\text{W} + \text{L} - 4 \cdot \text{etch})$$

$$C(\text{inst}) = C_j \cdot \text{Area_eff} + C_{jsw} \cdot \text{Perim_eff}$$

if C(model) is not given.

If the polynomial coefficients vector (coeffs=[c1 c2 ...]) is specified, the capacitor is nonlinear and the capacitance is

$$\begin{aligned} C(V) &= dQ(V) / dV \\ &= C(\text{inst}) \cdot (1 + c_1 \cdot V + c_2 \cdot V^2 + \dots) \end{aligned}$$

or

$$Q(V) = C(\text{inst}) \cdot V \cdot (1 + 1/2 \cdot c_1 \cdot V + 1/3 \cdot c_2 \cdot V^2 + \dots)$$

where c_k is the k th entry in the coefficient vector.

Spectre Circuit Simulator Reference

Component Statements Part II

The value of the capacitor as a function of the temperature is given by:

$$C(T) = C(tnom) * [1 + tc1 * (T - tnom) + tc2 * (T - tnom)^2].$$

where

$$T = trise(inst) + temp$$

if `trise(inst)` is given, and

$$T = trise(model) + temp$$

if `trise(inst)` is not given.

Sample Instance Statement

Without model:

```
c2 (1 0) capacitor c=2.5u tc1=1e-8
```

With model:

```
c2 (1 0) proc_cap c=2.5u tc1=1e-8
```

Sample Model Statement

```
model proc_cap capacitor c=2u tc1=1.2e-8 tnom=25
```

Instance Definition

```
Name 1 2 ModelName parameter=value ...
```

```
Name 1 2 capacitor parameter=value ...
```

Instance Parameters

1	<code>c</code>	(F)	Capacitance.
2	<code>w</code>	(m)	Capacitor width.
3	<code>l</code>	(m)	Capacitor length.
4	<code>m=1</code>		Multiplicity factor.
5	<code>scale=1</code>		Scale factor.

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Component Statements Part II

6	<code>trise</code> (C)	Temperature rise from ambient.
7	<code>tc1</code> (1/C)	Linear temperature coefficient.
8	<code>tc2</code> (C ⁻²)	Quadratic temperature coefficient.
9	<code>ic</code> (V)	Initial condition.
10	<code>area=1.0</code> m ²	capacitor area.
11	<code>perim=0.0</code> m	capacitor perimeter.

The instance parameter `scale`, if specified, overrides the value given by the `option` parameter `scale`. The `w` and `l` parameters are scaled by the resulting `scale`, and the `option` parameter `scalem`. The values of `w` and `l` printed by Spectre are those given in the input file, and these values might not have the correct units if the scaling factors are not unity. The actual capacitor dimensions are stored in the output parameters. You can obtain these dimensions with the `info` statement.

Model Definition

```
model modelName capacitor parameter=value ...
```

Model Parameters

1	<code>c=0</code> F	Default capacitance.
2	<code>tc1=0</code> 1/C	Linear temperature coefficient.
3	<code>tc2=0</code> C ⁻²	Quadratic temperature coefficient.
4	<code>trise=0</code> C	Default <code>trise</code> value for instance.
5	<code>tnom</code> (C)	Parameters measurement temperature. Default set by <code>options</code> .
6	<code>w=0</code> m	Default capacitor width.
7	<code>l=0</code> m	Default capacitor length.
8	<code>etch=0</code> m	Narrowing due to side etching.
9	<code>cj=0</code> F/m ²	Bottom capacitance density.

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Component Statements Part II

10	<code>cjsw=0</code>	F/m	Sidewall capacitance.
11	<code>scalec=1</code>		Capacitance scaling factor.
12	<code>coeffs=[...]</code>		Vector of polynomial capacitance coefficients.
13	<code>rforce=1</code>	Ω	Resistance used when forcing initial conditions.

Output Parameters

1	<code>leff</code>	(m)	Effective capacitor length.
2	<code>weff</code>	(m)	Effective capacitor width.
3	<code>ceff</code>	(F)	Effective capacitance.

Operating-Point Parameters

1	<code>cap</code>	(F)	Capacitance at operating point.
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Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

<code>area</code>	I-10	<code>coeffs</code>	M-12	<code>perim</code>	I-11	<code>tc2</code>	I-8
<code>c</code>	M-1	<code>etch</code>	M-8	<code>rforce</code>	M-13	<code>tnom</code>	M-5
<code>c</code>	I-1	<code>ic</code>	I-9	<code>scale</code>	I-5	<code>trise</code>	I-6
<code>cap</code>	OP-1	<code>l</code>	I-3	<code>scalec</code>	M-11	<code>trise</code>	M-4
<code>ceff</code>	O-3	<code>l</code>	M-7	<code>tc1</code>	M-2	<code>w</code>	I-2
<code>cj</code>	M-9	<code>leff</code>	O-1	<code>tc1</code>	I-7	<code>w</code>	M-6
<code>cjsw</code>	M-10	<code>m</code>	I-4	<code>tc2</code>	M-3	<code>weff</code>	O-2

Linear Current Controlled Current Source (cccs)

Description

A current-controlled source senses the current with a probe device. A valid probe is a component instance in the circuit that naturally computes current. For example, probes can be voltage sources (independent or controlled), inductors, transmission lines, microstrip lines, N-ports, and transformers. If the probe device computes more than one current (such as transmission lines, microstrip lines, and N-ports), the index of the probe port through which the controlling current flows needs to be specified. Positive current exits the source node and enters the sink node of the controlled source.

This device can also model ideal digital gates.

This device is supported within altergroups.

Sample Instance Statement

```
vcs (pos gnd) cccs gain=2.5 probe=v1 m=1 //Note that v1 is an instance of a voltage source
```

Instance Definition

```
Name sink src cccs parameter=value ...
```

Instance Parameters

1	<code>m=1</code>	Multiplicity factor.
2	<code>probe</code>	Device through which the controlling current flows.
3	<code>port=0</code>	Index of the probe port through which the controlling current flows.
4	<code>probes=[...]</code>	Devices through which the controlling currents flow. For multi-input digital gates only.
5	<code>ports=[...]</code>	Indices of the probe ports through which the controlling currents flow. For multi-input digital gates only.
6	<code>type=cccs</code>	Type of the source. Possible values are <code>cccs</code> , <code>and</code> , <code>nand</code> , <code>or</code> , <code>of</code> or <code>nor</code> .

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Component Statements Part II

7 `delta=0` Smoothing parameter. This may lead to circuit convergency. The smaller the delta is, the sharper the corner is.

Linear source parameters

8 `gain=0 A/A` Current gain.

9 `min (A)` Minimum output current.

10 `max (A)` Maximum output current.

11 `abs=off` Absolute output current.
Possible values are `off` or `on`.

PWL source parameters

12 `file` Name of file containing current/current pairs that define the PWL transfer function.

13 `pwl=[...]` Vector of current/current pairs that defines the PWL transfer function. The format of the vector is [`in1 out1 in2 out2 ...`].

14 `scale=1` Scale factor for the PWL output current.

15 `stretch=1` Scale factor for the PWL controlling current.

Temperature effects parameters

16 `tc1=0 1/C` Linear temperature coefficient.

17 `tc2=0 C-2` Quadratic temperature coefficient.

Operating-Point Parameters

1 `i (A)` Input current.

2 `v (V)` Output voltage.

3 `pwr (W)` Power dissipation.

Spectre Circuit Simulator Reference

Component Statements Part II

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

abs	I-11	m	I-1	probe	I-2	stretch	I-15
delta	I-7	max	I-10	probes	I-4	tc1	I-16
file	I-12	min	I-9	pwl	I-13	tc2	I-17
gain	I-8	port	I-3	pwr	OP-3	type	I-6
i	OP-1	ports	I-5	scale	I-14	v	OP-2

Linear Current Controlled Voltage Source (ccvs)

Description

A current-controlled source senses the current with a probe device. A valid probe is a component instance in the circuit that naturally computes current. For example, probes can be voltage sources (independent or controlled), inductors, transmission lines, microstrip lines, N-ports, and transformers. If the probe device computes more than one current (such as transmission lines, microstrip lines, and N-ports), the index of the probe port through which the controlling current flows needs to be specified. Current through the controlled voltage source is calculated and is defined to be positive if it flows from the positive terminal, through the source, to the negative terminal.

This device can also model ideal digital gates.

This device is supported within altergroups.

Sample Instance Statement

```
vvs (pos gnd) ccvs rm=1 probe=v1 m=1 //Note that v1 is an instance of a voltage source
```

Instance Definition

```
Name p n ccvs parameter=value ...
```

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Component Statements Part II

Instance Parameters

1	<code>m=1</code>	Multiplicity factor.
2	<code>probe</code>	Device through which the controlling current flows.
3	<code>port=0</code>	Index of the probe port through which the controlling current flows.
4	<code>probes=[...]</code>	Devices through which the controlling currents flow. For multi-input digital gates only.
5	<code>ports=[...]</code>	Indices of the probe ports through which the controlling currents flow. For multi-input digital gates only.
6	<code>type=ccvs</code>	Type of the source. Possible values are <code>ccvs</code> , <code>and</code> , <code>nand</code> , <code>or</code> , <code>or</code> or <code>nor</code> .
7	<code>delta=0</code>	Smoothing parameter. This may lead to circuit convergency. The smaller the delta is, the sharper the corner is.

Linear source parameters

8	<code>rm=0 Ω</code>	Transresistance.
9	<code>min (V)</code>	Minimum output voltage.
10	<code>max (V)</code>	Maximum output voltage.
11	<code>abs=off</code>	Absolute output voltage. Possible values are <code>off</code> or <code>on</code> .

PWL source parameters

12	<code>file</code>	Name of file containing current/voltage pairs that define the PWL transfer function.
13	<code>pwl=[...]</code>	Vector of current/voltage pairs that defines the PWL transfer function. The format of the vector is <code>[in1 out1 in2 out2 ...]</code> .
14	<code>scale=1</code>	Scale factor for the PWL output voltage.

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Component Statements Part II

15 `stretch=1` Scale factor for the PWL controlling current.

Temperature effects parameters

16 `tc1=0 1/C` Linear temperature coefficient.

17 `tc2=0 C-2` Quadratic temperature coefficient.

Operating-Point Parameters

1 `i (A)` Output current.

2 `v (V)` Output voltage.

3 `pwr (W)` Power dissipation.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

<code>abs</code>	I-11	<code>max</code>	I-10	<code>probes</code>	I-4	<code>stretch</code>	I-15
<code>delta</code>	I-7	<code>min</code>	I-9	<code>pwl</code>	I-13	<code>tc1</code>	I-16
<code>file</code>	I-12	<code>port</code>	I-3	<code>pwr</code>	OP-3	<code>tc2</code>	I-17
<code>i</code>	OP-1	<code>ports</code>	I-5	<code>rm</code>	I-8	<code>type</code>	I-6
<code>m</code>	I-1	<code>probe</code>	I-2	<code>scale</code>	I-14	<code>v</code>	OP-2

Circuit Reduced Order Model (cktrom)

Description

The circuit reduced order model is described by a set of partial differential equations in the form of:

Spectre Circuit Simulator Reference

Component Statements Part II

$$\dot{x} = Ax + Bu \quad (1)$$

$$y = Cx + Du \quad (2)$$

where Eqn.(1) is the state equation, Eqn.(2) is the output equation, A is nxn matrix, B is nxm, C is mxn, and D is an mxm matrix. x is a vector of state variables. Input u is a vector of voltages at all the ports and output y is a vector of electric current at all the ports. The number of inputs is always equal to the number of outputs. The order of the terminals in the input must be consistent with the matrix equations. In the input file, the matrices A, B, C and D are in the form of long vectors with row order.

This device is not supported within altergroup.

Sample Instance Statement

```
rom3 (net11 0) cktrom a=[ -2.022852e+14  2.583012e+13  9.553125e+13  9.627727e+13
1.533971e+13  9.987851e+13  4.592012e+13  -1.671024e+14  2.296589e+13  -2.719915e+14
7.668472e+12  -1.564519e+14  8.543123e+13  3.395689e+13  -3.863150e+14  -1.101618e+14
-5.415116e+14  -2.303841e+14  9.627728e+13  -1.711915e+14  -1.001818e+14  -
8.123120e+14  -2.272715e+14  -9.965181e+14  1.514961e+14  7.668372e+12  -6.415116e+14
-3.272715e+14  -2.852751e+15  -3.564466e+14  9.999851e+13  -1.761619e+14  -1.312841e+14
-8.967181e+14  -4.563456e+14  -4.068747e+15 ]
b=[3.366776e+06  5.932470e+05  -1.508475e+06  4.349182e+06  -3.128869e+06  -
2.995677e+06  -2.831481e+06  2.708942e+06  -4.968876e+06  -3.338945e+06  -3.278564e+06
3.925648e+06 ]
c=[-3.111296e+06  1.593292e+06  3.324594e+06  3.083731e+06  5.887179e+06
3.766094e+06  -5.049263e+05  -4.275158e+06  3.035578e+06  -3.666385e+06  3.424639e+06
-3.832285e+06]
d=[1.254627e-01  0.000000e+00  0.000000e+00  1.236790e-01 ]
```

Instance Definition

```
Name sink0 src0 [sink1] [src1] [sink2] [src2] [sink3] [src3] ... cktrom
parameter=value ...
```

Instance Parameters

- 1 m=1 Multiplicity factor.
- 2 a=[...] Coefficient matrix A of state equations.
- 3 b=[...] Coefficient Matrix B of state equations.
- 4 c=[...] Coefficient matrix C of output equations.

Spectre Circuit Simulator Reference

Component Statements Part II

5 $d=[\dots]$ Coefficient matrix D of output equations.

Operating-Point Parameters

1 $i=[\dots]$ A Port currents.
2 $v=[\dots]$ V Port voltages.
3 pwr (W) Power dissipation.

Parameter Index

In the following index, I refers to instance parameters, M refers to the model parameters section, O refers to the output parameters section, and OP refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of M-35 means the 35th model parameter.

a	I-2	c	I-4	i	OP-1	pwr	OP-3
b	I-3	d	I-5	m	I-1	v	OP-2

Magnetic Core with Hysteresis (core)

Description

This component models the magnetic hysteresis, with air gap, frequency, and temperature effects. The model is based on the AWB model for magnetic cores and windings. The user has to specify the cores material and geometric parameters to model the hysteresis.

The material parameters to specify are the B_r , B_m and H_c of the core. The geometric parameters are the area, magnetic path length and the air gap of the core.

You can specify the magnetic path length in one of the following ways:

Give the length directly in cm.

Or give the outer and inner diameter of the core.

Spectre Circuit Simulator Reference

Component Statements Part II

Cores without terminals represent complete magnetic loops. Cores with terminals are fragments that you can use as building blocks to build models of complicated core structures. For example, you can use the following set of core fragments to model an E core:

```
W1 e1p e1m winding turns=80 core=C1
```

```
W3a e2p e2c winding turns=80 core=C3
```

```
W3b e2c e2m winding turns=80 core=C3
```

```
C1 m1 0 permalloy area=1 len=2
```

```
C2 m1 0 permalloy area=2 len=2
```

```
C3 m1 0 permalloy area=1 len=2
```

```
model permalloy core ...
```

There are three parallel core fragments representing each of the three fingers on the E. One 80 turn winding is connected to core fragment C1. A center-tapped 160 turn winding (implemented as a pair of windings) are wrapped around core fragment C3. Node m1 is a magnetic node whose value is in magnetomotive force and flow is flux.

You can calculate the frequency and temperature dependency of the core model by specifying the frequency loss parameters and the temperature effects parameters. You can make all the core parameters vary in temperature, including the permeability, saturation flux, and core loss. For frequency losses, a static model refers to a value that you type in for frequency. This model does not adjust the shape of the B-H loop in response to power dissipation or rate of rise of the applied currents and voltages during transient analysis.

This device is not supported within altergroup.

The hysteresis is modeled by different regions whose equations are:

$$\phi = \phi_{ir} + (\phi_{is} - \phi_{ir}) F / (F + H_a) \quad \text{for region number 1}$$

$$\phi = \phi_{is} * (F - F_c) / (F - H_b) \quad \text{for region number 2}$$

where ϕ = flux density

F = magnetomotive force

ϕ_{ir} = residual flux density

ϕ_{is} = Saturated flux density

Spectre Circuit Simulator Reference

Component Statements Part II

F_c = Coercive magnetic force

H_a and H_b are shape parameters.

Sample Instance Statement

```
c1 (1 0) core_mod area=1.2 len=8.1 id=0.55 gap=0.25
```

Sample Model Statement

```
model core_mod core len=7.7 area=0.85 br=1e3 bm=5e3 hc_t1=0.2 p1_f1=2.08 f1=10e3  
p2_f2=50 f2=100K bflux=1e3 density=4.75
```

Instance Definition

Name ... ModelName parameter=value ...

Instance Parameters

1	area (cm ²)	Effective magnetic cross-sectional area of core.
2	len (cm)	Effective length of magnetic path.
3	id (cm)	Inner diameter of toroidal core.
4	od (cm)	Outer diameter of toroidal core.
5	gap (cm)	Gap length.
6	m=1	Multiplicity factor.

Model Definition

```
model modelName core parameter=value ...
```

Model Parameters

1	br=1 gauss	Residual flux density.
2	bm=1 gauss	Saturation flux density.
3	hc=1 oersteds	Coercive magnetizing force (value of H where B equals 0).

Spectre Circuit Simulator Reference

Component Statements Part II

4	area=1 cm ²	Effective magnetic cross-sectional area of core.
5	len=1 cm	Effective length of magnetic path.
6	id (cm)	Inner diameter of toroidal core.
7	od (cm)	Outer diameter of toroidal core.
8	gap=0.0 cm	Gap length.

Initial Conditions

9	b0 (gauss)	Initial condition for core.
---	------------	-----------------------------

Frequency Loss Parameters

10	freq (Hz)	Core operating frequency.
11	p1_f1 (W/Kg)	Core power loss at frequency f1.
12	f1 (Hz)	Reference frequency for power loss.
13	p2_f2 (W/Kg)	Core power loss at frequency f2.
14	f2 (Hz)	Reference frequency for power loss.
15	bflux (gauss)	Reference flux density.
16	density (g/cm ³)	Core density.

Temperature Effects Parameters

17	temp (C)	Core operating temperature.
18	bm_t1 (gauss)	Saturated flux density B _m at T1.
19	br_t1 (gauss)	Residual flux density B _r at T1.
20	hc_t1 (oersteds)	Coercive force H _c at T1.
21	t1 (C)	Reference temperature.

Spectre Circuit Simulator Reference

Component Statements Part II

Operating-Point Parameters

- 1 `b` (`gauss`) Flux density of the core.
- 2 `h` (`oersteds`) Magnetic field strength.

Parameter Index

In the following index, `I` refers to instance parameters, `M` refers to the model parameters section, `O` refers to the output parameters section, and `OP` refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of `M-35` means the 35th model parameter.

<code>area</code>	<code>I-1</code>	<code>br_t1</code>	<code>M-19</code>	<code>hc</code>	<code>M-3</code>	<code>od</code>	<code>I-4</code>
<code>area</code>	<code>M-4</code>	<code>density</code>	<code>M-16</code>	<code>hc_t1</code>	<code>M-20</code>	<code>p1_f1</code>	<code>M-11</code>
<code>b</code>	<code>OP-1</code>	<code>f1</code>	<code>M-12</code>	<code>id</code>	<code>I-3</code>	<code>p2_f2</code>	<code>M-13</code>
<code>b0</code>	<code>M-9</code>	<code>f2</code>	<code>M-14</code>	<code>id</code>	<code>M-6</code>	<code>t1</code>	<code>M-21</code>
<code>bflux</code>	<code>M-15</code>	<code>freq</code>	<code>M-10</code>	<code>len</code>	<code>M-5</code>	<code>temp</code>	<code>M-17</code>
<code>bm</code>	<code>M-2</code>	<code>gap</code>	<code>I-5</code>	<code>len</code>	<code>I-2</code>		
<code>bm_t1</code>	<code>M-18</code>	<code>gap</code>	<code>M-8</code>	<code>m</code>	<code>I-6</code>		
<code>br</code>	<code>M-1</code>	<code>h</code>	<code>OP-2</code>	<code>od</code>	<code>M-7</code>		

Logic-to-Analog Converter (d2a)

Description

The logic-to-analog converter converts a binary signal from a logic simulator to an analog waveform.

This device is not supported within altergroup.

Sample Instance Statement

```
d2a_1 (net1 net2) d2a src="99991" val0=0 val1=2.5 valx=1.25 rise=200p fall=200p m=2
//99991 is an analog net
```

Spectre Circuit Simulator Reference

Component Statements Part II

Instance Definition

Name p n d2a parameter=value ...

Instance Parameters

1	src	The foreign simulators name for the source of the analog signal.
2	nestlev=0	Number of nesting levels to ignore in the hierarchical name. This should be used skip over extra levels that do not exist in the co-simulator.
3	val0=0 V	Final value for logical 0.
4	val1=5 V	Final value for logical 1.
5	valx (V)	Final value for logical X.
6	valz (V)	Final value for logical Z.
7	rise=1ns s	Time for transition from val0 to val1.
8	fall=1ns s	Time for transition from val1 to val0.
9	ron=100 Ω	Output resistance when in active state.
10	m=1	Multiplicity factor.

Parameter Index

In the following index, I refers to instance parameters, M refers to the model parameters section, O refers to the output parameters section, and OP refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of M-35 means the 35th model parameter.

fall	I-8	rise	I-7	val0	I-3	valz	I-6
m	I-10	ron	I-9	val1	I-4		
nestlev	I-2	src	I-1	valx	I-5		

Delay Line (delay)

Description

The delay line model is a four terminal device with zero output impedance and infinite input impedance. The output between nodes *p* and *n* is the input voltage between nodes *ps* and *ns* delayed by the time delay *td* and scaled by *gain*

This device is not supported within altergroup.

Sample Instance Statement

```
dll(outp outn cntrlp cntrln) delay td=10n gain=1.5
```

Instance Definition

```
Name p n ps ns delay parameter=value ...
```

Instance Parameters

- | | | |
|---|------------------|----------------------|
| 1 | <i>td</i> =0.0 s | Time delay. |
| 2 | <i>gain</i> =1 | Gain parameter. |
| 3 | <i>m</i> =1 | Multiplicity factor. |

Operating-Point Parameters

- | | | |
|---|--------------|-----------------|
| 1 | <i>v</i> (V) | Output voltage. |
|---|--------------|-----------------|

Diode Level 500 (dio500)

Description

The *dio500* model provides a detailed description of the diode currents in forward and reverse biased Si-diodes. It is described in the Philips Bipolar Modelbook (Dec.93) as *Diode level 500*. Information on how to obtain this document can be found on Source Link by searching for Philips.

Spectre Circuit Simulator Reference

Component Statements Part II

(c) Philips Electronics N.V. 1994

In extension to the modelbook description a minimum conductance g_{min} is inserted between the diode nodes to aid convergence. The value of g_{min} is set by an options statement, default is $g_{min} = 1.0e-12$ S .

The i_{max} parameter is used to aid convergence and to prevent numerical overflow. The junction characteristics of the diode are accurately modeled for currents up to i_{max} . For currents above i_{max} , the junction is modeled as a linear resistor, and a warning is printed.

This device is supported within altergroups.

This device is dynamically loaded from the shared object /vobs/spectre_dev/tools.sun4v/spectre/lib/cmi/3.0.doc/libphilips_sh.so

Sample Instance Statement

```
d1 (pnode 0) phdiode area=2
```

Sample Model Statement

```
model phdiode dio500 is=3.5e-12 rs=26.3 n=2.7 imax=1e20 vlc=1.8 vbr=9.63 cj=2.65e-11 dta=12.88 tau=7.5e-10 tnom=25
```

Instance Definition

```
Name a k ModelName parameter=value ...
```

Instance Parameters

- | | | |
|---|------------|--|
| 1 | area=1.0 | Multiplication factor. |
| 2 | mult | Alias of area factor. |
| 3 | m=1.0 | Multiplicity factor. |
| 4 | region=fwd | Estimated DC operating region, used as a convergence aid. Possible values are fwd, rev or brk. |

Model Definition

```
model modelName dio500 parameter=value ...
```

Spectre Circuit Simulator Reference

Component Statements Part II

Model Parameters

1	$i_s=7.13e-13$ A	Saturation current.
2	$n=1.044$	Junction emission coefficient.
3	$v_{lc}=0.0$ V	Voltage dependence at low forward currents.
4	$v_{br}=7.459$ V	Breakdown voltage.
5	$emv_{br}=1.36e+06$ V/cm	Electric field at breakdown.
6	$csr_h=7.44e-07$ A/cm	Shockley-Read-Hall generation.
7	$cb_{bt}=3.255$ A/V	Band to band tunneling.
8	$ct_{at}=3.31e-06$ A/cm	Trap assisted tunneling.
9	$r_s=0.0$ Ω	Series resistance.
10	$\tau=500.0e-12$ s	Transit time.
11	$c_j=7.0e-12$ F	Zero-bias depletion capacitance.
12	$v_d=0.9$ V	Diffusion voltage.
13	$p=0.4$	Grading coefficient.
14	t_{ref} (C)	Reference temperature. Default set by option t_{nom} .
15	t_{nom} (C)	Alias of t_{ref} .
16	t_r (C)	Alias of t_{ref} .
17	$v_g=1.206$ V	Bandgap voltage.
18	$p_{trs}=0.0$	Power for temperature dependence of r_s .
19	$k_f=0.0$	Flickernoise coefficient.

Spectre Circuit Simulator Reference

Component Statements Part II

20	af=1.0	Flickernoise exponent.
21	dta=0.0 K	Difference between device temperature and ambient temperature.
22	trise (K)	Alias of dta.
23	imax=1.0 A	Explosion current.

Operating-Point Parameters

1	vak (V)	Diode voltage, measured from anode to cathode (including r_s).
2	id (A)	Total resistive diode current.
3	qd (Coul)	Diffusion charge.
4	qt (Coul)	Depletion charge.
5	rst (Ω)	Series resistance (temperature updated).
6	r1 (Ω)	AC linearized resistance.
7	c1 (F)	AC linearized capacitance.
8	pwr (W)	Power dissipation.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

af	M-20	id	OP-2	pwr	OP-8	tr	M-16
area	I-1	imax	M-23	qd	OP-3	tref	M-14
cbbt	M-7	is	M-1	qt	OP-4	trise	M-22
cj	M-11	kf	M-19	region	I-4	vak	OP-1
c1	OP-7	m	I-3	r1	OP-6	vbr	M-4

Spectre Circuit Simulator Reference Component Statements Part II

csr _h	M-6	mult	I-2	rs	M-9	vd	M-12
ctat	M-8	n	M-2	rst	OP-5	vg	M-17
dta	M-21	p	M-13	tau	M-10	vlc	M-3
emvbr	M-5	ptrs	M-18	tnom	M-15		

Junction Diode (diode)

Description

The junction diode model includes nonlinear junction capacitance and reverse breakdown.

This device is supported within altergroups.

Sample Instance Statement

```
d0 (dp dn) pdiode l=3e-4 w=2.5e-4 area=1
```

Sample Model Statement

```
model pdiode diode is=1.8e-5 rs=1.43 n=1.22 nz=2.31 gleak=6.2e-5 rsw=10 isw=6.1e-10  
ibv=0.95e-3 tgs=2 ik=1.2e7 fc=0.5 cj=1.43e-3 pb=0.967 mj=0.337 cjsw=2.76e-9  
vjsw=0.94 jmax=1e20
```

Instance Definition

```
Name a c ModelName parameter=value ...
```

In forward operation, the voltage on the anode (a) is more positive than the voltage on the cathode (c).

Instance Parameters

- | | | |
|---|----------|----------------------------|
| 1 | area | Junction area factor. |
| 2 | perim | Junction perimeter factor. |
| 3 | l=1e-6 m | Drawn length of junction. |
| 4 | w=1e-6 m | Drawn width of junction. |

Spectre Circuit Simulator Reference

Component Statements Part II

5	<code>m=1</code>	Multiplicity factor.
6	<code>scale=1</code>	Scale factor.
7	<code>region=on</code>	Estimated operating region. Spectre outputs number (0-2) in a rawfile. Possible values are <code>off</code> , <code>on</code> or <code>breakdown</code> .
8	<code>trise (C)</code>	Temperature rise from ambient.
9	<code>lm=0.0 m</code>	Length of metal capacitor.
10	<code>lp=0.0 m</code>	Length of polysilicon capacitor.
11	<code>wm=0.0 m</code>	Width of metal capacitor.
12	<code>wp=0.0 m</code>	Width of polysilicon capacitor.

The instance parameter `scale`, if specified, overrides the value given by the `option` parameter `scale`. If the model parameter `allow_scaling` is set to `yes` then, the `area`, `perim`, `l` and `w` parameters are scaled by `scale`. By default `allow_scaling` is set to `no` and no scaling of geometry parameters will occur. The values of `area`, `perim`, `l` and `w` printed out by spectre are those given in the input, and these values might not have the correct units if the scaling factors are not unity.

Model Definition

```
model modelName diode parameter=value ...
```

Model Parameters

Model selector parameters

1	<code>level=1</code>	Model selector. 1 = Junction ,2 = Fowler-Nordheim, 3 = Junction + additional metal and polysilicon capacitances.
2	<code>hcomp=0</code>	junction equations selector: 0 - Spectre equations 1-Hspice equations (level 3 only).
3	<code>dcap=1</code>	depletion capacitance equations selector (with level 3 and hspice compatibility flag only).

Spectre Circuit Simulator Reference

Component Statements Part II

Process parameters

4	<code>etch=0 m</code>	Narrowing due to etching per side.
5	<code>etchl=etch m</code>	Length reduction due to etching per side.
6	<code>shrink=1.0</code>	Shrink factor (level = 3 only).
7	<code>l=1e-6 m</code>	Drawn length of junction.
8	<code>w=1e-6 m</code>	Drawn width of junction.

Junction diode parameters

9	<code>js=1e-14 A</code>	Saturation current (*area).
10	<code>jsw=0 A</code>	Sidewall saturation current (*perim).
11	<code>n=1</code>	Emission coefficient.
12	<code>ns=1</code>	Sidewall emission coefficient.
13	<code>ik=∞ A</code>	High-level injection knee current (*area).
14	<code>ikp=ik A</code>	High-level injection knee current for sidewall (*area).
15	<code>ikr=∞ A</code>	Reverse high-level injection knee current (*area) (with level=3 and hspice compatibility flag only).
16	<code>area=1</code>	Junction area factor.
17	<code>perim=0</code>	Junction perimeter factor.
18	<code>allow_scaling=no</code>	Allow scale option and instance scale parameter to affect diode instance geometry parameters. Possible values are <code>no</code> or <code>yes</code> .

Capacitive parameters

19	<code>tt=0 s</code>	Transit time.
20	<code>cd=0 F</code>	Linear capacitance (*area).

Spectre Circuit Simulator Reference

Component Statements Part II

21	<code>cj0=0 F</code>	Zero-bias junction capacitance (*area).
22	<code>vj=1 V</code>	Junction potential.
23	<code>pb (V)</code>	Alias to <code>vj</code> .
24	<code>m=0.5</code>	Grading coefficient.
25	<code>cjsw=0 F</code>	Zero-bias sidewall junction capacitance (*perim).
26	<code>vjsw=1 V</code>	Sidewall junction potential.
27	<code>mjsw=0.33</code>	Sidewall grading coefficient.
28	<code>fc=0.5</code>	Forward-bias depletion capacitance threshold.
29	<code>fcsw=fc</code>	Coefficient for forward-bias depletion sidewall capacitance .
30	<code>lm=0.0 m</code>	Length of metal capacitor (level=3 only).
31	<code>lp=0.0 m</code>	Length of polysilicon capacitor(level=3 only).
32	<code>wm=0.0 m</code>	Width of metal capacitor(level=3 only).
33	<code>wp=0.0 m</code>	Width of polysilicon capacitor(level=3 only).
34	<code>xm=0.0 m</code>	XM accounts for masking and etching effects(level=3 only).
35	<code>xp=0.0 m</code>	XP accounts for masking and etching effects(level=3 only).
36	<code>xoi=0.0</code>	Thickness of the polysilicon to bulk oxide. Units - nAngstrom (level=3 only).
37	<code>xom=0.0 m</code>	Thickness of the metal to bulk oxide. Units - nAngstrom (level=3 only).
38	<code>xw=0.0 m</code>	Accounts for masking and etching effects(level=3 only).

Breakdown parameters

39	<code>bv=∞ V</code>	Reverse breakdown voltage. Note: <code>bv=0</code> is not the same as <code>bv=infinity</code> .
----	---------------------	--

Spectre Circuit Simulator Reference

Component Statements Part II

40	$v_b = \infty$ V	Alias to b_v .
41	$i_{bv} = 0.001$ A	Current at breakdown voltage (*area).
42	$n_z = 1$	Emission coefficient for Zener diode.
43	$b_{vj} = \infty$ V	Voltage at which junction breakdown warning is issued.

Parasitic resistance parameters

44	$r_s = 0$ Ω	Series resistance (/area).
45	$r_{sw} = 0$ Ω	Sidewall series resistance (/perim).
46	$g_{leak} = 0$ S	Bottom junction leakage conductance (*area).
47	$g_{leaksw} = 0$ S	Sidewall junction leakage conductance (*perim).
48	$minr = 0.1$ Ω	Minimum series resistance.

Temperature effects parameters

49	$t_{lev} = 0$	DC temperature selector.
50	$t_{levc} = 0$	AC temperature selector.
51	$eg = 1.124481$ V	Band gap. Note: when not specified, the default value is temperature dependent. It is 1.124481 at $temp = 27C$.
52	$gap1 = 7.02e-4$ V/C	Band gap temperature coefficient.
53	$gap2 = 1108$ C	Band gap temperature offset.
54	$x_{ti} = 3$	Saturation current temperature exponent.
55	$tbv1 = 0$ 1/C	Linear temperature coefficient for b_v .
56	$tbv2 = 0$ C ⁻²	Quadratic temperature coefficient for b_v .
57	t_{nom} (C)	Parameters measurement temperature. Default set by <code>options</code> .
58	$trise = 0$ C	Temperature rise from ambient.

Spectre Circuit Simulator Reference

Component Statements Part II

59	<code>trs=0</code>	$1/C$	Linear temperature coefficient for parasitic resistance.
60	<code>trs2=0</code>	C^{-2}	Quadratic temperature coefficient for parasitic resistance.
61	<code>tgs=0</code>	$1/C$	Linear temperature coefficient for leakage conductance.
62	<code>tgs2=0</code>	C^{-2}	Quadratic temperature coefficient for leakage conductance.
63	<code>cta=0</code>	$1/C$	Junction capacitance temperature coefficient.
64	<code>ctp=0</code>	$1/C$	Sidewall junction capacitance temperature coefficient.
65	<code>pta=0</code>	V/C	Junction potential temperature coefficient.
66	<code>ptp=0</code>	V/C	Sidewall junction potential temperature coefficient.

Junction diode model control parameters

67	<code>jmelt=jmax</code>	A	Explosion current (*area).
68	<code>jmax=1</code>	A	Maximum allowable current (*area).
69	<code>dskip=yes</code>		Use simple piece-wise linear model for diode currents below $0.1 \cdot i_{abstol}$. Possible values are <code>no</code> or <code>yes</code> .

Fowler-Nordheim diode parameters

70	<code>if=1e-10</code>	A/V^{nf}	Forward Fowler-Nordheim current coefficient (*area).
71	<code>ir=if</code>	A/V^{nr}	Reverse Fowler-Nordheim current coefficient (*area).
72	<code>ecrf=2.55e10</code>	V/m	Forward critical field.
73	<code>ecrr=ecrf</code>	V/m	Reverse critical field.
74	<code>nf=2</code>		Forward voltage power.
75	<code>nr=nf</code>		Reverse voltage power.
76	<code>tox=1e-8</code>	m	Thickness of insulating layer.

Spectre Circuit Simulator Reference

Component Statements Part II

Noise model parameters

77 `kf=0` Flicker noise (1/f) coefficient.

78 `af=1` Flicker noise (1/f) exponent.

Both of these parameters have current density counterparts, `jmax` and `jmelt`, that you can specify if you want the absolute current values to depend on the device area.

Operating-Point Parameters

1 `region=on` Estimated operating region. Spectre outputs number (0-2) in a rawfile.
Possible values are `off`, `on` or `breakdown`.

2 `v (V)` Extrinsic diode voltage.

3 `i (A)` Resistive diode current.

4 `pwr (W)` Power dissipation.

5 `res (Ω)` Resistance of intrinsic diode.

6 `cap (F)` Junction capacitance.

7 `resp (Ω)` Resistance of intrinsic sidewall diode.

8 `capp (F)` Sidewall junction capacitance.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

<code>af</code>	M-78	<code>gleaksw</code>	M-47	<code>n</code>	M-11	<code>tnom</code>	M-57
<code>allow_scaling</code>	M-18	<code>hcomp</code>	M-2	<code>nf</code>	M-74	<code>tox</code>	M-76

Spectre Circuit Simulator Reference Component Statements Part II

area	I-1	i	OP-3	nr	M-75	trise	M-58
area	M-16	ibv	M-41	ns	M-12	trise	I-8
bv	M-39	if	M-70	nz	M-42	trs	M-59
bvj	M-43	ik	M-13	pb	M-23	trs2	M-60
cap	OP-6	ikp	M-14	perim	I-2	tt	M-19
capp	OP-8	ikr	M-15	perim	M-17	v	OP-2
cd	M-20	ir	M-71	pta	M-65	vb	M-40
cjo	M-21	jmax	M-68	ptp	M-66	vj	M-22
cjsw	M-25	jmelt	M-67	pwr	OP-4	vjsw	M-26
cta	M-63	js	M-9	region	OP-1	w	I-4
ctp	M-64	jsw	M-10	region	I-7	w	M-8
dcap	M-3	kf	M-77	res	OP-5	wm	M-32
dskip	M-69	l	M-7	resp	OP-7	wm	I-11
ecrf	M-72	l	I-3	rs	M-44	wp	I-12
ecrr	M-73	level	M-1	rsw	M-45	wp	M-33
eg	M-51	lm	M-30	scale	I-6	xm	M-34
etch	M-4	lm	I-9	shrink	M-6	xoi	M-36
etch1	M-5	lp	M-31	tbv1	M-55	xom	M-37
fc	M-28	lp	I-10	tbv2	M-56	xp	M-35
fcs	M-29	m	M-24	tgs	M-61	xti	M-54
gap1	M-52	m	I-5	tgs2	M-62	xw	M-38
gap2	M-53	minr	M-48	tlev	M-49		
gleak	M-46	mjsw	M-27	tlevc	M-50		

EKV MOSFET Transistor (ekv)

Description

The EPFL-EKV mosfet model was developed by Electronics Laboratories, Swiss Federal Institute of Technology (EPFL), Switzerland. The detailed description of the model and equations can be found in the Affirma Spectre Circuit Simulator Device Model Equations manual. EKV transistors require that you use a model statement.

Spectre Circuit Simulator Reference

Component Statements Part II

This device is supported within altergroups.

Sample Instance Statement

```
mn1 (dn gn sn 0) ekvnmos w=1.5u l=1u ad=2.6p as=2.6p pd=6.6p ps=6.6p nrd=1.54  
nrs=1.54
```

Sample Model Statement

```
model ekvnmos ekv type=n update=2.6 xqc=0.4 cox=3.4e-3 xj=0.145e-6 vto=0.6  
gamma=0.71 phi=0.967 kp=155e-6 e0=88e6 iba=200e6 ibb=350e6 tnom=25 tcv=1.55e-3  
bex=-1.45 kf=1e-27 af=1 hdif=0.94e-6 rsh=512 jsw=1.5e-10
```

Instance Definition

Name d g s b ModelName parameter=value ...

Instance Parameters

1	w (m)	Channel width.
2	l (m)	Channel length.
3	as (m ²)	Area of source diffusion.
4	ad (m ²)	Area of drain diffusion.
5	ps (m)	Perimeter of source diffusion.
6	pd (m)	Perimeter of drain diffusion.
7	nrd (m/m)	Number of squares of drain diffusion.
8	nrs (m/m)	Number of squares of source diffusion.
9	rdc (Ω)	Drain contact resistance.
10	rsc (Ω)	Source contact resistance.
11	m=1	Multiplicity factor (number of MOSFETs in parallel).
12	ns=1	Series Multiplicity factor (number of MOSFETs in series).

Spectre Circuit Simulator Reference

Component Statements Part II

- 13 `region=triode` Estimated operating region. Spectre outputs number (0-3) in a rawfile.
Possible values are `off`, `triode`, `sat`, or `subth`.
- 14 `trise` Temperature rise from ambient.

Model Definition

`model modelName ekv parameter=value ...`

Model Parameters

Device type parameters

- 1 `type=n` Transistor type.
Possible values are `n` or `p`.

Process parameters

- 2 `tox=2e-8 m` Gate oxide thickness.
- 3 `cox=7e-4 F/m2` Gate oxide capacitance. (Overrides `Tox`).
- 4 `xj=1.0e-7 m` Metallurgical junction depth.
- 5 `dw=0 m` Channel Width Correction.
- 6 `dl=0 m` Channel Length Correction.
- 7 `nfs=0 cm-2` Fast surface state density.
- 8 `nsub=1.13e16 cm-3` Channel doping concentration.

Drain current model parameters

- 9 `vto=0.5 V` Threshold voltage at zero body bias.
- 10 `gamma=1.0 \sqrt{V}` Body-effect parameter.
- 11 `phi=0.7 V` Surface potential at strong inversion.

Spectre Circuit Simulator Reference

Component Statements Part II

12	$k_p=5.0e-5 \text{ A/V}^2$	Transconductance parameter.
13	$e_0=1.0e12 \text{ V/m}$	Vertical Critical Field.
14	$u_{crit}=2.0e6 \text{ V/cm}$	Longitudinal Critical field for mobility degradation.
15	$\theta=0.0 \text{ 1/V}$	Mobility reduction coefficient.
16	$u_0 \text{ (cm}^2/\text{V s)}$	Carrier surface mobility.
17	$v_{max} \text{ (m/s)}$	Carrier saturation velocity.
18	$v_{fb} \text{ (V)}$	Flat-band voltage.
19	$\lambda=0.5$	Channel length modulation parameter.
20	$\eta=0.25$	Narrow Channel Effect Coefficient.
21	$\epsilon=0.1$	Short Channel Effect Coefficient.
22	$x_w=0 \text{ m}$	Width variation due to masking and etching.
23	$x_l=0 \text{ m}$	Length variation due to masking and etching.
24	$m_{eto}=0 \text{ m}$	Metal overlap in fringing field.

Impact ionization parameters

25	$i_{ba}=0 \text{ 1/m}$	First Impact Ionization Coefficient.
26	$i_{bb}=3.0e8 \text{ V/m}$	Second Impact Ionization Coefficient.
27	$i_{bc}=0$	Third Impact Ionization Coefficient.
28	$i_{bn}=1.0$	Saturation velocity factor for impact ionization.

Reverse Short Channel parameters

29	$q_0=0 \text{ A s/m}^2$	Reverse short channel peak charge density.
30	$l_k=2.9e-7 \text{ m}$	Reverse short channel characteristic length.

Spectre Circuit Simulator Reference

Component Statements Part II

Charge model selection parameters

31 `xqc=0.0` Drain/source channel charge partition in saturation for charge models, e.g. use 0.4 for 40/60, 0.5 for 50/50, 0 for 0/100.

Junction diode model parameters

32 `is=1e-14 A` Bulk junction reverse saturation current.

33 `js (A/m2)` Bulk junction reverse saturation current density.

34 `jsw=0 A/m` Bulk junction reverse saturation sidewall current density.

35 `n=1` Junction emission coefficient.

36 `dskip=yes` Use simple piece-wise linear model for diode currents below $0.1 \cdot i_{abstol}$.
Possible values are `no` or `yes`.

37 `imelt='imaxA'` Explosion current, diode is linearized beyond this current to aid convergence.

Junction capacitance model parameters

38 `cbd=0 F` Bulk-drain zero-bias p-n capacitance.

39 `cbs=0 F` Bulk-source zero-bias p-n capacitance.

40 `cj=0 F/m2` Zero-bias junction bottom capacitance density.

41 `cjsw=0 F/m` Zero-bias junction sidewall capacitance density.

42 `mj=0.5` Bulk junction bottom grading coefficient.

43 `mjsw=0.33` Bulk junction sidewall grading coefficient.

44 `cjswg=0 F/m` Gate-side zero-bias junction sidewall capacitance density.

45 `mjswg=0.33` Gate-side bulk junction sidewall grading coefficient.

46 `pbswg=0.8 V` Gate-side junction built-in potential.

Spectre Circuit Simulator Reference

Component Statements Part II

47	$f_c=0.5$	Forward-bias capacitance coefficient.
48	$p_b=0.8$ V	Bulk p-n bottom contact potential.
49	$p_{bsw}=0.8$ V	Side-wall contact potential.
50	$t_t=0.0$ V	Bulk p-n transit time.
51	$f_{csw}=0.5$	Side-wall forward-bias depletion capacitance threshold.

Overlap capacitance parameters

52	$c_{gso}=0$ F/m	Gate-source overlap capacitance.
53	$c_{gdo}=0$ F/m	Gate-drain overlap capacitance.
54	$c_{gbo}=0$ F/m	Gate-bulk overlap capacitance.

Parasitic resistance parameters

55	$r_s=0$ Ω	Source resistance.
56	$r_d=0$ Ω	Drain resistance.
57	$r_{sh}=0$ Ω/sqr	Source/drain diffusion sheet resistance.
58	$r_{ss}=0$ Ω m	Scalable source resistance.
59	$r_{dd}=0$ Ω m	Scalable drain resistance.
60	$r_{sc}=0$ Ω	Source contact resistance.
61	$r_{dc}=0$ Ω	Drain contact resistance.
62	$minr=0.1$ Ω	Minimum source/drain resistance.
63	$ldif=0$ m	Lateral diffusion beyond the gate.
64	$hdif=0$ m	Length of heavily doped diffusion.

Spectre Circuit Simulator Reference

Component Statements Part II

Short distance matching parameters

- 65 `avto=0` V m Area related threshold voltage mismatch parameter.
- 66 `akp=0` m Area related gain mismatch parameter.
- 67 `agamma=0` $\sqrt{\text{V}}$ m Area related body effect mismatch parameter.

Operating region warning control parameters

- 68 `alarm=none` Forbidden operating region.
Possible values are `none`, `off`, `triode`, `sat`, `subth`, or `rev`.
- 69 `imax=1` A Maximum current, currents above this limit generate a warning.
- 70 `jmax=1e8` A/m² Maximum current density, currents above this limit generate a warning.
- 71 `vbox=1e9` `tox` V Oxide breakdown voltage.
- 72 `bvj= ∞` V Junction reverse breakdown voltage.

Temperature effects parameters

- 73 `tnom` (C) Parameters measurement temperature. Default set by `options`.
- 74 `trise=0` C Temperature rise from ambient.
- 75 `tcv=1.0e-3` V/C Threshold voltage temperature coefficient.
- 76 `bex=-1.5` Mobility temperature exponent.
- 77 `ucex=0.8` Longitudinal critical field temp. exponent.
- 78 `ibbt=9.0e-4` 1/C Temperature coefficient for IBB.
- 79 `xti=3` Saturation current temperature exponent.
- 80 `tlev=0` DC temperature selector.
- 81 `tlevc=0` AC temperature selector.

Spectre Circuit Simulator Reference

Component Statements Part II

82	<code>eg=1.12452 V</code>	Energy band gap.
83	<code>gap1=7.02e-4 V/C</code>	Band gap temperature coefficient.
84	<code>gap2=1108 C</code>	Band gap temperature offset.
85	<code>tr1=0.6</code>	First source-drain resistance temperature coefficient.
86	<code>tr2=0.6</code>	Second source-drain resistance temperature coefficient.
87	<code>ptc=0 V/C</code>	Surface potential temperature coefficient.
88	<code>pta=0 V/C</code>	Junction potential temperature coefficient.
89	<code>ptp=0 V/C</code>	Sidewall junction potential temperature coefficient.
90	<code>cta=0 1/C</code>	Junction capacitance temperature coefficient.
91	<code>ctp=0 1/C</code>	Sidewall junction capacitance temperature coefficient.

Default instance parameters

92	<code>w=3e-6 m</code>	Default channel width.
93	<code>l=3e-6 m</code>	Default channel length.
94	<code>as=0 m²</code>	Default area of source diffusion.
95	<code>ad=0 m²</code>	Default area of drain diffusion.
96	<code>ps=0 m</code>	Default perimeter of source diffusion.
97	<code>pd=0 m</code>	Default perimeter of drain diffusion.
98	<code>nrd=0 m/m</code>	Default number of squares of drain diffusion.
99	<code>nrs=0 m/m</code>	Default number of squares of source diffusion.

Noise model parameters

100	<code>noisemod=1</code>	Noise model selector.
-----	-------------------------	-----------------------

Spectre Circuit Simulator Reference

Component Statements Part II

101	<code>kf=0</code>	Flicker (1/f) noise coefficient.
102	<code>af=1</code>	Flicker (1/f) noise exponent.
103	<code>ef=1</code>	Flicker (1/f) noise frequency exponent.

Model selection parameters

104	<code>nqs=0</code>	Nonquasi-static flag.
105	<code>satlim=exp(4)</code>	Ratio defining saturation limit.
106	<code>ekvint=0.0</code>	Interpolation function selector.
107	<code>scalem=1.0</code>	Model scaling factor.
108	<code>update=2.6</code>	Model version selector.

Auto Model Selector parameters

109	<code>wmax=1.0 m</code>	Maximum channel width for which the model is valid.
110	<code>wmin=0.0 m</code>	Minimum channel width for which the model is valid.
111	<code>lmax=1.0 m</code>	Maximum channel length for which the model is valid.
112	<code>lmin=0.0 m</code>	Minimum channel length for which the model is valid.

`Imax` and `Imelt`

The `imax` parameter aids convergence and prevents numerical overflow. The junction characteristics of the device are accurately modeled for current up to `imax`. If `imax` is exceeded during iterations, the linear model is substituted until the current drops below `imax` or until convergence is achieved. If convergence is achieved with the current exceeding `imax`, the results are inaccurate, and Spectre prints a warning.

A separate model parameter, `imelt`, is used as a limit warning for the junction current. This parameter can be set to the maximum current rating of the device. When any component of the junction current exceeds `imelt`, note that base and collector currents are composed of many exponential terms, Spectre issues a warning and the results become inaccurate. The junction current is linearized above the value of `imelt` to prevent arithmetic exception, with the exponential term replaced by a linear equation at `imelt`.

Spectre Circuit Simulator Reference

Component Statements Part II

Both of these parameters have current density counterparts, j_{max} and j_{melt} , that you can specify if you want the absolute current values to depend on the device area.

Auto Model Selection

Many models need to be characterized for different geometries in order to obtain accurate results for model development. The model selector program automatically searches for a model with the length and width range specified in the instance statement and uses this model in the simulations.

For the auto model selector program to find a specific model, the models to be searched should be grouped together within braces. Such a group is called a model group. An opening brace is required at the end of the line defining each model group. Every model in the group is given a name followed by a colon and the list of parameters. Also, the four geometric parameters l_{max} , l_{min} , w_{max} , and w_{min} should be given. The selection criteria to choose a model is as follows:

$$l_{min} \leq inst_length < l_{max} \text{ and } w_{min} \leq inst_width < w_{max}$$

Example:

```
model ModelName ModelType {  
    1: <model parameters> lmin=2 lmax=4 wmin=1 wmax=2  
    2: <model parameters> lmin=1 lmax=2 wmin=2 wmax=4  
    3: <model parameters> lmin=2 lmax=4 wmin=4 wmax=6  
}
```

Then for a given instance

```
M1 1 2 3 4 ModelName w=3 l=1.5
```

the program would search all the models in the model group with the name `ModelName` and then pick the first model whose geometric range satisfies the selection criteria. In the preceding example, the auto model selector program would choose `ModelName.2`.

You must specify both length (l) and width (w) on the device instance line to enable automatic model selection.

Spectre Circuit Simulator Reference

Component Statements Part II

Output Parameters

1	<code>w_{eff}</code> (m)	Effective channel width.
2	<code>l_{eff}</code> (m)	Effective channel length.
3	<code>r_{seff}</code> (Ω)	Effective source resistance.
4	<code>r_{deff}</code> (Ω)	Effective drain resistance.
5	<code>r_{deff}</code> (Ω)	Effective drain resistance.

Operating-Point Parameters

1	<code>type=n</code>	Transistor type. Possible values are <code>n</code> or <code>p</code> .
2	<code>region=triode</code>	Estimated operating region. Spectre outputs number (0-3) in a rawfile. Possible values are <code>off</code> , <code>triode</code> , <code>sat</code> , or <code>subth</code> .
3	<code>reversed</code>	Reverse mode indicator. Possible values are <code>no</code> or <code>yes</code> .
4	<code>ids</code> (A)	Resistive drain-to-source current.
5	<code>v_{gs}</code> (V)	Gate-source voltage.
6	<code>v_{ds}</code> (V)	Drain-source voltage.
7	<code>v_{bs}</code> (V)	Bulk-source voltage.
8	<code>v_p</code> (V)	Pinchoff voltage.
9	<code>v_{th}</code> (V)	Threshold voltage.
10	<code>v_{dss}</code> (V)	Drain-source saturation voltage.
11	<code>g_m</code> (S)	Common-source transconductance.
12	<code>g_{ds}</code> (S)	Common-source output conductance.
13	<code>g_{mbs}</code> (S)	Body-transconductance.

Spectre Circuit Simulator Reference

Component Statements Part II

14	<code>nfac</code>	Slope factor.
15	<code>if (A)</code>	Forward current.
16	<code>ir (A)</code>	Reverse current.
17	<code>irprime (A)</code>	Reverse current.
18	<code>isub (A)</code>	Substrate Current.
19	<code>ibd (A)</code>	Bulk-drain junction current.
20	<code>ibs (A)</code>	Bulk-source junction current.
21	<code>pwr (W)</code>	Power at op point.
22	<code>gmoverid (1/V)</code>	G_m/I_{ds} .
23	<code>gamma (\sqrt{V})</code>	Body-effect parameter.
24	<code>cjd (F)</code>	Drain-bulk junction capacitance.
25	<code>cjs (F)</code>	Source-bulk junction capacitance.
26	<code>cgg (F)</code>	Gate-gate capacitance.
27	<code>cgd (F)</code>	Gate-drain capacitance.
28	<code>cgs (F)</code>	Gate-source capacitance.
29	<code>cgb (F)</code>	Gate-bulk capacitance.
30	<code>cdg (F)</code>	Drain-gate capacitance.
31	<code>cdd (F)</code>	Drain-drain capacitance.
32	<code>cds (F)</code>	Drain-source capacitance.
33	<code>cdb (F)</code>	Drain-bulk capacitance.
34	<code>csg (F)</code>	Source-gate capacitance.
35	<code>csd (F)</code>	Source-drain capacitance.

Spectre Circuit Simulator Reference Component Statements Part II

36	<code>css</code> (F)	Source-source capacitance.
37	<code>csb</code> (F)	Source-bulk capacitance.
38	<code>cbg</code> (F)	Bulk-gate capacitance.
39	<code>cbd</code> (F)	Bulk-drain capacitance.
40	<code>cbs</code> (F)	Bulk-source capacitance.
41	<code>cbb</code> (F)	Bulk-bulk capacitance.
42	<code>vm</code> (V)	Early voltage.
43	<code>vovrdr</code> (V)	Overdrive voltage.
44	<code>tau</code> (s)	NQS time constant.
45	<code>tau0</code> (s)	Intrinsic time constant.
46	<code>ron</code> (Ω)	On-resistance.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

<code>ad</code>	I-4	<code>eg</code>	M-82	<code>mj</code>	M-42	<code>tau</code>	OP-44
<code>ad</code>	M-95	<code>ekvint</code>	M-106	<code>mjsw</code>	M-43	<code>tau0</code>	OP-45
<code>af</code>	M-102	<code>fc</code>	M-47	<code>mjswg</code>	M-45	<code>tcv</code>	M-75
<code>agamma</code>	M-67	<code>fcs</code>	M-51	<code>n</code>	M-35	<code>theta</code>	M-15
<code>akp</code>	M-66	<code>gamma</code>	M-10	<code>nfac</code>	OP-14	<code>tlev</code>	M-80
<code>alarm</code>	M-68	<code>gamma</code>	OP-23	<code>nfs</code>	M-7	<code>tlevc</code>	M-81
<code>as</code>	I-3	<code>gap1</code>	M-83	<code>noisemod</code>	M-100	<code>tnom</code>	M-73
<code>as</code>	M-94	<code>gap2</code>	M-84	<code>nqs</code>	M-104	<code>tox</code>	M-2

Spectre Circuit Simulator Reference Component Statements Part II

avto	M-65	gds	OP-12	nrd	I-7	tr1	M-85
bex	M-76	gm	OP-11	nrd	M-98	tr2	M-86
bvj	M-72	gmbs	OP-13	nrs	M-99	trise	I-14
cbb	OP-41	gmoverid	OP-22	nrs	I-8	trise	M-74
cbd	M-38	hdif	M-64	ns	I-12	tt	M-50
cbd	OP-39	iba	M-25	nsub	M-8	type	M-1
cbg	OP-38	ibb	M-26	pb	M-48	type	OP-1
cbs	M-39	ibbt	M-78	pbsw	M-49	ucex	M-77
cbs	OP-40	ibc	M-27	pbswg	M-46	ucrit	M-14
cdb	OP-33	ibd	OP-19	pd	I-6	uo	M-16
cdd	OP-31	ibn	M-28	pd	M-97	update	M-108
cdg	OP-30	ibs	OP-20	phi	M-11	vbox	M-71
cds	OP-32	ids	OP-4	ps	I-5	vbs	OP-7
cgb	OP-29	if	OP-15	ps	M-96	vds	OP-6
cgbo	M-54	imax	M-69	pta	M-88	vdss	OP-10
cgd	OP-27	imelt	M-37	ptc	M-87	vfb	M-18
cgdo	M-53	ir	OP-16	ptp	M-89	vgs	OP-5
cgg	OP-26	irprime	OP-17	pwr	OP-21	vm	OP-42
cgs	OP-28	is	M-32	q0	M-29	vmax	M-17
cgso	M-52	isub	OP-18	rd	M-56	vovrdr	OP-43
cj	M-40	jmax	M-70	rdc	I-9	vp	OP-8
cjd	OP-24	js	M-33	rdc	M-61	vth	OP-9
cjs	OP-25	jsw	M-34	rdd	M-59	vto	M-9
cjsw	M-41	kf	M-101	rdeff	O-4	w	M-92
cjswg	M-44	kp	M-12	rdeff	O-5	w	I-1
cox	M-3	l	M-93	region	I-13	weff	O-1
csb	OP-37	l	I-2	region	OP-2	weta	M-20
csd	OP-35	lambda	M-19	reversed	OP-3	wmax	M-109
csg	OP-34	ldif	M-63	ron	OP-46	wmin	M-110
css	OP-36	leff	O-2	rs	M-55	xj	M-4
cta	M-90	leta	M-21	rsc	M-60	xl	M-23
ctp	M-91	lk	M-30	rsc	I-10	xqc	M-31

Spectre Circuit Simulator Reference

Component Statements Part II

dl	M-6	lmax	M-111	rseff	O-3	xti	M-79
dskip	M-36	lmin	M-112	rsh	M-57	xw	M-22
dw	M-5	m	I-11	rss	M-58		
e0	M-13	meto	M-24	satlim	M-105		
ef	M-103	minr	M-62	scalem	M-107		

Ratiometric Fourier Analyzer (fourier)

Description

The ratiometric Fourier analyzer measures the Fourier coefficients of two different signals at a specified fundamental frequency without loading the circuit. The algorithm used is based on the Fourier integral rather than the discrete Fourier transform and therefore is not subject to aliasing. Even on broad-band signals, it computes a small number of Fourier coefficients accurately and efficiently. Therefore, this Fourier analyzer is suitable on clocked sinusoids generated by sigma-delta converters, pulse-width modulators, digital-to-analog converters, sample-and-holds, and switched-capacitor filters as well as on the traditional low-distortion sinusoids produced by amplifiers or filters.

The analyzer is active only during a transient analysis. For each signal, the analyzer prints the magnitude and phase of the harmonics along with the total harmonic distortion at the end of the transient analysis. The total harmonic distortion is found by summing the power in all of the computed harmonics except DC and the fundamental. Consequently, the distortion is not accurate if you request an insufficient number of harmonics. The Fourier analyzer also prints the ratio the spectrum of the first signal to the fundamental of the second, so you can use the analyzer to compute large signal gains and immittances directly.

If you are concerned about accuracy, perform an additional Fourier transform on a pure sinusoid generated by an independent source. Because both transforms use the same time points, the relative errors measured with the known pure sinusoid are representative of the errors in the other transforms. In practice, this second Fourier transform is performed on the reference signal. To increase the accuracy of the Fourier transform, use the `points` parameter to increase the number of points. Tightening `reltol` and setting `errpreset=conservative` are two other measures to consider.

The accuracy of the magnitude and phase for each harmonic is independent of the number of harmonics computed. Thus, increasing the number of harmonics (while keeping `points` constant) does not change the magnitude and phase of the low order harmonics, but it does improve the accuracy of the total harmonic distortion computation. However, if you do not

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specify `points`, you can increase accuracy by requesting more harmonics, which creates more points.

The large number of points required for accurate results is not a result of aliasing. Many points are needed because a quadratic polynomial interpolates the waveform between the time-points. If you use too few time-points the polynomials deviate slightly from the true waveform between time-points and all of the computed Fourier coefficients are slightly in error. The algorithm that computes the Fourier integral does accept unevenly spaced time-points, but because it uses quadratic interpolation, it is usually more accurate using time-steps that are small and nearly evenly spaced.

This device is not supported within `altergroup`.

Sample Instance Statement

```
four1 (1 0) fourmod harms=50
```

Sample Model Statement

```
model fourmod fourier fund=900M points=2500 order=2
```

Instance Definition

```
Name [p] [n] [pr] [nr] ModelName parameter=value ...
```

Name [p] [n] [pr] [nr] fourier parameter=value ...

The signal between terminals `p` and `n` is the test or numerator signal. The signal between terminals `pr` and `nr` is the reference or denominator signal. Fourier analysis is performed on terminal currents by specifying the `term` or `refterm` parameters. If both `term` and `p` or `n` are specified, then the terminal current becomes the numerator and the node voltages become the denominator. By mixing voltages and currents, it is possible to compute large signal immittances.

Instance Parameters

- | | | |
|---|---|---|
| 1 | <code>fund</code> (Hz) | Fundamental frequency. |
| 2 | <code>points=20</code> <code>maxharm</code> | Minimum number of time points. |
| 3 | <code>active=yes</code> | Whether Fourier analysis should be performed or skipped.
Possible values are <code>no</code> or <code>yes</code> . |

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4	<code>order=2</code>	Order of interpolation.
5	<code>term</code>	Terminal used to measure current for test (numerator) channel.
6	<code>refterm</code>	Terminal used to measure current for reference (denominator) channel.
7	<code>harmsvec=[...]</code>	Array of desired harmonics for test (numerator) channel.
8	<code>harms=9</code>	Number of harmonics for test (numerator) channel, if an array is not given. The harmonics start from <code>firstharm</code> and go up to <code>firstharm + harms - 1</code> .
9	<code>refharmsvec=[...]</code>	Array of desired harmonics for reference (denominator) channel.
10	<code>refharms=9</code>	Number of harmonics for reference (denominator) channel, if an array is not given. The harmonics start from <code>reffirstharm</code> and go up to <code>reffirstharm + harms - 1</code> .
11	<code>scale=1</code>	Scale factor for ratioed results.
12	<code>firstharm=1</code>	First harmonic computed for test (numerator) channel.
13	<code>reffirstharm=1</code>	First harmonic computed for reference (denominator) channel.
14	<code>normharm=1</code>	Normalizing harmonic for test (numerator) channel.
15	<code>refnormharm=1</code>	Normalizing harmonic for reference (denominator) channel.
16	<code>where=logfile</code>	Where Fourier results should be printed. Possible values are <code>screen</code> , <code>logfile</code> or <code>both</code> .

Model Definition

```
model modelName fourier parameter=value ...
```

Model Parameters

1	<code>fund (Hz)</code>	Fundamental frequency.
2	<code>points=20 maxharm</code>	Minimum number of time points.

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3	<code>harms=9</code>	Desired number of harmonics.
4	<code>active=yes</code>	Whether Fourier analysis should be performed or skipped. Possible values are <code>no</code> or <code>yes</code> .
5	<code>order=2</code>	Order of interpolation.
6	<code>firstharm=1</code>	First harmonic computed for test (numerator) channel.
7	<code>reffirstharm=1</code>	First harmonic computed for reference (denominator) channel.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

<code>active</code> I-3	<code>harms</code> I-8	<code>points</code> I-2	<code>refnormharm</code> I-15
<code>active</code> M-4	<code>harms</code> M-3	<code>points</code> M-2	<code>refterm</code> I-6
<code>firstharm</code> M-6	<code>harmsvec</code> I-7	<code>reffirstharm</code> M-7	<code>scale</code> I-11
<code>firstharm</code> I-12	<code>normharm</code> I-14	<code>reffirstharm</code> I-13	<code>term</code> I-5
<code>fund</code> I-1	<code>order</code> M-5	<code>refharms</code> I-10	<code>where</code> I-16
<code>fund</code> M-1	<code>order</code> I-4	<code>refharmsvec</code> I-9	

GaAs MESFET (gaas)

Description

The GaAs MESFET model was derived from the model by H. Statz and others at Raytheon. This model is completely symmetric and is modified slightly to make it charge conserving. GaAs MESFET instances require that you use a model statement.

This device is supported within altergroups.

There are some convergence problems with this model because of `Cgs` going to zero beyond pinchoff. The problems occur when the gate is driven from an inductive source, and there is

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no other capacitance at the gate. To prevent these problems, avoid setting C_{gd} to zero and add sidewall capacitance to the gate-source and gate-drain junctions. A good estimate for these capacitors is $C = \pi \cdot \epsilon \cdot w/2$ where w is the gate width in microns and $\epsilon = 0.116$ fF/micron.

Sample Instance Statement

```
m1 (1 2 0) nmes area=1 m=2
```

Sample Model Statement

```
model nmes gaas type=n vto=-2 beta=0.06 lambda=0 b=0.25 rs=3.65 alpha=1.9 rd=1.98  
is=1.1e-9 n=1.28 fc=0.5 cgs=0.365e-12
```

Instance Definition

```
Name d g s ModelName parameter=value ...
```

Instance Parameters

- | | | |
|---|------------|--|
| 1 | area=1 | Junction area factor. |
| 2 | m=1 | Multiplicity factor. |
| 3 | region=fwd | Estimated operating region. Spectre outputs number (0-4) in a rawfile.
Possible values are off, triode, sat, subth, or breakdown. |

Model Definition

```
model modelName gaas parameter=value ...
```

Model Parameters

Device type parameters

- | | | |
|---|--------|---|
| 1 | type=n | Transistor type.
Possible values are n or p. |
|---|--------|---|

Drain current parameters

- | | | |
|---|----------|--------------------|
| 2 | vto=-2 V | Pinch-off voltage. |
|---|----------|--------------------|

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- 3 `beta=0.0001 A/V2` Transconductance parameter.
- 4 `lambda=0 1/V` Channel length modulation parameter.
- 5 `b=0.3 1/V` Doping tail extending parameter.
- 6 `alpha=2 1/V` Saturation voltage parameter.

Parasitic resistance parameters

- 7 `rd=0 Ω` Drain resistance (/area).
- 8 `rs=0 Ω` Source resistance (/area).
- 9 `rg=0 Ω` Gate resistance (/area).
- 10 `minr=0.1 Ω` Minimum source/drain/gate resistance.

Junction diode model parameters

- 11 `is=1e-14 A` Gate saturation current (*area).
- 12 `n=1` Emission coefficient for the gate junction.
- 13 `imelt=`imaxA'` Explosion current (*area).
- 14 `dskip=yes` Use simple piece-wise linear model for diode currents below $0.1 \cdot i_{abstol}$.
Possible values are `no` or `yes`.

Junction capacitance model parameters

- 15 `capmod=2` Charge model selector.
- 16 `cgs=0 F` Gate-source zero-bias junction capacitance (*area).
- 17 `cgd=0 F` Gate-drain zero-bias junction capacitance (*area).
- 18 `pb=1 V` Gate junction potential.
- 19 `fc=0.5` Junction capacitor forward-bias threshold.

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20 `delta=0.2 V` Gate capacitance pinch-off transition width.

Temperature effects parameters

21 `tnom (C)` Parameters measurement temperature. Default set by options.

22 `trise=0 C` Temperature rise from ambient.

23 `xti=3` Temperature exponent for effect on *i_s*.

Operating region warning control parameters

24 `alarm=none` Forbidden operating region.
Possible values are `none`, `off`, `triode`, `sat`, `subth`, or `rev`.

25 `imax=1 A` Maximum allowable current (*area).

26 `bvj=∞ V` Junction reverse breakdown voltage.

Noise model parameters

27 `kf=0` Flicker noise (1/f) coefficient.

28 `af=1` Flicker noise (1/f) exponent.

I_{max} and *I_{melt}*

The *i_{max}* parameter aids convergence and prevents numerical overflow. The junction characteristics of the device are accurately modeled for current up to *i_{max}*. If *i_{max}* is exceeded during iterations, the linear model is substituted until the current drops below *i_{max}* or until convergence is achieved. If convergence is achieved with the current exceeding *i_{max}*, the results are inaccurate, and Spectre prints a warning.

A separate model parameter, *i_{melt}*, is used as a limit warning for the junction current. This parameter can be set to the maximum current rating of the device. When any component of the junction current exceeds *i_{melt}*, note that base and collector currents are composed of many exponential terms, Spectre issues a warning and the results become inaccurate. The junction current is linearized above the value of *i_{melt}* to prevent arithmetic exception, with the exponential term replaced by a linear equation at *i_{melt}*.

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The `bv` parameter detects the junction breakdown only. The breakdown currents of the junctions are not modeled.

Operating-Point Parameters

1	<code>type=n</code>	Transistor type. Possible values are <code>n</code> or <code>p</code> .
2	<code>region=fwd</code>	Estimated operating region. Spectre outputs number (0-4) in a rawfile. Possible values are <code>off</code> , <code>triode</code> , <code>sat</code> , <code>subth</code> , or <code>breakdown</code> .
3	<code>ids (A)</code>	Resistive drain current.
4	<code>vth (V)</code>	Threshold voltage.
5	<code>vgs (V)</code>	Gate-source voltage.
6	<code>vds (V)</code>	Drain-source voltage.
7	<code>vdsat (V)</code>	Drain saturation voltage.
8	<code>gm (S)</code>	Common-source transconductance.
9	<code>gds (S)</code>	Common-source output conductance.
10	<code>cgs (F)</code>	Gate-source capacitance.
11	<code>cgd (F)</code>	Gate-drain capacitance.
12	<code>ig (A)</code>	Resistive gate current.
13	<code>pwr (W)</code>	Power at operating point.

Parameter Index

In the following index, `I` refers to instance parameters, `M` refers to the model parameters section, `O` refers to the output parameters section, and `OP` refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the

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description for that parameter. For example, a reference of M-35 means the 35th model parameter.

af	M-28	cgs	M-16	kf	M-27	rs	M-8
alarm	M-24	delta	M-20	lambda	M-4	tnom	M-21
alpha	M-6	dskip	M-14	m	I-2	trise	M-22
area	I-1	fc	M-19	minr	M-10	type	M-1
b	M-5	gds	OP-9	n	M-12	type	OP-1
beta	M-3	gm	OP-8	pb	M-18	vds	OP-6
bvj	M-26	ids	OP-3	pwr	OP-13	vdsat	OP-7
capmod	M-15	ig	OP-12	rd	M-7	vgs	OP-5
cgd	OP-11	imax	M-25	region	OP-2	vth	OP-4
cgd	M-17	imelt	M-13	region	I-3	vto	M-2
cgs	OP-10	is	M-11	rg	M-9	xti	M-23

Hetero-Junction Bipolar Transistor (hbt)

Description

The HBT (Hetero-junction Bipolar Transistor) model was developed by UCSD as part of the ARPA High Speed Circuit Design Program. The model has four external electrical nodes, one thermal node, and up to nine internals depending on the complexity of the model users specified. Detailed description of the model and equations can be found in the Affirma Spectre Circuit Simulator Device Model Equations manual.

This device is supported within altergroups.

Sample Instance Statement

```
q7 (net5 net2 0) hbtmod m=1 top=25
```

Sample Model Statement

```
model hbtmod hbt type=npn bf=500 br=1000 xtb=-2.4 xti=0 xcjc=0.83 mje=0.34 fc=0.5  
eg=1.2 ise=5.5e-15 vjc=0.84 vaf=40 cjc=5.1e-15
```

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Instance Definition

Name c b e [t] [s] ModelName parameter=value ...

It is not necessary to specify the substrate and thermal terminal. If left unspecified, the substrate node is connected to ground while the thermal node is fixed to the ambient temperature. However, you must specify the thermal node if you specify the substrate node.

Instance Parameters

- | | | |
|---|------------|--|
| 1 | area=1 | Transistor area factor. |
| 2 | m=1 | Multiplicity factor. |
| 3 | top (C) | Average device operating temperature. |
| 4 | region=fwd | Estimated operating region. Spectre outputs number (0-3) in a rawfile.
Possible values are off, fwd, rev, or sat. |

Model Definition

model modelName hbt parameter=value ...

Model Parameters

Structural parameters

- | | | |
|---|----------|---|
| 1 | type=npn | Transistor type.
Possible values are npn or pnp. |
|---|----------|---|

Saturation current parameters

- | | | |
|---|--------------|--|
| 2 | is=1e-25 A | Saturation value for forward collector current (*area). |
| 3 | ise=1e-25 A | Saturation value for nonideal base current. (*area). |
| 4 | isex=1e-25 A | Saturation current for emitter leakage diode (*area). |
| 5 | isc=1e-20 A | Saturation value for intrinsic BC junction current. (*area). |
| 6 | iscx=1e-20 A | Saturation value for extrinsic B-C junction current (*area). |

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7 `ics=1e-30 A` Saturation value for C-S junction current (*area).

Emission coefficient parameters

8 `nf=1` Forward collector current ideality factor.

9 `nr=1` Reverse ideality factor.

10 `ne=2` Nonideal base forward current ideality factor.

11 `nex=2` Ideality factor for emitter leakage diode.

12 `nc=2` Intrinsic B-C junction ideality factor.

13 `ncx=2` Ideality factor for extrinsic B-C junction.

14 `ncs=2` Ideality factor for C-S junction.

Current gain parameters

15 `bf=1000 A/A` Forward ideal current gain (beta).

16 `br=1000 A/A` Reverse ideal current gain.

17 `isa=1e10 A` Collector E-B barrier limiting current (*area).

18 `na=2` Collector E-B barrier ideality factor.

19 `isb=1e10 A` Collector B-C barrier limiting current (*area).

20 `nb=2` Collector B-C barrier ideality factor.

21 `ik=1e10 A` Knee current for dc high-level injection effect (*area).

Early voltage parameters

22 `vaf=500 V` Forward Early voltage.

23 `var=500 V` Reverse Early voltage.

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Breakdown voltage parameters

24	<code>bkdn=no</code>	Flag denoting B-C breakdown should be included. Possible values are <code>no</code> or <code>yes</code> .
25	<code>bvc=50 A</code>	Collector-base breakdown voltage BV_{cbo} .
26	<code>nbc=8</code>	Exponent for B-C multiplication factor versus voltage.
27	<code>fa=0.9</code>	Factor for specification of avalanche voltage.
28	<code>imax=1 A</code>	Maximum allowable base current (*area).
29	<code>imelt=10 A</code>	Explosion current (*area).

Parasitic resistance parameters

30	<code>rbi=0 Ω</code>	Intrinsic base resistance (/area).
31	<code>rbx=0 Ω</code>	Extrinsic base resistance (/area).
32	<code>rci=0 Ω</code>	Intrinsic collector resistance (/area).
33	<code>rcx=0 Ω</code>	Extrinsic collector resistance (/area).
34	<code>re=0 Ω</code>	Emitter resistance (/area).
35	<code>rex=0 Ω</code>	Extrinsic emitter leakage diode series resistance (/area).

Junction capacitance parameters

36	<code>cje=0 F</code>	B-E depletion capacitance at zero bias (*area).
37	<code>vje=1.6 V</code>	B-E built-in potential for C_j .
38	<code>mje=0.5</code>	Exponent for voltage variation of B-E C_j .
39	<code>cemin=0 F</code>	Minimum B-E capacitance (*area).
40	<code>fce=0.8</code>	Factor for start of high bias B-E C_j approximation.
41	<code>cjc=0 F</code>	Intrinsic B-C depletion capacitance at zero bias (*area).

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42	<code>vjc=1.4 V</code>	Intrinsic B-C built-in potential for Cj.
43	<code>mjc=0.33</code>	Exponent for voltage variation of Intrinsic B-C Cj.
44	<code>ccmin=0 F</code>	Minimum B-C capacitance (*area).
45	<code>fc=0.8</code>	Factor for start of high bias B-C Cj approximation.
46	<code>cjcx=0 F</code>	Extrinsic B-C depletion capacitance at zero bias (*area).
47	<code>vjcx=1.4 V</code>	Extrinsic B-C built-in potential for Cj.
48	<code>mjcx=0.33</code>	B-C junction exponent.
49	<code>cxmin=0 F</code>	Minimum extrinsic B-C capacitance (*area).
50	<code>xcjc=0</code>	Fraction of B-C capacitance tied to external base node.
51	<code>cjs=0 F</code>	B-S depletion capacitance at zero bias (*area).
52	<code>vjs=1.4 V</code>	B-S built-in potential for Cj.
53	<code>mjs=0.5</code>	Exponent for voltage variation of C-S Cj.

Transit time and excess phase parameters

54	<code>tfb=0 s</code>	Base transit time.
55	<code>tbexs=0</code>	Excess B-E heterojunction transit time.
56	<code>tbcxs=0</code>	Excess B-C heterojunction transit time.
57	<code>tfc0=0 s</code>	Collector forward transit time.
58	<code>icrit0=1e3 A</code>	Critical current for intrinsic Cj variation.
59	<code>itc=0 A</code>	Characteristic current for Tfc.
60	<code>itc2=0 A</code>	Characteristic current for Tfc.
61	<code>vtc=1e3 V</code>	Characteristic voltage for Tfc.
62	<code>tkrk=0 s</code>	Forward transit time for Kirk effect.

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63	<code>vkrk=1e3 V</code>	Characteristic voltage for Kirk effect.
64	<code>ikrk=1e3 A</code>	Characteristic voltage for Kirk effect.
65	<code>tr=0 s</code>	Reverse charge storage time for intrinsic B-C junction.
66	<code>trx=0 s</code>	Reverse charge storage time for extrinsic B-C junction.
67	<code>fex=0 s</code>	Factor to determine excess phase.

Temperature effects parameters

68	<code>selft=no</code>	Flag denoting self-heating. Possible values are <code>no</code> or <code>yes</code> .
69	<code>tnom (C)</code>	Parameters measurement temperature. Default set by options.
70	<code>top=27 C</code>	Average device operating temperature.
71	<code>rth=0 Ω</code>	Thermal resistance of device.
72	<code>cth=0 F</code>	Thermal capacitance of device.
73	<code>xti=2</code>	Exponent for <code>is</code> temperature dependence.
74	<code>xtb=2</code>	Exponent for beta temperature dependence.
75	<code>tne=0</code>	Coefficient for <code>ne</code> temperature dependence.
76	<code>tnc=0</code>	Coefficient for <code>nc</code> temperature dependence.
77	<code>tnex=0</code>	Coefficient for <code>nex</code> temperature dependence.
78	<code>eaе=0 V</code>	Activation energy for <code>isa</code> temperature dependence.
79	<code>eac=0 V</code>	Activation energy for <code>isb</code> temperature dependence.
80	<code>eaa=0 V</code>	Activation energy for <code>ise</code> temperature dependence.
81	<code>eab=0 V</code>	Activation energy for <code>isc</code> temperature dependence.
82	<code>eax=0 V</code>	Activation energy for <code>isex</code> temperature dependence.

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83	<code>xre=0</code>	Exponent for r_e temperature dependence.
84	<code>xrex=0</code>	Exponent for r_{ex} temperature dependence.
85	<code>xrb=0</code>	Exponent for r_b temperature dependence.
86	<code>xrc=0</code>	Exponent for r_c temperature dependence.
87	<code>tvje=0 V/C</code>	Coefficient for v_{je} temperature dependence.
88	<code>tvjc=0 V/C</code>	Coefficient for v_{jc} temperature dependence.
89	<code>tvjcx=0 V/C</code>	Coefficient for v_{jcx} temperature dependence.
90	<code>tvjs=0 V/C</code>	Coefficient for v_{js} temperature dependence.
91	<code>xtitc=0</code>	Exponent for i_{tc} temperature dependence.
92	<code>xtitc2=0</code>	Exponent for i_{tc2} temperature dependence.
93	<code>xttf=0</code>	Exponent for t_f temperature dependence.
94	<code>xttkrk=0</code>	Exponent for t_{krk} temperature exponent.
95	<code>xtvkrk=0</code>	Exponent for v_{krk} temperature dependence.
96	<code>xtikrk=0</code>	Exponent for i_{krk} temperature dependence.
97	<code>xrt=0</code>	Exponent for r_{th} temperature dependence.
98	<code>eg=1.5 V</code>	Activation energy for i_s temperature dependence.
99	<code>dtmax=1000 C</code>	Maximum expected temperature rise above heat sink.

Noise model parameters

100	<code>kfn=0</code>	Flicker (1/f) noise coefficient.
101	<code>afn=1</code>	Flicker (1/f) noise exponent.
102	<code>bfm=1</code>	Flicker noise frequency exponent.

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Component Statements Part II

Imax and Imelt

The `imax` parameter aids convergence and prevents numerical overflow. The junction characteristics of the device are accurately modeled for current up to `imax`. If `imax` is exceeded during iterations, the linear model is substituted until the current drops below `imax` or until convergence is achieved. If convergence is achieved with the current exceeding `imax`, the results are inaccurate, and Spectre prints a warning.

A separate model parameter, `imelt`, is used as a limit warning for the junction current. This parameter can be set to the maximum current rating of the device. When any component of the junction current exceeds `imelt`, note that base and collector currents are composed of many exponential terms, Spectre issues a warning and the results become inaccurate. The junction current is linearized above the value of `imelt` to prevent arithmetic exception, with the exponential term replaced by a linear equation at `imelt`.

Operating-Point Parameters

1	<code>type=npn</code>	Transistor type. Possible values are <code>npn</code> or <code>pnP</code> .
2	<code>region=fwd</code>	Estimated operating region. Spectre outputs number (0-3) in a rawfile. Possible values are <code>off</code> , <code>fwd</code> , <code>rev</code> , or <code>sat</code> .
3	<code>vbe (V)</code>	Base-emitter voltage.
4	<code>vbc (V)</code>	Base-collector voltage.
5	<code>vce (V)</code>	Collector-emitter voltage.
6	<code>vcs (V)</code>	XC-substrate voltage.
7	<code>temp (C)</code>	Device temperature.
8	<code>ith (A)</code>	Thermal source.
9	<code>ice (A)</code>	Intrinsic B-C current.
10	<code>ibe (A)</code>	Intrinsic B-E current.
11	<code>ics (A)</code>	C-S junction current.
12	<code>ibei (A)</code>	B-E junction current.

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13	<code>ibci</code> (A)	B-C junction current.
14	<code>ibex</code> (A)	XB-E junction current.
15	<code>ibcx</code> (A)	XB-C junction current.
16	<code>ibk</code> (A)	Breakdown current.
17	<code>dice_dvbe</code> (S)	Intrinsic dl_{ce}/dV_{be} .
18	<code>dice_dvbc</code> (S)	Intrinsic dl_{ce}/dV_{bc} .
19	<code>dibe_dvbe</code> (S)	Intrinsic dl_{be}/dV_{be} .
20	<code>dibe_dvbc</code> (S)	Intrinsic dl_{be}/dV_{bc} .
21	<code>dqbe_dvbe</code> (F)	Intrinsic dQ_{be}/dV_{be} .
22	<code>dqbe_dvbc</code> (F)	Intrinsic dQ_{be}/dV_{bc} .
23	<code>dqbc_dvbe</code> (F)	Intrinsic dQ_{bc}/dV_{be} .
24	<code>dqbc_dvbc</code> (F)	Intrinsic dQ_{bc}/dV_{bc} .
25	<code>cbcx</code> (F)	XB-C junction capacitance.
26	<code>cbcxx</code> (F)	EXTB-C junction capacitance.
27	<code>ccs</code> (F)	Substrate junction capacitance.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

<code>afn</code>	M-101	<code>fc</code>	M-45	<code>nb</code>	M-20	<code>tvjcx</code>	M-89
<code>area</code>	I-1	<code>fce</code>	M-40	<code>nbc</code>	M-26	<code>tvje</code>	M-87

Spectre Circuit Simulator Reference Component Statements Part II

bf	M-15	fex	M-67	nc	M-12	tvjs	M-90
bfm	M-102	ibci	OP-13	ncs	M-14	type	M-1
bkdn	M-24	ibcx	OP-15	ncx	M-13	type	OP-1
br	M-16	ibe	OP-10	ne	M-10	vaf	M-22
bvc	M-25	ibei	OP-12	nex	M-11	var	M-23
cbcx	OP-25	ibex	OP-14	nf	M-8	vbc	OP-4
cbcxx	OP-26	ibk	OP-16	nr	M-9	vbe	OP-3
ccmin	M-44	ice	OP-9	rbi	M-30	vce	OP-5
ccs	OP-27	icrit0	M-58	rbx	M-31	vcs	OP-6
cemin	M-39	ics	M-7	rci	M-32	vjc	M-42
cjc	M-41	ics	OP-11	rcx	M-33	vjcx	M-47
cjcx	M-46	ik	M-21	re	M-34	vje	M-37
cje	M-36	ikrk	M-64	region	I-4	vjs	M-52
cjs	M-51	imax	M-28	region	OP-2	vkrk	M-63
cth	M-72	imelt	M-29	rex	M-35	vtc	M-61
cxmin	M-49	is	M-2	rth	M-71	xcjc	M-50
dibe_dvbc	OP-20	isa	M-17	selft	M-68	xrb	M-85
dibe_dvbe	OP-19	isb	M-19	tbcxs	M-56	xrc	M-86
dice_dvbc	OP-18	isc	M-5	tbexs	M-55	xre	M-83
dice_dvbe	OP-17	iscx	M-6	temp	OP-7	xrex	M-84
dqbc_dvbc	OP-24	ise	M-3	tfb	M-54	xrt	M-97
dqbc_dvbe	OP-23	isex	M-4	tfc0	M-57	xtb	M-74
dqbe_dvbc	OP-22	itc	M-59	tkrk	M-62	xti	M-73
dqbe_dvbe	OP-21	itc2	M-60	tnc	M-76	xtikrk	M-96
dtmax	M-99	ith	OP-8	tne	M-75	xtitc	M-91
ea	M-80	kfn	M-100	tnex	M-77	xtitc2	M-92
eab	M-81	m	I-2	tnom	M-69	xttf	M-93
eac	M-79	mjc	M-43	top	M-70	xttkrk	M-94
eae	M-78	mjcx	M-48	top	I-3	xtvkrk	M-95
eax	M-82	mje	M-38	tr	M-65		
eg	M-98	mjs	M-53	trx	M-66		
fa	M-27	na	M-18	tvjc	M-88		

HiSIM1 Field Effect Transistor (hisim)

Description

HiSIM MOS IGFET DEVICE

The HiSIM (Hiroshima-university STARC IGFET Model) model was developed at Hiroshima University in collaboration with the STARC research center. The model is based on the drift-diffusion approximation. All device characteristics are described by channel-surface potentials at the source side and at the drain side. These surface potentials are implicit functions of applied voltages, therefore additional iteration procedures are required to global iterations in circuit simulation.

This device is supported within altergroups.

Sample Instance Statement

```
m4 (0 2 1 1) nch w=2u l=0.8u as=250p ad=250p pd=168p ps=168p temp=350
```

Sample Model Statement

```
model nch hisim version=111 type=n
```

Instance Definition

```
Name d g s b ModelName parameter=value ...
```

Instance Parameters

1	w (m)	Channel width.
2	l (m)	Channel length.
3	as (m ²)	Area of source diffusion.
4	ad (m ²)	Area of drain diffusion.
5	ps (m)	Perimeter of source diffusion.
6	pd (m)	Perimeter of drain diffusion.
7	temp (K)	Lattice temperature.

Spectre Circuit Simulator Reference

Component Statements Part II

8 `dtemp` (K) Lattice temperature rise from ambient.

Model Definition

```
model modelName hisim parameter=value ...
```

Model Parameters

Device type parameters

1	<code>type=n</code>	Transistor type. Possible values are n or p.
2	<code>version=101</code>	Version of HiSIM.
3	<code>noise=5</code>	Noise model selector.
4	<code>corsrd=0</code>	Flag for accounting Rs and Rd.
5	<code>cocgso=0</code>	Flag for calculate cgso.
6	<code>cocgdo=0</code>	Flag for calculate cgdo.
7	<code>cocgbo=0</code>	Flag for calculate cgbo.
8	<code>coadov=1</code>	Flag for overlap charges/capacitances.
9	<code>coisub=0</code>	Flag for calculate isub.
10	<code>cogidl=0</code>	Flag for calculate I _{gidl} current.
11	<code>coiigs=0</code>	Flag for calculate I _{gate} current.
12	<code>coovlp=0</code>	Flag for calculate overlap charge.
13	<code>conois=0</code>	Flag for calculate 1/f noise.
14	<code>coisti=0</code>	Flag for calculate STI HiSIM1.1.
15	<code>coiprv=1</code>	Flag for use <code>ids_prv</code> as initial guess of I _{ds} .
16	<code>copprrv=1</code>	Flag for use <code>ps{0/l}_prv</code> as initial guess of P _{s{0/l}} .

Spectre Circuit Simulator Reference

Component Statements Part II

Default for instance parameters

17	$w=5e-6$ m	Default channel width.
18	$l=5e-6$ m	Default channel length.
19	$a_s=0$ m ²	Default area of source diffusion.
20	$a_d=0$ m ²	Default area of drain diffusion.
21	$p_s=0$ m	Default perimeter of source diffusion.
22	$p_d=0$ m	Default perimeter of drain diffusion.
23	$t_{temp}=300.15$ K	Default lattice temperature.
24	$dt_{temp}=0$ K	Default lattice temperature rise from ambient.

Technological parameters

25	$t_{ox}=5.0e-9$ m	Gate oxide thickness.
26	$x_{ld}=0$ m	Gate-overlap length.
27	$x_{wd}=0$ m	Gate-overlap width.
28	$x_{polyd}=0$ m	Difference between gate-poly and design lengths.
29	$t_{poly}=0$ m	Height of the gate poly-si.
30	$n_{subc}=1.0e17$ cm ⁻³	Constant part of Nsub.
31	$n_{subp}=1.0e17$ cm ⁻³	Peak of the pocket concentration.
32	$r_s=8.0e-5$ Ω m	Source contact resistance.
33	$r_d=8.0e-5$ Ω m	Drain contact resistance.
34	$v_{fbc}=-1.0$ V	Constant part of flat-band voltage.
35	$l_p=1.5e-8$ m	Length of the pocket penetration into the channel.

Spectre Circuit Simulator Reference

Component Statements Part II

- 36 $x_j=0.0$ m Junction depth, used only in HiSIM1.0.0.
- 37 $x_{qy}=0.0$ m Distance from channel/drain junction to maximum electric field point, used only in HiSIM1.1.0.

Temperature dependence effects

- 38 $b_{gtmp1}=9.02e-5$ eV/K First order temperature coefficient for band gap.
- 39 $b_{gtmp2}=1.0e-7$ eV/K² Second order temperature coefficient for band gap.

Quantum Mechanical Effects

- 40 $q_{me1}=4.0e-11$ m/V² Coefficient for quantum mechanical effect.
- 41 $q_{me2}=3.0e-10$ V Coefficient for quantum mechanical effect.
- 42 $q_{me3}=0.0$ m Coefficient for quantum mechanical effect.

Poly Depletion Effects

- 43 $pgd1=0.01$ V Strength of poly depletion.
- 44 $pgd2=1.0$ V Threshold voltage of poly depletion.
- 45 $pgd3=0.8$ V_{ds} dependence of poly depletion.

Short Channel Effects

- 46 $par11=1.0$ Strength of lateral-electric-field gradient.
- 47 $par12=0.0$ m Depletion width of channel/contact junction.
- 48 $sc1=0.0$ 1/V Short-channel coefficient 1.
- 49 $sc2=0.0$ 1/V² Short-channel coefficient 2.
- 50 $sc3=0.0$ m/V² Short-channel coefficient 3.

Spectre Circuit Simulator Reference

Component Statements Part II

- 51 `scp1=0.0 1/V` Short-channel coefficient 1 for pocket.
- 52 `scp2=0.0 1/V2` Short-channel coefficient 2 for pocket.
- 53 `scp3=0.0 m/V2` Short-channel coefficient 3 for pocket.

Narrow channel effects

- 54 `wfc=0.0 m F/cm2` Threshold voltage reduction.
- 55 `mueph2=0.0` Mobility reduction.
- 56 `w0=0.0 log(cm)` Minimum gate width.
- 57 `wvthsc=0.0` Short-channel effect at the STI edge.
- 58 `nsti=1.0e17 cm-3` Substrate-impurity concentration at the SIT edge.
- 59 `wsti=0.0 m` Width of the high-field region at STI.

Mobility Effects

- 60 `vds0=0.05 V` Drain voltage for extracting the low-field mobility.
- 61 `muecb0=300.0 cm2/(V s)`
Coulomb scattering.
- 62 `muecb1=30.0 cm2/(V s)`
Coulomb scattering.
- 63 `mueph0=0.3 cm2 (V/cm)(Muesr1)/(V s)`
Phonon scattering.
- 64 `mueph1=2.5e4` Phonon scattering.
- 65 `muetmp=1.5` Temperature dependence of phonon scattering.
- 66 `muesr0=2.0 cm2 (V/cm)(Muesr1)/(V s)`
Surface-roughness scattering.
- 67 `muesr1=2.0e15` Surface-roughness scattering.

Spectre Circuit Simulator Reference

Component Statements Part II

- 68 `ndep=1.0` Coefficient of effective electric field.
- 69 `ninv=0.5` Coefficient of effective electric field.
- 70 `ninvd=1.0e-9 1/V` Modification of `Ninv`.
- 71 `bb=2.0` High-field mobility degradation.
- 72 `vmax=7.0e6 cm/s` Maximum of electron saturation velocity.
- 73 `vover=0.01 cm^(voverp)`
Parameter for velocity overshoot.
- 74 `voverp=0.1` Lgate dependence of velocity overshoot.
- 75 `rpock1=0.001 V2 μm^(Rpocp2-1)/A^(Rpocp1)`
Pocket technology parameter for `Ids`.
- 76 `rpock2=0.1 V` Pocket technology parameter for `Ids`.
- 77 `rpocp1=1.0` Pocket technology parameter for `Ids`(HiSIM1.1.0).
- 78 `rpocp2=0.5` Pocket technology parameter for `Ids`(HiSIM1.1.0).

Channel Length Modulation Effects

- 79 `clm1=0.7` First parameter for CLM.
- 80 `clm2=2.0 1/m` Second parameter for CLM.
- 81 `clm3=1.0` Third parameter for CLM.

Substrate Current Effects

- 82 `sub1=10.0 1/V` First parameter for `Isub`.
- 83 `sub2=20.0 V` Second parameter for `Isub`.
- 84 `sub3=0.8` Third parameter for `Isub`.

Spectre Circuit Simulator Reference

Component Statements Part II

Gate Current Effects

85 $g_{leak1}=1.0e4 \text{ A}/(V^{(3/2)} c^{(1/2)})$
First gate current coefficient.

86 $g_{leak2}=2.0e7 \text{ 1}/(V^{(1/2)} c^{(3/2)} m)$
Second gate current coefficient.

87 $g_{leak3}=0.3$ Third gate current coefficient.

GIDL Current Effects

88 $gidl1=5.0e-6 \text{ A m}/(V^{(3/2)} c^{(1/2)})$
First parameter for GIDL.

89 $gidl2=1.0e6 \text{ 1}/(V^{(1/2)} c^{(3/2)} m)$
Second parameter for GIDL.

90 $gidl3=0.3$ Third parameter for GIDL.

Noise 1/f Effects

91 $nfalp=1.0e-16$ Flicker (1/f) noise contribution of the mobility fluctuation.

92 $nftrp=1.0e10$ Flicker (1/f) noise ratio of trap density to attenuation coefficient.

93 $cit=0.0$ Flicker (1/f) noise interface trapped carriers capacitance.

94 $kf=0$ Flicker (1/f) noise coefficient.

95 $af=1.0$ Flicker (1/f) noise exponent.

96 $ef=0.0$ Flicker (1/f) noise frequency exponent.

Symmetry for short-channel mosfet

97 $vzadd0=1.0e-2 \text{ V}$ Vzadd at Vds=0.

98 $pzadd0=5.0e-3 \text{ V}$ Pzadd at Vds=0.

Spectre Circuit Simulator Reference

Component Statements Part II

P-N junctions parameters

99	$j_{s0}=1.0e-4 \text{ A/m}^2$	Junction saturation current density.
100	$j_{s0sw}=0.0 \text{ A/m}$	Side-wall saturation current density.
101	$n_j=1.0$	Junction emission coefficient.
102	$n_{jsw}=1.0$	Junction emission coefficient (sidewall).
103	$x_{ti}=3.0$	Junction saturation current temperature exponent coefficient.
104	$c_j=5.0e-4 \text{ F/m}^2$	Bottom junction capacitance per unit area at zero bias.
105	$c_{jsw}=5e-10 \text{ F/m}$	Source/drain sidewall junction capacitance per unit length at zero bias.
106	$c_{jswg}=5e-10 \text{ F/m}$	Source/drain sidewall junction capacitance grading coefficient per unit length at zero bias.
107	$m_j=0.5$	Bulk junction bottom grading coefficient.
108	$m_{jsw}=0.33$	Source/drain sidewall junction capacitance grading coefficient.
109	$m_{jswg}=0.33$	Bottom junction capacitance grading coefficient.
110	$p_b=1.0 \text{ V}$	Bottom junction build-in potential.
111	$p_{bsw}=1.0 \text{ V}$	Source/drain sidewall junction build-in potential.
112	$p_{bswg}=1.0 \text{ V}$	Source/drain gate sidewall junction build-in potential.

Overlap capacitance parameters

113	$c_{gso} \text{ (F/m)}$	Gate-source overlap capacitance.
114	$c_{gdo} \text{ (F/m)}$	Gate-drain overlap capacitance.
115	$c_{gbo} \text{ (F/m)}$	Gate-bulk overlap capacitance.

Spectre Circuit Simulator Reference

Component Statements Part II

Auto Model Selector parameters

116	<code>wmax=1 m</code>	Maximum channel width for which the model is valid.
117	<code>wmin=0 m</code>	Minimum channel width for which the model is valid.
118	<code>lmax=1 m</code>	Maximum channel length for which the model is valid.
119	<code>lmin=0 m</code>	Minimum channel length for which the model is valid.

Output Parameters

1	<code>w_{eff} (m)</code>	Effective channel width.
2	<code>l_{eff} (m)</code>	Effective channel length.
3	<code>r_{seff} (Ω)</code>	Effective source resistance.
4	<code>r_{deff} (Ω)</code>	Effective drain resistance.

Operating-Point Parameters

1	<code>reversed</code>	Reverse mode indicator. Possible values are <code>no</code> or <code>yes</code> .
2	<code>ids (A)</code>	Resistive drain-to-source current.
3	<code>vgs (V)</code>	Gate-source voltage.
4	<code>vds (V)</code>	Drain-source voltage.
5	<code>vbs (V)</code>	Bulk-source voltage.
6	<code>vth (V)</code>	Threshold voltage.
7	<code>gm (S)</code>	Common-source transconductance.
8	<code>gds (S)</code>	Common-source output conductance.
9	<code>gmbs (S)</code>	Body-transconductance.
10	<code>Qb (Coul)</code>	Qb.

Spectre Circuit Simulator Reference

Component Statements Part II

11	Qd (Coul)	Qd.
12	Qg (Coul)	Qg.
13	Qs (Coul)	Qs.
14	cjd (F)	Drain-bulk junction capacitance.
15	cjs (F)	Source-bulk junction capacitance.
16	cgg (F)	dQg_dVg.
17	cgd (F)	dQg_dVd.
18	cgs (F)	dQg_dVs.
19	cgb (F)	dQg_dVbk.
20	cdg (F)	dQd_dVg.
21	cdd (F)	dQd_dVd.
22	cds (F)	dQd_dVs.
23	cdb (F)	dQd_dVb.
24	csg (F)	dQs_dVg.
25	csd (F)	dQs_dVd.
26	css (F)	dQs_dVs.
27	csb (F)	dQs_dVb.
28	cbg (F)	dQb_dVg.
29	cbd (F)	dQb_dVd.
30	cbs (F)	dQb_dVs.
31	cbb (F)	dQb_dVb.
32	id (A)	Resistive drain current.

Spectre Circuit Simulator Reference

Component Statements Part II

33	<code>is</code> (A)	Resistive source current.
34	<code>ibulk</code> (A)	Resistive bulk current.
35	<code>pwr</code> (W)	Power at op point.
36	<code>ps0</code> (V)	Surface potential at source side.
37	<code>ps1</code> (V)	Surface potential at drain side.
38	<code>pds</code> (V)	Delta between <code>ps1</code> and <code>ps0</code> .
39	<code>isub</code> (A)	Substrate current <code>Isub</code> .
40	<code>gbds</code> (S)	Substrate trans conductance (dI_{sub}/dV_{ds}).
41	<code>gbgs</code> (S)	Substrate trans conductance (dI_{sub}/dV_{gs}).
42	<code>gbbs</code> (S)	Substrate transconductance (dI_{sub}/dV_{bs}).

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

<code>Qb</code>	OP-10	<code>coisub</code>	M-9	<code>mueph1</code>	M-64	<code>rseff</code>	O-3
<code>Qd</code>	OP-11	<code>conois</code>	M-13	<code>mueph2</code>	M-55	<code>sc1</code>	M-48
<code>Qg</code>	OP-12	<code>coovlp</code>	M-12	<code>muesr0</code>	M-66	<code>sc2</code>	M-49
<code>Qs</code>	OP-13	<code>copprv</code>	M-16	<code>muesr1</code>	M-67	<code>sc3</code>	M-50
<code>ad</code>	I-4	<code>corsrd</code>	M-4	<code>muetmp</code>	M-65	<code>scp1</code>	M-51
<code>ad</code>	M-20	<code>csb</code>	OP-27	<code>ndep</code>	M-68	<code>scp2</code>	M-52
<code>af</code>	M-95	<code>csd</code>	OP-25	<code>nfalp</code>	M-91	<code>scp3</code>	M-53
<code>as</code>	M-19	<code>csg</code>	OP-24	<code>nftrp</code>	M-92	<code>sub1</code>	M-82
<code>as</code>	I-3	<code>css</code>	OP-26	<code>ninv</code>	M-69	<code>sub2</code>	M-83
<code>bb</code>	M-71	<code>dtemp</code>	I-8	<code>ninvd</code>	M-70	<code>sub3</code>	M-84

Spectre Circuit Simulator Reference Component Statements Part II

bgtmp1	M-38	dtemp	M-24	nj	M-101	temp	I-7
bgtmp2	M-39	ef	M-96	njsw	M-102	temp	M-23
cbb	OP-31	gbbs	OP-42	noise	M-3	tox	M-25
cbd	OP-29	gbds	OP-40	nsti	M-58	tpoly	M-29
cbg	OP-28	gbgs	OP-41	nsubc	M-30	type	M-1
cbs	OP-30	gds	OP-8	nsubp	M-31	vbs	OP-5
cdb	OP-23	gidl1	M-88	parl1	M-46	vds	OP-4
cdd	OP-21	gidl2	M-89	parl2	M-47	vds0	M-60
cdg	OP-20	gidl3	M-90	pb	M-110	version	M-2
cds	OP-22	gleak1	M-85	pbsw	M-111	vfbc	M-34
cgb	OP-19	gleak2	M-86	pbswg	M-112	vgs	OP-3
cgbo	M-115	gleak3	M-87	pd	M-22	vmax	M-72
cgd	OP-17	gm	OP-7	pd	I-6	vover	M-73
cgdo	M-114	gmbs	OP-9	pds	OP-38	voverp	M-74
cgg	OP-16	ibulk	OP-34	pgd1	M-43	vth	OP-6
cgs	OP-18	id	OP-32	pgd2	M-44	vzadd0	M-97
cgso	M-113	ids	OP-2	pgd3	M-45	w	M-17
cit	M-93	is	OP-33	ps	M-21	w	I-1
cj	M-104	isub	OP-39	ps	I-5	w0	M-56
cjd	OP-14	js0	M-99	ps0	OP-36	weff	O-1
cjs	OP-15	js0sw	M-100	ps1	OP-37	wfc	M-54
cjsw	M-105	kf	M-94	pwr	OP-35	wmax	M-116
cjswg	M-106	l	M-18	pzadd0	M-98	wmin	M-117
clm1	M-79	l	I-2	qme1	M-40	wsti	M-59
clm2	M-80	leff	O-2	qme2	M-41	wvthsc	M-57
clm3	M-81	lmax	M-118	qme3	M-42	xj	M-36
coadv	M-8	lmin	M-119	rd	M-33	xld	M-26
cocgbo	M-7	lp	M-35	rdef	O-4	xpolyd	M-28
cocgdo	M-6	mj	M-107	reversed	OP-1	xqy	M-37
cocgso	M-5	mjsw	M-108	rpock1	M-75	xti	M-103
cogidl	M-10	mjswg	M-109	rpock2	M-76	xwd	M-27
coiigs	M-11	muecb0	M-61	rpocp1	M-77		

Spectre Circuit Simulator Reference Component Statements Part II

coiprv	M-15	muecb1	M-62	rpocp2	M-78
coisti	M-14	mueph0	M-63	rs	M-32

HV MOS Transistor (hvmos)

Description

HV (High-Voltage) MOS transistor model is a deep submicron, high-voltage MOSFET model. It is based on the BSIM3v3 version 3.1. Major enhancements include current-crowding effect at high gate bias, asymmetric source-drain structure, mobility reduction, transconductance reduction under high Vgs at saturation region, forward and reverse mode, self-heating, and more flexible gate-dependent output characteristics. HVMOS can be used for high voltage IC design applications such as Flash memory with asymmetric LDD structures, LCD drivers, CCD, E2PROM and LDMOS applications.

Like BSIM3v3, the HVMOS transistor model also allows the binning option to achieve even higher accuracy. The binning equation is given by

$$P = P_0 + P_I / L_{eff} + P_w / W_{eff} + P_p / (L_{eff} * W_{eff})$$

Only the P0 parameters are listed. PI, Pw, and Pp are not shown but can be recognized. The names of PI, Pw, and Pp are identical to that of P0 but with a prefix of I, w, and p, respectively. HVMOS transistors require that you use a model statement.

This device is supported within altergroups.

Sample Instance Statement

```
m1 (1 2 0 0) hvmos w=1.5u l=1u ad=2.6p as=2.6p pd=6.6p ps=6.6p nrd=1.54 nrs=1.54
```

Sample Model Statement

```
model hvmos hvmos vtho=0.53 w0=2.14e-6 nlx=1.8e-7 nch=2.3e18 xj=0.22e-6 k1=0.48  
k2=-0.02 drout=1.1 rsh=10 cgso=2.4e-10 cgdo=2.4e-10
```

Instance Definition

```
Name d g s b ModelName parameter=value ...
```

Spectre Circuit Simulator Reference

Component Statements Part II

Instance Parameters

1	<code>w</code> (m)	Channel width.
2	<code>l</code> (m)	Channel length.
3	<code>as</code> (m ²)	Area of source diffusion.
4	<code>ad</code> (m ²)	Area of drain diffusion.
5	<code>ps</code> (m)	Perimeter of source diffusion.
6	<code>pd</code> (m)	Perimeter of drain diffusion.
7	<code>nrd</code> (m/m)	Number of squares of drain diffusion.
8	<code>nrs</code> (m/m)	Number of squares of source diffusion.
9	<code>ld</code> (m)	Length of drain diffusion region.
10	<code>ls</code> (m)	Length of source diffusion region.
11	<code>m=1</code>	Multiplicity factor (number of MOSFETs in parallel).
12	<code>region=triode</code>	Estimated operating region. Spectre outputs number (0-3) in a rawfile. Possible values are <code>off</code> , <code>triode</code> , <code>sat</code> , or <code>subth</code> .
13	<code>nqsmod</code>	NQS flag.
14	<code>trise</code>	Temperature rise from ambient.

Model Definition

```
model modelName hvmos parameter=value ...
```

Model Parameters

Device type parameters

1	<code>type=n</code>	Transistor type. Possible values are <code>n</code> or <code>p</code> .
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Spectre Circuit Simulator Reference

Component Statements Part II

- 2 `hvmosver=1.0` HVMOS Model version selector. The available versions are 1.0, 2.0.
- 3 `version=3.1` model version.

Threshold voltage parameters

- 4 `vtho (V)` Threshold voltage at zero body bias for long-channel devices. For enhancement-mode devices, $v_{tho} > 0$ for n-channel and $v_{tho} < 0$ for p-channel.
- 5 `k1=0.5 \sqrt{V}` Body-effect coefficient, default is 0.53 for `hvmosver` ≥ 2.0 .
- 6 `k2=-0.0186` Charge-sharing parameter.
- 7 `k3=80` Narrow width coefficient.
- 8 `k3b=0 1/V` Narrow width coefficient.
- 9 `w0=2.5e-6 m` Narrow width coefficient.
- 10 `nlx=1.74e-7 m` Lateral nonuniform doping coefficient.
- 11 `gamma1=1.0 \sqrt{V}` Body-effect coefficient near the surface.
- 12 `gamma2=0 \sqrt{V}` Body-effect coefficient in the bulk.
- 13 `vbx=-3 V` Threshold voltage transition body voltage.
- 14 `vbm=-3 V` Maximum applied body voltage.
- 15 `dvt0=2.2` First coefficient of short-channel effects.
- 16 `dvt1=0.53` Second coefficient of short-channel effects.
- 17 `dvt2=-0.032 1/V` Body-bias coefficient of short-channel effects.
- 18 `a0f=1` Forward nonuniform depletion width effect coefficient.
- 19 `a0r=a0f` Reverse nonuniform depletion width effect coefficient.
- 20 `b0=0 m` Bulk charge coefficient due to narrow width effect.

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Component Statements Part II

21	$b1=0$ m	Bulk charge coefficient due to narrow width effect.
22	$a1=0$	No-saturation coefficient.
23	$a2=1$	No-saturation coefficient.
24	$ags=0$ F/m ² V	Gate-bias dependence of A_{bulk} .
25	$ketaf=-0.047$ 1/V	Body-bias coefficient for non-uniform depletion width effect.
26	$ketar=ketaf$ 1/V	Reverse body-bias coefficient for non-uniform depletion width effect.

Process parameters

27	$n_{sub}=6e16$ cm ⁻³	Substrate doping concentration.
28	$n_{ch}=1.7e17$ cm ⁻³	Peak channel doping concentration.
29	$x_j=0.15e-6$ m	Source/drain junction depth.
30	$l_{int}=0$ m	Lateral diffusion for one side.
31	$w_{int}=0$ m	Width reduction for one side.
32	$l1=0$ m	Length dependence of δL .
33	$l1n=1$	Length exponent of δL .
34	$lw=0$ m	Width dependence of δL .
35	$lwn=1$	Width exponent of δL .
36	$lw1=0$ m ²	Area dependence of δL .
37	$l_{min}=0$ m	The minimum channel length for which the model is still valid.
38	$l_{max}=1$ m	The maximum channel length for which the model is still valid.
39	$w1=0$ m	Length dependence of δW .
40	$w1n=1$	Length exponent of δW .

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Component Statements Part II

41	<code>ww=0</code>	m	Width dependence of delta W.
42	<code>wwn=1</code>		Width exponent of delta W.
43	<code>wwl=0</code>	m ²	Area dependence of delta W.
44	<code>wmin=0</code>	m	The minimum channel width for which the model is still valid.
45	<code>wmax=1</code>	m	The maximum channel width for which the model is still valid.
46	<code>dwg=0</code>	m/v	Gate-bias dependence of channel width.
47	<code>dwb=0</code>	m/√v	Body-bias dependence of channel width.
48	<code>tox=1.5e-8</code>	m	Gate oxide thickness.
49	<code>xt=1.55e-7</code>	m	Doping depth.
50	<code>rd0=0</code>	Ω	Fixed drain resistance.
51	<code>rs0=0</code>	Ω	Fixed source resistance.
52	<code>rdw=0</code>	Ω μm	Width dependence of drain resistance.
53	<code>rsw=0</code>	Ω μm	Width dependence of source resistance.
54	<code>rdsw=0</code>	Ω μm	Width dependence of drain-source resistance.
55	<code>prwb=0</code>	1/√v	Body-effect coefficient for Rds.
56	<code>prwg=0</code>	1/√v	Gate-effect coefficient for Rds.
57	<code>wr=1</code>		Width offset for parasitic resistance.
58	<code>binunit=2</code>		Bin parameter unit selector. 1 for microns and 2 for meters.

Mobility parameters

59	<code>mobmod=1</code>		Mobility model selector.
60	<code>u0f=670</code>	cm ² /V s	Forward low-field surface mobility at <code>t_{nom}</code> . Default is 250 for PMOS.

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Component Statements Part II

- 61 $u0r=u0f \text{ cm}^2/\text{V s}$ Reverse low-field surface mobility at t_{nom} .
- 62 $vsatf=8e4 \text{ m/s}$ Forward carrier saturation velocity at t_{nom} .
- 63 $dvsatf=0 \text{ m/s}$ Forward gate-bias dependence of saturation velocity.
- 64 $dvsatbf=0 \text{ m/s}$ Forward body-bias dependence of saturation velocity.
- 65 $vsatr=vsatf \text{ m/s}$ Reverse carrier saturation velocity at t_{nom} .
- 66 $dvsatr=dvsatf \text{ m/s}$ Reverse gate-bias dependence of saturation velocity.
- 67 $dvsatbr=dvsatbf \text{ m/s}$
Reverse body-bias dependence of saturation velocity.
- 68 $uaf=2.25e-9 \text{ m/v}$ Forward first-order mobility reduction coefficient.
- 69 $ubf=5.87e-19 \text{ m}^2/\text{v}^2$
Forward second-order mobility reduction coefficient.
- 70 $ucf=-4.65e-11 \text{ m/v}^2$
Forward body-bias dependence of mobility. Default is -0.046 and unit is 1/V for $mobmod=3$.
- 71 $udf=0 \text{ m/v}^2$ Forward source-resistance dependence of mobility.
- 72 $uar=uaf \text{ m/v}$ Reverse first-order mobility reduction coefficient.
- 73 $ubr=ubf \text{ m}^2/\text{v}^2$ Reverse second-order mobility reduction coefficient.
- 74 $ucr=ucf \text{ m/v}^2$ Reverse body-bias dependence of mobility.
- 75 $udr=udf \text{ m/v}^2$ Reverse source-resistance dependence of mobility.

Output resistance parameters

- 76 $drout=0.56$ DIBL effect on output resistance coefficient.
- 77 $pclmf=1.3$ Forward channel length modulation coefficient.
- 78 $pclmr=pclmf$ Reverse channel length modulation coefficient.
- 79 $pdiblc1f=0.39$ Forward first coefficient of drain-induced barrier lowering.

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- 80 `pdiblc1r=pdiblc1f` Reverse first coefficient of drain-induced barrier lowering.
- 81 `pdiblc2f=8.6e-3` Forward second coefficient of drain-induced barrier lowering.
- 82 `pdiblc2r=pdiblc2f` Reverse second coefficient of drain-induced barrier lowering.
- 83 `pdiblc1bf=0 1/V` Body-effect coefficient for DIBL.
- 84 `pdiblc1br=pdiblc1bf 1/V` Reverse body-effect coefficient for DIBL.
- 85 `pscbe1f=4.24e8 V/m` First coefficient of substrate current body effect.
- 86 `pscbe2f=1e-5 m/v` Second coefficient of substrate current body effect.
- 87 `pscbe3f=0 V/m` Third coefficient of substrate current body effect.
- 88 `pscbe1r=pscbe1f V/m` Reverse first coefficient of substrate current body effect.
- 89 `pscbe2r=pscbe2f m/v` Reverse second coefficient of substrate current body effect.
- 90 `pclmgf=0` Forward gate dependence of V_{a1m} .
- 91 `pclmgr=pc1mgf` Reverse gate dependence of V_{a1m} .
- 92 `pclmbf=0` Forward body dependence of V_{a1m} .
- 93 `pclmbr=pc1mbf` Reverse body dependence of V_{a1m} .
- 94 `pdiblgf=0` Forward gate dependence of V_{d1bl} .
- 95 `pdiblgr=pdiblgf` Reverse gate dependence of V_{d1bl} .
- 96 `delta=0.01 V` Effective drain voltage smoothing parameter.

Subthreshold parameters

- 97 `cdsc=2.4e-4 F/m2` Source/drain and channel coupling capacitance.

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Component Statements Part II

98	$cdscb=0 \text{ F/m}^2 \text{ V}$	Body-bias dependence of $cdsc$.
99	$cdscd=0 \text{ F/m}^2 \text{ V}$	Drain-bias dependence of $cdsc$.
100	$nfactor=1$	Subthreshold swing coefficient.
101	$cit=0 \text{ F}$	Interface trap parameter for subthreshold swing.
102	$voff=-0.08 \text{ V}$	Threshold voltage offset.
103	$dsub=drout$	DIBL effect in subthreshold region.
104	$eta0f=0.08$	DIBL coefficient subthreshold region.
105	$etabf=-0.07 \text{ 1/V}$	Body-bias dependence of $et0$.
106	$eta0r=eta0f$	Reverse DIBL coefficient subthreshold region.
107	$etabr=etabf \text{ 1/V}$	Body-bias dependence of $eta0r$.

Substrate current parameters

108	$alpha0=0 \text{ m/v}$	Substrate current impact ionization coefficient.
109	$alpha1=0 \text{ 1/V}$	substrate current model parameter.
110	$beta0=30 \text{ 1/V}$	Substrate current impact ionization exponent.

Parasitic resistance parameters

111	$rsh=0 \text{ } \Omega/\text{sqr}$	Source/drain diffusion sheet resistance.
112	$rs=0 \text{ } \Omega$	Source resistance.
113	$rd=0 \text{ } \Omega$	Drain resistance.
114	$lgcs=0 \text{ m}$	Gate-to-contact length of source side.
115	$lgcd=0 \text{ m}$	Gate-to-contact length of drain side.
116	$rsc=0 \text{ } \Omega$	Source contact resistance.

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Component Statements Part II

117	$r_{dc}=0 \ \Omega$	Drain contact resistance.
118	$r_{ss}=0 \ \Omega \ m$	Scalable source resistance.
119	$r_{dd}=0 \ \Omega \ m$	Scalable drain resistance.
120	$s_c=\infty \ m$	Spacing between contacts.
121	$l_{dif}=0 \ m$	Lateral diffusion beyond the gate.
122	$h_{dif}=0 \ m$	Length of heavily doped diffusion .
123	$minr=0.1 \ \Omega$	Minimum source/drain resistance.

Junction diode model parameters

124	$j_s \ (A/m^2)$	Bulk junction reverse saturation current density.
125	$j_{sw}=0 \ A/m$	Sidewall junction reverse saturation current density.
126	$i_s=1e-14 \ A$	Bulk junction reverse saturation current.
127	$n=1$	Junction emission coefficient.
128	$dskip=yes$	Use simple piece-wise linear model for diode currents below $0.1*i_{abstol}$. Possible values are <code>no</code> or <code>yes</code> .
129	$imelt='imaxA'$	Explosion current, default is 0.1 for <code>hvmosver >=2.0</code> .
130	$ijth='imax' \ A$	Junction threshold current.
131	$jmelt='jmaxA/m'^2$	Explosion current density.

Overlap capacitance parameters

132	$c_{gso} \ (F/m)$	Gate-source overlap capacitance.
133	$c_{gdo} \ (F/m)$	Gate-drain overlap capacitance.
134	$c_{gbo} \ (F/m)$	Gate-bulk overlap capacitance.

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Component Statements Part II

135	$meto=0$	m	Metal overlap in fringing field.
136	$cgs1=0$	F/m	Gate-source overlap capacitance in LDD region.
137	$cgd1=0$	F/m	Gate-drain overlap capacitance in LDD region.
138	$ckappa=0.6$		Overlap capacitance fitting parameter.
139	$deltaacc=0.1$	V	Capacitance smoothing parameter.

Junction capacitance model parameters

140	$cbs=0$	F	Bulk-source zero-bias junction capacitance.
141	$cbd=0$	F	Bulk-drain zero-bias junction capacitance.
142	$cj=5e-4$	F/m ²	Zero-bias junction bottom capacitance density.
143	$mj=1/2$		Bulk junction bottom grading coefficient.
144	$pb=0.8$	V	Bulk junction built-in potential, default is 1.0 for hvmosver ≥ 2.0 .
145	$fc=0.5$		Forward-bias depletion capacitance threshold.
146	$cjsw=5e-10$	F/m	Zero-bias junction sidewall capacitance density.
147	$mjsw=1/3$		Bulk junction sidewall grading coefficient.
148	$pbsw=0.8$	V	Side-wall junction built-in potential, default is 1.0 for hvmosver ≥ 2.0 .
149	$fcsw=0.5$		Side-wall forward-bias depletion capacitance threshold.
150	$cjswg=cjsw$	F/m	Zero-bias gate-side junction capacitance density.
151	$mjswg=mjsw$		Gate-side junction grading coefficient.
152	$pbswg=pbsw$	V	Gate-side junction built-in potential.

Charge model selection parameters

153	$capmod=2$		Intrinsic charge model.
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Spectre Circuit Simulator Reference

Component Statements Part II

154	<code>nqsmod=0</code>	Non-quasi static model selector. Set to 1 to turn on nqs.
155	<code>dwc=wint m</code>	Delta W for capacitance model.
156	<code>dlc=lint m</code>	Delta L for capacitance model.
157	<code>clc=1e-7 m</code>	Intrinsic capacitance fitting parameter.
158	<code>cle=0.6</code>	Intrinsic capacitance fitting parameter.
159	<code>cf (F/m)</code>	Fringe capacitance parameter.
160	<code>a0cvf=a0f</code>	A0 for C-V calculation.
161	<code>a0cvr=a0r</code>	Reverse A0 for C-V calculation.
162	<code>qgvd0f=1</code>	Cgd fitting parameter.
163	<code>qgvd0r=qgvd0f</code>	Reverse Cgd fitting parameter.
164	<code>elm=5</code>	Elmore constant of the channel.
165	<code>vfbcv=-1</code>	Flat-band voltage for capmod=0.
166	<code>acde=0.5 1/V</code>	Exponential coefficient for finite charge thickness.
167	<code>moin=15 1/V</code>	Coefficient for Vgs dependent surface potential.
168	<code>xpart=0</code>	Drain/source channel charge partition in saturation for BSIM charge model, use 0.0 for 40/60, 0.5 for 50/50, or 1.0 for 0/100.

Default instance parameters

169	<code>w=5e-6 m</code>	Default channel width.
170	<code>l=5e-6 m</code>	Default channel length.
171	<code>as=0 m²</code>	Default area of source diffusion.
172	<code>ad=0 m²</code>	Default area of drain diffusion.
173	<code>ps=0 m</code>	Default perimeter of source diffusion.

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Component Statements Part II

174	<code>pd=0 m</code>	Default perimeter of drain diffusion.
175	<code>nrd=0 m/m</code>	Default number of squares of drain diffusion.
176	<code>nrs=0 m/m</code>	Default number of squares of source diffusion.
177	<code>xw=0 m</code>	Width variation due to masking and etching.
178	<code>xl=0 m</code>	Length variation due to masking and etching.

Temperature effects parameters

179	<code>tnom (C)</code>	Parameters measurement temperature. Default set by <code>options</code> .
180	<code>trise=0 C</code>	Temperature rise from ambient.
181	<code>tlev=0</code>	DC temperature selector.
182	<code>tlevc=0</code>	AC temperature selector.
183	<code>eg=1.12452 V</code>	Energy band gap.
184	<code>gap1=7.02e-4 V/C</code>	Band gap temperature coefficient.
185	<code>gap2=1108 C</code>	Band gap temperature offset.
186	<code>kt1=-0.11 V</code>	Temperature coefficient for threshold voltage.
187	<code>kt1l=0 v m</code>	Temperature coefficient for threshold voltage.
188	<code>kt2=0.022</code>	Temperature coefficient for threshold voltage.
189	<code>atf=3.3e4 m/s</code>	Temperature coefficient for <code>vsatf</code> .
190	<code>atr=atf m/s</code>	Temperature coefficient for <code>vsatr</code> .
191	<code>at1f=0 m/s</code>	Temperature coefficient for <code>dvsatf</code> .
192	<code>at1r=at1f m/s</code>	Temperature coefficient for <code>dvsatr</code> .
193	<code>ua1f=4.31e-9 m/v</code>	Temperature coefficient for <code>uaf</code> .

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194	$ub1f=-7.61e-18 \text{ m}^2/\text{v}^2$	Temperature coefficient for ubf .
195	$uc1f=-5.5e-11 \text{ m}/\text{v}^2$	Temperature coefficient for ucf . Default is -0.056 for $mobmod=3$.
196	$ud1f=0 \text{ m}/\text{v}^2$	Temperature coefficient for udf .
197	$ualr=ualf \text{ m}/\text{v}$	Temperature coefficient for uar .
198	$ub1r=ub1f \text{ m}^2/\text{v}^2$	Temperature coefficient for ubr .
199	$uc1r=uc1f \text{ m}/\text{v}^2$	Temperature coefficient for ucr .
200	$ud1r=0 \text{ m}/\text{v}^2$	Temperature coefficient for udr .
201	$rth=0 \text{ } \Omega$	Self-heating thermal resistance.
202	$rthg=0 \text{ 1}/\text{V}$	Gate-effect coefficient for Rth .
203	$rthb=0 \text{ 1}/\sqrt{\text{v}}$	Body-effect coefficient for Rth .
204	$prt=0 \text{ } \Omega$	Temperature coefficient for Rds .
205	$trs=0 \text{ 1}/\text{C}$	Temperature parameter for source resistance.
206	$trd=0 \text{ 1}/\text{C}$	Temperature parameter for drain resistance.
207	$ute=-1.5$	Mobility temperature exponent.
208	$xTi=3$	Saturation current temperature exponent.
209	$ptc=0 \text{ V}/\text{C}$	Surface potential temperature coefficient.
210	$tcv=0 \text{ V}/\text{C}$	Threshold voltage temperature coefficient.
211	$pta=0 \text{ V}/\text{C}$	Junction potential temperature coefficient.
212	$tpb=0 \text{ V}/\text{C}$	Junction potential temperature coefficient.
213	$ptp=0 \text{ V}/\text{C}$	Sidewall junction potential temperature coefficient.
214	$tpbsw=0 \text{ V}/\text{C}$	Sidewall junction potential temperature coefficient.

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Component Statements Part II

215	<code>cta=0 1/C</code>	Junction capacitance temperature coefficient.
216	<code>tcj=0 1/C</code>	Junction capacitance temperature coefficient.
217	<code>ctp=0 1/C</code>	Sidewall junction capacitance temperature coefficient.
218	<code>tcjsw=0 1/C</code>	Sidewall junction capacitance temperature coefficient.
219	<code>tcjswg=0 1/C</code>	Temperature coefficient for <code>cjswg</code> .
220	<code>tpbswg=0 V/C</code>	Temperature coefficient for <code>pbswg</code> .

Noise model parameters

221	<code>noimod=1</code>	Noise model selector.
222	<code>kf=0</code>	Flicker (1/f) noise coefficient.
223	<code>af=1</code>	Flicker (1/f) noise exponent.
224	<code>ef=1</code>	Flicker (1/f) noise frequency exponent.
225	<code>noia=1e20</code>	Oxide trap density coefficient. Default is 9.9e18 for pmos.
226	<code>noib=5e4</code>	Oxide trap density coefficient. Default is 2.4e3 for pmos.
227	<code>noic=-1.4e-8</code>	Oxide trap density coefficient. Default is 1.4e-8 for pmos. Default is -1.4-12 for hvmosver >=2.0.

Operating region warning control parameters

228	<code>alarm=none</code>	Forbidden operating region. Possible values are <code>none</code> , <code>off</code> , <code>triode</code> , <code>sat</code> , <code>subth</code> , or <code>rev</code> .
229	<code>imax=1 A</code>	Maximum allowable current, default is 0.1 for hvmosver >=2.0 .
230	<code>jmax=1e8 A/m²</code>	Maximum allowable current density.
231	<code>bvj=∞ V</code>	Junction reverse breakdown voltage.
232	<code>vbox=1e9 tox V</code>	Oxide breakdown voltage.

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Component Statements Part II

Length dependent parameters (Not listed)

Width dependent parameters (Not listed)

Cross-term dependent parameters (Not listed)

The i_{\max} (j_{\max}) parameter is used to aid convergence and prevent numerical overflow. The junction characteristics of the FET are accurately modeled for current (density) up to i_{\max} (j_{\max}). For currents (density) above i_{\max} (j_{\max}), the junction is modeled as a linear resistor and a warning is printed.

Output Parameters

1	<code>weff</code> (m)	Effective channel width.
2	<code>leff</code> (m)	Effective channel length.
3	<code>rseff</code> (Ω)	Effective source resistance.
4	<code>rdeff</code> (Ω)	Effective drain resistance.

Operating-Point Parameters

1	<code>type=n</code>	Transistor type. Possible values are <code>n</code> or <code>p</code> .
2	<code>region=triode</code>	Estimated operating region. Spectre outputs number (0-3) in a rawfile. Possible values are <code>off</code> , <code>triode</code> , <code>sat</code> , or <code>subth</code> .
3	<code>reversed</code>	Reverse mode indicator. Possible values are <code>no</code> or <code>yes</code> .
4	<code>ids</code> (A)	Resistive drain-to-source current.
5	<code>vgs</code> (V)	Gate-source voltage.
6	<code>vds</code> (V)	Drain-source voltage.
7	<code>vbs</code> (V)	Bulk-source voltage.

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8	<code>vth</code> (V)	Threshold voltage.
9	<code>vdsat</code> (V)	Drain-source saturation voltage.
10	<code>gm</code> (S)	Common-source transconductance.
11	<code>gds</code> (S)	Common-source output conductance.
12	<code>gmbs</code> (S)	Body-transconductance.
13	<code>betaeff</code> (A/V ²)	Effective beta.
14	<code>cjd</code> (F)	Drain-bulk junction capacitance.
15	<code>cjs</code> (F)	Source-bulk junction capacitance.
16	<code>cgg</code> (F)	Cgg.
17	<code>cgd</code> (F)	Cgd.
18	<code>cgs</code> (F)	Cgs.
19	<code>cgb</code> (F)	Cgb.
20	<code>cdg</code> (F)	Cdg.
21	<code>cdd</code> (F)	Cdd.
22	<code>cds</code> (F)	Cds.
23	<code>cdb</code> (F)	Cdb.
24	<code>csg</code> (F)	Csg.
25	<code>csd</code> (F)	Csd.
26	<code>css</code> (F)	Css.
27	<code>csb</code> (F)	Csb.
28	<code>cbg</code> (F)	Cbg.
29	<code>cbd</code> (F)	Cbd.

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Component Statements Part II

30	cbs (F)	Cbs.
31	cbb (F)	Cbb.
32	ron (Ω)	On-resistance.
33	id (A)	Resistive drain current.
34	ibulk (A)	Resistive bulk current.
35	pwr (W)	Power at op point.
36	gmoverid (1/V)	Gm/Ids.

Parameter Index

In the following index, I refers to instance parameters, M refers to the model parameters section, O refers to the output parameters section, and OP refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of M-35 means the 35th model parameter.

a0cvf	M-160	dvsatbr	M-67	moin	M-167	sc	M-120
a0cvr	M-161	dvsatf	M-63	n	M-127	tcj	M-216
a0f	M-18	dvsatr	M-66	nch	M-28	tcjsw	M-218
a0r	M-19	dvt0	M-15	nfactor	M-100	tcjswg	M-219
a1	M-22	dvt1	M-16	nlx	M-10	tcv	M-210
a2	M-23	dvt2	M-17	noia	M-225	tlev	M-181
acde	M-166	dwb	M-47	noib	M-226	tlevc	M-182
ad	I-4	dwc	M-155	noic	M-227	tnom	M-179
ad	M-172	dwg	M-46	noimod	M-221	tox	M-48
af	M-223	ef	M-224	nqsmod	I-13	tpb	M-212
ags	M-24	eg	M-183	nqsmod	M-154	tpbsw	M-214
alarm	M-228	elm	M-164	nrd	M-175	tpbswg	M-220
alpha0	M-108	eta0f	M-104	nrd	I-7	trd	M-206
alpha1	M-109	eta0r	M-106	nrs	M-176	trise	I-14

Spectre Circuit Simulator Reference Component Statements Part II

as	I-3	etabf	M-105	nrs	I-8	trise	M-180
as	M-171	etabr	M-107	nsub	M-27	trs	M-205
atlf	M-191	fc	M-145	pb	M-144	type	OP-1
atlr	M-192	fcs w	M-149	p bsw	M-148	type	M-1
atf	M-189	gamma1	M-11	p bswg	M-152	u0f	M-60
atr	M-190	gamma2	M-12	p clmbf	M-92	u0r	M-61
b0	M-20	gap1	M-184	p clmbr	M-93	ualf	M-193
b1	M-21	gap2	M-185	p clmf	M-77	ualr	M-197
beta0	M-110	gds	OP-11	p clmgf	M-90	uaf	M-68
betaeff	OP-13	gm	OP-10	p clmgr	M-91	uar	M-72
binunit	M-58	gmbs	OP-12	p clmr	M-78	ublf	M-194
bvj	M-231	gmoverid	OP-36	pd	M-174	ublr	M-198
capmod	M-153	hdif	M-122	pd	I-6	ubf	M-69
cbb	OP-31	hvmosver	M-2	pdiblc1f	M-79	ubr	M-73
cbd	M-141	ibulk	OP-34	pdiblc1r	M-80	uclf	M-195
cbd	OP-29	id	OP-33	pdiblc2f	M-81	uclr	M-199
cbg	OP-28	ids	OP-4	pdiblc2r	M-82	ucf	M-70
cbs	OP-30	ijth	M-130	pdiblcbf	M-83	ucr	M-74
cbs	M-140	imax	M-229	pdiblcbr	M-84	udlf	M-196
cdb	OP-23	imelt	M-129	pdiblgf	M-94	udlr	M-200
cdd	OP-21	is	M-126	pdiblgr	M-95	udf	M-71
cdg	OP-20	jmax	M-230	prt	M-204	udr	M-75
cds	OP-22	jmelt	M-131	prwb	M-55	ute	M-207
cdsc	M-97	js	M-124	prwg	M-56	vbm	M-14
cdscb	M-98	jsw	M-125	ps	I-5	vbox	M-232
cdscd	M-99	k1	M-5	ps	M-173	vbs	OP-7
cf	M-159	k2	M-6	pscbelf	M-85	vbx	M-13
cgb	OP-19	k3	M-7	pscbelr	M-88	vds	OP-6
cgbo	M-134	k3b	M-8	pscbe2f	M-86	vdsat	OP-9
cgd	OP-17	ketaf	M-25	pscbe2r	M-89	version	M-3
cgdl	M-137	ketar	M-26	pscbeg	M-87	vfbcv	M-165
cgdo	M-133	kf	M-222	pta	M-211	vgs	OP-5

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cgg	OP-16	kt1	M-186	ptc	M-209	voff	M-102
cgs	OP-18	kt11	M-187	ptp	M-213	vsatf	M-62
cgs1	M-136	kt2	M-188	pwr	OP-35	vsatr	M-65
cgs0	M-132	l	M-170	qgvd0f	M-162	vth	OP-8
cit	M-101	l	I-2	qgvd0r	M-163	vtho	M-4
cj	M-142	ld	I-9	rd	M-113	w	M-169
cjd	OP-14	ldif	M-121	rd0	M-50	w	I-1
cjs	OP-15	leff	O-2	rdc	M-117	w0	M-9
cjsw	M-146	lgcd	M-115	rdd	M-119	weff	O-1
cjswg	M-150	lgcs	M-114	rdeff	O-4	wint	M-31
ckappa	M-138	lint	M-30	rdsw	M-54	wl	M-39
clc	M-157	ll	M-32	rdw	M-52	wln	M-40
cle	M-158	lln	M-33	region	I-12	wmax	M-45
csb	OP-27	lmax	M-38	region	OP-2	wmin	M-44
csd	OP-25	lmin	M-37	reversed	OP-3	wr	M-57
csg	OP-24	ls	I-10	ron	OP-32	ww	M-41
css	OP-26	lw	M-34	rs	M-112	ww1	M-43
cta	M-215	lw1	M-36	rs0	M-51	wwn	M-42
ctp	M-217	lwn	M-35	rsc	M-116	xj	M-29
delta	M-96	m	I-11	rseff	O-3	x1	M-178
deltaacc	M-139	meto	M-135	rsh	M-111	xpart	M-168
dlc	M-156	minr	M-123	rss	M-118	xt	M-49
drout	M-76	mj	M-143	rsw	M-53	xti	M-208
dskip	M-128	mjsw	M-147	rth	M-201	xw	M-177
dsub	M-103	mjswg	M-151	rthb	M-203		
dvsatbf	M-64	mobmod	M-59	rthg	M-202		

Two Terminal Inductor (inductor)

Description

The inductance of this component can be a function of temperature or branch current. If you do not specify the inductance in the instance statement, it is taken from the model.

This device is supported within altergroups.

If the polynomial coefficients vector (coeffs=[c1 c2 ...]) is specified, the inductor is nonlinear and the inductance is

$$L(I) = L(\text{inst}) * (1 + c1 * I + c2 * I^2 + \dots).$$

The branch flux as a function of current is

$$\text{Flux}(I) = L(\text{inst}) * I * (1 + 1/2 * c1 * I + 1/3 * c2 * I^2 + \dots)$$

where c_k is the k th entry in the coefficient vector.

The value of the inductor as a function of the temperature is given by:

$$L(T) = L(\text{tnom}) * [1 + tc1 * (T - \text{tnom}) + tc2 * (T - \text{tnom})^2].$$

where

$$T = \text{trise}(\text{inst}) + \text{temp}$$

if $\text{trise}(\text{inst})$ is given, and

$$T = \text{trise}(\text{model}) + \text{temp}$$

otherwise.

Sample Instance Statement

without model:

```
133 (0 net29) inductor l=10e-9 r=1 m=1
```

with model:

```
133 (0 net29) ind l=10e-9 r=1 m=1
```

Spectre Circuit Simulator Reference

Component Statements Part II

Sample Model Statement

```
model ind inductor l=6e-9 r=1 tc1=1e-12 tc2=1e-12 tnom=25
```

Instance Definition

```
Name 1 2 ModelName parameter=value ...
```

```
Name 1 2 inductor parameter=value ...
```

Instance Parameters

1	<code>l</code> (H)	Inductance.
2	<code>r</code> (Ω)	Resistance.
3	<code>m=1</code>	Multiplicity factor.
4	<code>trise</code>	Temperature rise from ambient.
5	<code>ic</code> (A)	Initial condition.
6	<code>isnoisy=yes</code>	Should inductor resistance generate noise. Possible values are <code>no</code> or <code>yes</code> .

Model Definition

```
model modelName inductor parameter=value ...
```

Model Parameters

1	<code>l=0</code> H	Default inductance.
2	<code>r=0</code> Ω	Default resistance.
3	<code>tc1=0</code> 1/C	Linear temperature coefficient.
4	<code>tc2=0</code> C ⁻²	Quadratic temperature coefficient.
5	<code>trise=0</code> C	Default <code>trise</code> value for instance.
6	<code>tnom</code> (C)	Parameters measurement temperature. Default set by <code>options</code> .
7	<code>rforce=1e9</code> Ω^2	Resistance used when forcing nodesets and initial conditions.

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Component Statements Part II

8 `coeffs=[...]` Vector of polynomial inductance coefficients.

9 `scalei=1` Inductance scaling factor.

Noise model parameters

10 `kf=0` Flicker (1/f) noise coefficient.

11 `af=2` Flicker (1/f) noise exponent.

Output Parameters

1 `indefeff (H)` Effective inductance.

Operating-Point Parameters

1 `ind (H)` Inductance at operating point.

2 `i (A)` Current at operating point.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

<code>af</code>	M-11	<code>indefeff</code>	O-1	<code>m</code>	I-3	<code>tc1</code>	M-3
<code>coeffs</code>	M-8	<code>isnoisy</code>	I-6	<code>r</code>	M-2	<code>tc2</code>	M-4
<code>i</code>	OP-2	<code>kf</code>	M-10	<code>r</code>	I-2	<code>tnom</code>	M-6
<code>ic</code>	I-5	<code>l</code>	M-1	<code>rforce</code>	M-7	<code>trise</code>	I-4
<code>ind</code>	OP-1	<code>l</code>	I-1	<code>scalei</code>	M-9	<code>trise</code>	M-5

Interconnect Capacitance (intcap)

Description

Intcap is a model for the calculation of the interconnect capacitance, which takes into account the local layer composition and the tracks spacing width. It is described in the Philips MOST Modelbook (Dec.96) as INTCAP model.

(c) Philips Electronics N.V. 1993,1996

The model is extended by the device parameters `lxbelps`, `lxbelin` and `lxbelins`, according to a specification by H.Okel (I&A Hamburg).

This device is supported within altergroups.

This device is dynamically loaded from the shared object `/vobs/spectre_dev/tools.sun4v/spectre/lib/cmi/3.0.doc/libphilips_sh.so`

Sample Instance Statement

```
intc (net9 net12) intconcap m=1 ael=2.5e-15 ain=2e-15 aps=1.8e-15
```

Sample Model Statement

```
model intconcap intcap cbps=1.5e-13 cebpsm=0.9e-15 cebpsi=0.83e-15 cbin=1.45e-13  
cbins=1.4e-13
```

Instance Definition

```
Name n1 n2 ModelName parameter=value ...
```

Instance Parameters

- | | | |
|---|-------------------------------------|--|
| 1 | <code>m=1</code> | Multiplicity factor. |
| 2 | <code>ael=0.0 m²</code> | The common area of EL track of the reference electrode. |
| 3 | <code>ain=0.0 m²</code> | The common area of IN track of the reference electrode. |
| 4 | <code>ains=0.0 m²</code> | The common area of INS track of the reference electrode. |
| 5 | <code>aps=0.0 m²</code> | The common area of PS track of the reference electrode. |

Spectre Circuit Simulator Reference

Component Statements Part II

6	lbel=0.0 m	The sum of periphery length of EL-segments common to node n2 downwards.
7	lbin=0.0 m	The sum of periphery length of IN-segments to node n2 downwards.
8	lbins=0.0 m	The sum of periphery length of INS-segments common to node n2 downwards.
9	lbsps=0.0 m	The sum of periphery length of PS-segments common to node n2 downwards.
10	lfbel=0.0 m	The sum of periphery length-factor products EL downwards.
11	lfbins=0.0 m	The sum of periphery length-factor products IN downwards.
12	lfbins=0.0 m	The sum of periphery length-factor products INS downwards.
13	lfbps=0.0 m	The sum of periphery length-factor products PS downwards.
14	lftel=0.0 m	The sum of periphery length-factor products EL upwards.
15	lftin=0.0 m	The sum of periphery length-factor products IN upwards.
16	lftins=0.0 m	The sum of periphery length-factor products INS upwards.
17	lftps=0.0 m	The sum of periphery length-factor products PS upwards.
18	ltel=0.0 m	The sum of periphery length of EL-segments common to node n2 upwards.
19	ltin=0.0 m	The sum of periphery length of IN-segments common to node n2 upwards.
20	ltins=0.0 m	The sum of periphery length of INS-segments common to node n2 upwards.
21	ltps=0.0 m	The sum of periphery length of PS-segments common to node n2 upwards.
22	ldsel=0.0 m	The sum of Li/Si quotients for EL tracks.
23	ldsins=0.0 m	The sum of Li/Si quotients for IN tracks.

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Component Statements Part II

24	<code>ldsins=0.0 m</code>	The sum of Li/Si quotients for <code>INS</code> tracks.
25	<code>ldsps=0.0 m</code>	The sum of Li/Si quotients for <code>PS</code> tracks.
26	<code>lxbinsps=0.0 m</code>	The sum of Li/Si quotients for an <code>IN</code> track in parallel with an <code>PS</code> track.
27	<code>lxbinsin=0.0 m</code>	The sum of Li/Si quotients for an <code>INS</code> track in parallel with an <code>IN</code> track.
28	<code>lxbinsps=0.0 m</code>	The sum of Li/Si quotients for an <code>INS</code> track in parallel with an <code>PS</code> track.
29	<code>lxbelps=0.0 m</code>	The sum of Li/Si quotients for an <code>EL</code> track in parallel with an <code>PS</code> track.
30	<code>lxbelin=0.0 m</code>	The sum of Li/Si quotients for an <code>EL</code> track in parallel with an <code>IN</code> track.
31	<code>lxbelins=0.0 m</code>	The sum of Li/Si quotients for an <code>EL</code> track in parallel with an <code>INS</code> track.

The Spectre option `scale`, default value is 1.0, scales the geometric parameters. The actual areas (parameters starting with letter `a`) are equal

$$axxx * scale ^ 2$$

The actual lengths (parameters starting with letter `l`) are equal

$$lxxx * scale$$

Model Definition

```
model modelName intcap parameter=value ...
```

Model Parameters

1	<code>cbps=0.0 F/m²</code>	Bottom capacitance, <code>PS</code> to node <code>n2</code> .
2	<code>cebpsm=0.0 F/m</code>	Edge to bottom capacitance (<code>PS</code>), 1.0um spacing.
3	<code>cebpsi=0.0 F/m</code>	Edge to bottom capacitance (<code>PS</code>), single track.

Spectre Circuit Simulator Reference

Component Statements Part II

4	<code>cetpsm=0.0 F/m</code>	Edge to top capacitance (PS), 1.0um spacing.
5	<code>cetpsi=0.0 F/m</code>	Edge to top capacitance (PS), single track.
6	<code>cbin=0.0 F/m²</code>	Bottom capacitance, IN to node n2.
7	<code>cebinm=0.0 F/m</code>	Edge to bottom capacitance (IN), 1.0um spacing.
8	<code>cebini=0.0 F/m</code>	Edge to bottom capacitance (IN), single track.
9	<code>cetinm=0.0 F/m</code>	Edge to top capacitance (IN), 1.0um spacing.
10	<code>cetini=0.0 F/m</code>	Edge to top capacitance (IN), single track.
11	<code>cbins=0.0 F/m²</code>	Bottom capacitance, INS to node n2.
12	<code>cebinsm=0.0 F/m</code>	Edge to bottom capacitance (INS), 1.0um spacing.
13	<code>cebinsi=0.0 F/m</code>	Edge to bottom capacitance (INS), single track.
14	<code>cetinsm=0.0 F/m</code>	Edge to top capacitance (INS), 1.0um spacing.
15	<code>cetinsi=0.0 F/m</code>	Edge to top capacitance (INS), single track.
16	<code>cbel=0.0 F/m²</code>	Bottom capacitance, EL to node n2.
17	<code>cebelm=0.0 F/m</code>	Edge to bottom capacitance (EL), 1.0um spacing.
18	<code>cebeli=0.0 F/m</code>	Edge to bottom capacitance (EL), single track.
19	<code>cetelm=0.0 F/m</code>	Edge to top capacitance (EL), 1.0um spacing.
20	<code>ceteli=0.0 F/m</code>	Edge to top capacitance (EL), single track.
21	<code>cecps=0.0 F/m</code>	Lateral capacitance (PS), 1.0um spacing.
22	<code>cecin=0.0 F/m</code>	Lateral capacitance (IN), 1.0um spacing.
23	<code>cecins=0.0 F/m</code>	Lateral capacitance (INS), 1.0um spacing.
24	<code>cecel=0.0 F/m</code>	Lateral capacitance (EL), 1.0um spacing.

Spectre Circuit Simulator Reference

Component Statements Part II

Output Parameters

1 `cap` (F) Total Capacitance.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

<code>ael</code>	I-2	<code>cebinsm</code>	M-12	<code>cetpsm</code>	M-4	<code>lftin</code>	I-15
<code>ain</code>	I-3	<code>cebpsi</code>	M-3	<code>lbel</code>	I-6	<code>lftins</code>	I-16
<code>ains</code>	I-4	<code>cebpsm</code>	M-2	<code>lbin</code>	I-7	<code>lftps</code>	I-17
<code>aps</code>	I-5	<code>cecel</code>	M-24	<code>lbins</code>	I-8	<code>ltel</code>	I-18
<code>cap</code>	O-1	<code>cecin</code>	M-22	<code>lbsps</code>	I-9	<code>ltin</code>	I-19
<code>cbel</code>	M-16	<code>cecins</code>	M-23	<code>ldsel</code>	I-22	<code>ltins</code>	I-20
<code>cbin</code>	M-6	<code>cecps</code>	M-21	<code>ldsins</code>	I-23	<code>ltps</code>	I-21
<code>cbins</code>	M-11	<code>ceteli</code>	M-20	<code>ldsins</code>	I-24	<code>lxbelin</code>	I-30
<code>cbps</code>	M-1	<code>cetelm</code>	M-19	<code>ldsps</code>	I-25	<code>lxbelins</code>	I-31
<code>cebeli</code>	M-18	<code>cetini</code>	M-10	<code>lfbel</code>	I-10	<code>lxbelps</code>	I-29
<code>cebelm</code>	M-17	<code>cetinm</code>	M-9	<code>lfbins</code>	I-11	<code>lxbinsps</code>	I-26
<code>cebini</code>	M-8	<code>cetinsi</code>	M-15	<code>lfbins</code>	I-12	<code>lxbinsin</code>	I-27
<code>cebinm</code>	M-7	<code>cetinsm</code>	M-14	<code>lfbps</code>	I-13	<code>lxbinsps</code>	I-28
<code>cebinsi</code>	M-13	<code>cetpsi</code>	M-5	<code>lftel</code>	I-14	<code>m</code>	I-1

Current Probe (`iprobe`)

Description

Current through the probe is computed and is defined to be positive if it flows from the input node, through the probe, to the output node. The current variable is given the name of the `iprobe` instance, so you cannot create an `iprobe` with the same name as a circuit node.

This device is not supported within altergroup.

Sample Instance Statement

```
ip (1 0) iprobe
```

Instance Definition

```
Name in out iprobe
```

Independent Current Source (isource)

Description

The value of the DC current as a function of the temperature is given by:

$$I(T) = I(tnom) * [1 + tc1 * (T - tnom) + tc2 * (T - tnom)^2].$$

Sample Instance Statement

```
i1 (in 0) isource dc=0 type=pulse delay=10n val0=0 vall=500u period=500n rise=1n  
fall=1n width=250n
```

Instance Definition

```
Name sink src isource parameter=value ...
```

Positive current exits the source node and enters the sink node.

Instance Parameters

1 dc=0 A DC value.

General waveform parameters

2 type=dc Waveform type.
Possible values are dc, pulse, pwl, sine, or exp.

3 fundname Name of the fundamental frequency. Must be specified if the source is active during a pdisto analysis or it is the active clock during an envlp analysis.

Spectre Circuit Simulator Reference

Component Statements Part II

4 `delay=0 s` Waveform delay time.

Pulse waveform parameters

5 `val0=0 A` Zero value used in pulse and exponential waveforms.

6 `val1=1 A` One value used in pulse and exponential waveforms.

7 `period= ∞ s` Period of waveform.

8 `rise (s)` Rise time for pulse waveform (time for transition from `val0` to `val1`).

9 `fall (s)` Fall time for pulse waveform (time for transition from `val1` to `val0`).

10 `width= ∞ s` Pulse width (duration of `val1`).

PWL waveform parameters

11 `file` Name of file containing waveform.

12 `wave=[...]` Vector of time/value pairs that defines waveform.

13 `offset=0 A` DC offset for the PWL waveform.

14 `scale=1` Scale factor for the PWL waveform.

15 `stretch=1` Scale factor for time given for the PWL waveform.

16 `allbrkpts` All the points in the PWL waveform are breakpoints if set to yes. Default is yes if the number of points is less than 20. Possible values are `no` or `yes`.

17 `pwlperiod (s)` Period of the periodic PWL waveform.

18 `twidth=pwlperiod/1000 s` Transition width used when making PWL waveforms periodic.

Sinusoidal waveform parameters

19 `sinedc=dc A` DC level for sinusoidal waveforms.

Spectre Circuit Simulator Reference

Component Statements Part II

20	<code>amp1=1 A</code>	Peak amplitude of sinusoidal waveform.
21	<code>freq=0 Hz</code>	Frequency of sinusoidal waveform.
22	<code>sinephase=0 °</code>	Phase of sinusoid when <code>t=delay</code> .
23	<code>amp12=1 A</code>	Peak amplitude of second sinusoidal waveform.
24	<code>freq2=0 Hz</code>	Frequency of second sinusoidal waveform.
25	<code>sinephase2=0 °</code>	Phase of second sinusoid when <code>t=delay</code> .
26	<code>fundname2</code>	Name of the fundamental frequency associated with <code>freq2</code> . Must be specified if <code>freq2</code> is used in a <code>pdisto</code> analysis.
27	<code>fmodindex=0</code>	FM index of modulation for sinusoidal waveform.
28	<code>fmodfreq=0 Hz</code>	FM modulation frequency for sinusoidal waveform.
29	<code>ammodindex=0</code>	AM index of modulation for sinusoidal waveform.
30	<code>ammodfreq=0 Hz</code>	AM modulation frequency for sinusoidal waveform.
31	<code>ammodphase=0 °</code>	AM phase of modulation for sinusoidal waveform.
32	<code>damp=0 1/s</code>	Damping factor for sinusoidal waveform.

Exponential waveform parameters

33	<code>td1=0 s</code>	Rise start time for exponential wave.
34	<code>tau1 (s)</code>	Rise time constant for exponential wave.
35	<code>td2 (s)</code>	Fall start time for exponential wave.
36	<code>tau2 (s)</code>	Fall time constant for exponential wave.

Noise Parameters

37	<code>noisefile</code>	Name of file containing excess spot noise data in the form of frequency-noise pairs.
----	------------------------	--

Spectre Circuit Simulator Reference

Component Statements Part II

38 `noisevec=[...]` A^2/Hz Excess spot noise as a function of frequency in the form of frequency-noise pairs.

Small signal parameters

- 39 `mag=0` A Small signal current.
- 40 `phase=0` ° Small signal phase.
- 41 `xfmag=1` A/A Transfer function analysis magnitude.
- 42 `pacmag=0` A Periodic AC analysis magnitude.
- 43 `pacphase=0` ° Periodic AC analysis phase.

Multiplication factor parameters

- 44 `m=1` Multiplicity factor.

Temperature effects parameters

- 45 `tc1=0` 1/C First order temperature coefficient.
- 46 `tc2=0` C^{-2} Second order temperature coefficient.
- 47 `tnom=27` C Parameter measurement temperature. Default set by options.

If you do not specify the DC value, it is assumed to be the `time=0` value of the waveform.

Sinusoidal waveform in the time interval $0 < t < \text{delay}$ has constant value equal to that at `t=delay`.

In DC analyses, the only active parameters are `dc`, `m`, and the temperature coefficient parameters. In AC analyses, the only active parameters are `m`, `mag` and `phase`. In transient analyses, all parameters are active except the small signal parameters and the noise parameters. The `type` parameter selects which type of waveform is generated. You may specify parameters for more than one waveform type, and use the `alter` statement to change the waveform type between analyses.

A vector of time-value pairs describes the piecewise linear waveform. As an alternative, you can read the waveform from a file. In this case, you give time-value pairs one pair per line with a space or tab between the time and the value.

Spectre Circuit Simulator Reference

Component Statements Part II

If you set `allbrkpts` to `yes`, you force the simulator to place time points at each point specified in a PWL waveform during a transient analysis. This can be very expensive for waveforms with many points. If you set `allbrkpts` to `no`, Spectre inspects the waveform, looking for abrupt changes, and forces time points only at those changes.

The PWL waveform is periodic if you specify `pwlperiod`. If the value of the waveform specified is not exactly the same at both its beginning and its end, then you must provide a nonzero value `twidth`. Before repeating, the waveform changes linearly in an interval of `twidth` from its value at `(period - twidth)` to its value at the beginning of the waveform. Thus `twidth` must always be less than `period`.

You can give the excess noise of the source as a file or specify it with a vector of frequency-noise pairs. For a file, give the frequency-noise pairs one pair per line with a space or tab between the frequency and noise values.

Operating-Point Parameters

1	<code>i</code> (A)	Current through the source.
2	<code>v</code> (V)	Voltage across the source.
3	<code>pwr</code> (W)	Power dissipation.

Parameter Index

In the following index, `I` refers to instance parameters, `M` refers to the model parameters section, `O` refers to the output parameters section, and `OP` refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of `M-35` means the 35th model parameter.

<code>allbrkpts</code>	I-16	<code>freq</code>	I-21	<code>phase</code>	I-40	<code>td1</code>	I-33
<code>ammodfreq</code>	I-30	<code>freq2</code>	I-24	<code>pwlperiod</code>	I-17	<code>td2</code>	I-35
<code>ammodindex</code>	I-29	<code>fundname</code>	I-3	<code>pwr</code>	OP-3	<code>tnom</code>	I-47
<code>ammodphase</code>	I-31	<code>fundname2</code>	I-26	<code>rise</code>	I-8	<code>twidth</code>	I-18
<code>amp1</code>	I-20	<code>i</code>	OP-1	<code>scale</code>	I-14	<code>type</code>	I-2
<code>amp12</code>	I-23	<code>m</code>	I-44	<code>sinedc</code>	I-19	<code>v</code>	OP-2
<code>damp</code>	I-32	<code>mag</code>	I-39	<code>sinephase</code>	I-22	<code>val0</code>	I-5

Spectre Circuit Simulator Reference Component Statements Part II

dc	I-1	noisefile	I-37	sinephase2	I-25	vall	I-6
delay	I-4	noisevec	I-38	stretch	I-15	wave	I-12
fall	I-9	offset	I-13	tau1	I-34	width	I-10
file	I-11	pacmag	I-42	tau2	I-36	xfmag	I-41
fmodfreq	I-28	pacphase	I-43	tc1	I-45		
fmodindex	I-27	period	I-7	tc2	I-46		

Junction Field Effect Transistor (jfet)

Description

The JFET model is derived from the FET model of Shichman and Hodges. JFETs require that you use a model statement.

This device is supported within altergroups.

Sample Instance Statement

```
jf1 (net1 net2 0) jmod area=1
```

Sample Model Statement

```
model jmod jfet beta=9e-5 lambda=0 type=n vt0=-18.7 rd=10 rs=10 cgs=1.3e-13 pb=0.65
```

Instance Definition

```
Name d g s [b] ModelName parameter=value ...
```

You do not have to specify the back gate terminal when you use the four-terminal model. If left unspecified, the substrate is connected to ground.

Instance Parameters

- 1 area=1 Junction area factor.
- 2 m=1 Multiplicity factor.

Spectre Circuit Simulator Reference

Component Statements Part II

3 `region=triode` Estimated operating region. Spectre outputs number (0-4) in a rawfile.
Possible values are `off`, `triode`, `sat`, `subth`, or `breakdown`.

Model Definition

`model modelName jfet parameter=value ...`

Model Parameters

Device type parameters

1 `type=n` Transistor type.
Possible values are `n` or `p`.

Drain current model parameters

2 `level=1` Drain current model level selector.

3 `vto=-2 V` Pinchoff voltage.

4 `beta=0.0001 A/V2` Transconductance parameter.

5 `lambda=0 1/V` Channel length modulation parameter.

6 `lambda1=0 1/V` Gate dependence of channel length modulation parameter.

7 `np=2` Power-law exponent.

8 `alpha=2` Triode-to-saturation transition parameter.

9 `io=0 A` Subthreshold current parameter.

10 `ns=1` Subthreshold swing parameter.

11 `ai=0 1/V` Impact ionization current coefficient.

12 `bi=0 V` Impact ionization current exponent.

Spectre Circuit Simulator Reference

Component Statements Part II

Four terminal threshold voltage parameters

- 13 `vtop=0.6 V` Back gate to channel junction potential.
- 14 `vtos=1.2 V` Threshold voltage slope.
- 15 `vtoe=0.33` Threshold voltage exponent.
- 16 `vtoc=-3.3 V` Threshold voltage constant.

Parasitic resistance parameters

- 17 `rd=0 Ω` Drain resistance (/area).
- 18 `rs=0 Ω` Source resistance (/area).
- 19 `rg=0 Ω` Gate resistance (/area).
- 20 `rb=0 Ω` Back gate resistance (/area).
- 21 `minr=0.1 Ω` Minimum source/drain/gate resistance.

Junction diode model parameters

- 22 `is=1e-14 A` Gate saturation current (*area).
- 23 `n=1` Emission coefficient for G-D and G-S junctions.
- 24 `imelt=`imaxA'` Explosion current (*area).
- 25 `dskip=yes` Use simple piece-wise linear model for diode currents below $0.1 \cdot i_{abstol}$.
Possible values are `no` or `yes`.

Junction capacitance model parameters

- 26 `tt=0 s` Transit time.
- 27 `cgs=0 F` Gate-source zero-bias junction capacitance (*area).
- 28 `cgd=0 F` Gate-drain zero-bias junction capacitance (*area).

Spectre Circuit Simulator Reference

Component Statements Part II

- 29 `mj=1/2` Junction grading coefficient.
- 30 `pb=1 V` Gate-junction potential.
- 31 `fc=0.5` Junction capacitor forward-bias threshold.

Four terminal junction parameters

- 32 `isb=1e-14 A` Back gate-saturation current (*area).
- 33 `nb=1` Emission coefficient for back gate-junctions.
- 34 `cgbs=0 F` Back gate-source zero-bias junction capacitance (*area).
- 35 `cgbd=0 F` Back gate-drain zero-bias junction capacitance (*area).
- 36 `mjb=1/2` Back gate-junction grading coefficient.
- 37 `pbb=1 V` Back gate-junction potential.

Temperature effect parameters

- 38 `tnom (C)` Parameters measurement temperature. Default set by `options`.
- 39 `trise=0 C` Temperature rise from ambient.
- 40 `xti=3` Temperature exponent for effect on `is`.
- 41 `tlev=0` DC temperature selector.
- 42 `tlevc=0` AC temperature selector.
- 43 `eg=1.12452 V` Energy band gap.
- 44 `gap1=7.02e-4 V/C` Band gap temperature coefficient.
- 45 `gap2=1108 C` Band gap temperature offset.
- 46 `tcv=0 1/C` Threshold voltage temperature coefficient.
- 47 `bto=0 C` Transconductance parameter temperature offset.

Spectre Circuit Simulator Reference

Component Statements Part II

48	<code>bte=0</code>	Transconductance parameter temperature exponent.
49	<code>lto=0 C</code>	Channel length modulation parameters temperature offset.
50	<code>lte=0</code>	Channel length modulation parameters temperature exponent.
51	<code>tc1=0 1/C</code>	Linear temperature coefficient for parasitic resistors.
52	<code>tc2=0 C⁻²</code>	Quadratic temperature coefficient for parasitic resistors.

Operating region warning control parameters

53	<code>alarm=none</code>	Forbidden operating region. Possible values are <code>none</code> , <code>off</code> , <code>triode</code> , <code>sat</code> , <code>subth</code> , or <code>rev</code> .
54	<code>imax=1 A</code>	Maximum allowable current (*area).
55	<code>bvj=∞ V</code>	Junction reverse breakdown voltage.

Noise parameters

56	<code>kf=0</code>	Flicker noise (1/f) coefficient.
57	<code>af=1</code>	Flicker noise (1/f) exponent.
58	<code>kfd=0</code>	Flicker noise (1/f) coefficient for gate diodes.
59	<code>afg=1</code>	Flicker noise (1/f) exponent for gate diodes.

Imax and Imelt

The `imax` parameter aids convergence and prevents numerical overflow. The junction characteristics of the device are accurately modeled for current up to `imax`. If `imax` is exceeded during iterations, the linear model is substituted until the current drops below `imax` or until convergence is achieved. If convergence is achieved with the current exceeding `imax`, the results are inaccurate, and Spectre prints a warning.

A separate model parameter, `imelt`, is used as a limit warning for the junction current. This parameter can be set to the maximum current rating of the device. When any component of the junction current exceeds `imelt`, note that base and collector currents are composed of

Spectre Circuit Simulator Reference

Component Statements Part II

many exponential terms, Spectre issues a warning and the results become inaccurate. The junction current is linearized above the value of `imelt` to prevent arithmetic exception, with the exponential term replaced by a linear equation at `imelt`.

The `bv` parameter is used to detect the junction breakdown only. The breakdown currents of the junctions are not modeled.

Operating-Point Parameters

1	<code>type=n</code>	Transistor type. Possible values are <code>n</code> or <code>p</code> .
2	<code>region=triode</code>	Estimated operating region. Spectre outputs number (0-4) in a rawfile. Possible values are <code>off</code> , <code>triode</code> , <code>sat</code> , <code>subth</code> , or <code>breakdown</code> .
3	<code>ids (A)</code>	Resistive drain current.
4	<code>vgs (V)</code>	Gate-source voltage.
5	<code>vds (V)</code>	Drain-source voltage.
6	<code>vth (V)</code>	Threshold at op point.
7	<code>vdsat (V)</code>	Drain saturation voltage.
8	<code>gm (S)</code>	Common-source transconductance.
9	<code>gds (S)</code>	Common-source output conductance.
10	<code>cgs (F)</code>	Gate-source capacitance.
11	<code>cgd (F)</code>	Gate-drain capacitance.
12	<code>ig (A)</code>	Resistive gate current.
13	<code>pwr (W)</code>	Power at op point.
14	<code>qd (V)</code>	Threshold at op point.
15	<code>qg (V)</code>	Threshold at op point.
16	<code>qs (V)</code>	Threshold at op point.

Spectre Circuit Simulator Reference Component Statements Part II

17 `qb` (V) Threshold at op point.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

<code>af</code>	M-57	<code>gap1</code>	M-44	<code>mj</code>	M-29	<code>tc2</code>	M-52
<code>afg</code>	M-59	<code>gap2</code>	M-45	<code>mjb</code>	M-36	<code>tcv</code>	M-46
<code>ai</code>	M-11	<code>gds</code>	OP-9	<code>n</code>	M-23	<code>tlev</code>	M-41
<code>alarm</code>	M-53	<code>gm</code>	OP-8	<code>nb</code>	M-33	<code>tlevc</code>	M-42
<code>alpha</code>	M-8	<code>ids</code>	OP-3	<code>np</code>	M-7	<code>tnom</code>	M-38
<code>area</code>	I-1	<code>ig</code>	OP-12	<code>ns</code>	M-10	<code>trise</code>	M-39
<code>beta</code>	M-4	<code>imax</code>	M-54	<code>pb</code>	M-30	<code>tt</code>	M-26
<code>bi</code>	M-12	<code>imelt</code>	M-24	<code>pbb</code>	M-37	<code>type</code>	M-1
<code>bte</code>	M-48	<code>io</code>	M-9	<code>pwr</code>	OP-13	<code>type</code>	OP-1
<code>bto</code>	M-47	<code>is</code>	M-22	<code>qb</code>	OP-17	<code>vds</code>	OP-5
<code>bvj</code>	M-55	<code>isb</code>	M-32	<code>qd</code>	OP-14	<code>vdsat</code>	OP-7
<code>cgbd</code>	M-35	<code>kf</code>	M-56	<code>qg</code>	OP-15	<code>vgs</code>	OP-4
<code>cgbs</code>	M-34	<code>kfd</code>	M-58	<code>qs</code>	OP-16	<code>vth</code>	OP-6
<code>cgd</code>	OP-11	<code>lambda</code>	M-5	<code>rb</code>	M-20	<code>vto</code>	M-3
<code>cgd</code>	M-28	<code>lambda1</code>	M-6	<code>rd</code>	M-17	<code>vtoc</code>	M-16
<code>cgs</code>	M-27	<code>level</code>	M-2	<code>region</code>	OP-2	<code>vtoe</code>	M-15
<code>cgs</code>	OP-10	<code>lte</code>	M-50	<code>region</code>	I-3	<code>vtop</code>	M-13
<code>dskip</code>	M-25	<code>lto</code>	M-49	<code>rg</code>	M-19	<code>vtos</code>	M-14
<code>eg</code>	M-43	<code>m</code>	I-2	<code>rs</code>	M-18	<code>xti</code>	M-40
<code>fc</code>	M-31	<code>minr</code>	M-21	<code>tc1</code>	M-51		

Junction Capacitor (juncap)

Description

The juncap model is intended to describe the behavior of the diodes that are formed by the source, drain or well-to-bulk junctions in MOS devices. It is described in the Philips MOST Modelbook (Dec.93) as JUNCAP model. Information on how to obtain this document can be found on Source Link by searching for Philips.

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In extension to the modelbook description a minimum conductance g_{min} is inserted between the juncap nodes, to aid convergence. The value of g_{min} is set by an options statement, default = 1e-12 S.

The i_{max} parameter is used to aid convergence and to prevent numerical overflow. The junction characteristics of the junction capacitor are accurately modeled for currents up to i_{max} . For currents above i_{max} , the junction is modeled as a linear resistor and a warning is printed.

This device is supported within altergroups.

This device is dynamically loaded from the shared object /vobs/spectre_dev/tools.sun4v/spectre/lib/cmi/3.0.doc/libphilips_sh.so

Sample Instance Statement

```
c2 (1 2) capmod ab=7e-12 lg=5e-6 region=rev
```

Sample Model Statement

```
model capmod juncap type=n cjbr=0.2 cjgr=0.2 cjsr=0.2 tref=25 jsgbr=2e-3 jsdbr=0.28e-3  
jsggr=1e-5 jsdgr=0.33e-6 vdsr=0.8 vdgr=0.8 vdbr=0.8
```

Instance Definition

```
Name n [b] ModelName parameter=value ...
```

Instance Parameters

```
1 ab=1.0 scale2 m2
```

Diffusion area. Scale set by option scale.

Spectre Circuit Simulator Reference

Component Statements Part II

- | | | |
|---|-----------------------------|--|
| 2 | <code>ls=1.0 scale m</code> | Length of the sidewall of the diffusion area ab which is not under the gate. Scale set by option scale. |
| 3 | <code>lg=1.0 scale m</code> | Length of the sidewall of the diffusion area ab which is under the gate. Scale set by option scale. |
| 4 | <code>m=1.0</code> | Multiplicity factor. |
| 5 | <code>region=rev</code> | Estimated DC operating region, used as a convergence aid. Possible values are <code> fwd </code> or <code> rev </code> . |

Model Definition

`model modelName juncap parameter=value ...`

Model Parameters

Structural parameters

- | | | |
|---|---------------------|---|
| 1 | <code>type=n</code> | Type of the juncap device. Possible values are <code> n </code> or <code> p </code> . |
| 2 | <code>vb (V)</code> | Not used for juncap model. |
| 3 | <code>bv (V)</code> | Alias of <code> vb </code> . |
| 4 | <code>level</code> | Not used for juncap model. |

Current parameters

- | | | |
|---|---|---|
| 5 | <code>jsgbr=1.0e-3 A/m²</code> | Bottom saturation-current density due to electron-hole generation at reference voltage. |
| 6 | <code>jsdbr=1.0e-3 A/m²</code> | Bottom saturation-current density due to diffusion from back contact. |
| 7 | <code>jsgsr=1.0e-3 A/m</code> | Sidewall saturation-current density due to electron-hole generation at reference voltage. |

Spectre Circuit Simulator Reference

Component Statements Part II

- 8 `jsdsr=1.0e-3 A/m` Sidewall saturation-current density due to diffusion from back contact.
- 9 `jsggr=1.0e-3 A/m` Gate edge saturation-current density due to electron-hole generation at reference voltage.
- 10 `jsdgr=1.0e-3 A/m` Gate edge saturation-current density due to diffusion from back contact.
- 11 `imax=1.0 A` Explosion current.

Temperature effects parameters

- 12 `dta=0.0 K` Temperature offset of the juncap element with respect to ambient temperature.
- 13 `trise=0.0 K` Alias of `dta`.
- 14 `tr (C)` Temperature at which the parameters have been determined. Default set by option `tnom`.
- 15 `tref (C)` Alias of `tr`. Default set by option `tnom`.
- 16 `tnom (C)` Alias of `tr`. Default set by option `tnom`.

Junction capacitance parameters

- 17 `cjbr=1.0e-12 F/m2` Bottom junction capacitance at reference voltage.
- 18 `cjsr=1.0e-12 F/m` Sidewall junction capacitance at reference voltage.
- 19 `cjgr=1.0e-12 F/m` Gate edge junction capacitance at reference voltage.

Emission coefficient parameters

- 20 `nb=1.0` Emission coefficient of the bottom forward current.
- 21 `ns=1.0` Emission coefficient of the sidewall forward current.
- 22 `ng=1.0` Emission coefficient of the gate-edge forward current.

Spectre Circuit Simulator Reference

Component Statements Part II

Voltage parameters

- | | | |
|----|-----------------|---|
| 23 | $v_r=0.0$ V | Voltage at which parameters have been determined. |
| 24 | $v_{dbr}=1.0$ V | Diffusion voltage of the bottom junction at reference temperature. |
| 25 | $v_{dsr}=1.0$ V | Diffusion voltage of the sidewall junction at reference temperature. |
| 26 | $v_{dgr}=1.0$ V | Diffusion voltage of the gate edge junction at reference temperature. |

Grading coefficient parameters

- | | | |
|----|-----------|---|
| 27 | $p_b=0.4$ | Bottom-junction grading coefficient. |
| 28 | $p_s=0.4$ | Sidewall-junction grading coefficient. |
| 29 | $p_g=0.4$ | Gate edge-junction grading coefficient. |

Output Parameters

- | | | |
|----|---------------|--|
| 1 | c_{jb} (F) | Capacitance of bottom area a_b . |
| 2 | c_{js} (F) | Capacitance of locos-edge l_s . |
| 3 | c_{jg} (F) | Capacitance of gate-edge l_g . |
| 4 | i_{sdb} (A) | Diffusion saturation-current of bottom area a_b . |
| 5 | i_{sds} (A) | Diffusion saturation-current of locos-edge l_s . |
| 6 | i_{sdg} (A) | Diffusion saturation-current of gate-edge l_g . |
| 7 | i_{sgb} (A) | Generation saturation-current of bottom area a_b . |
| 8 | i_{sgs} (A) | Generation saturation-current of locos-edge l_s . |
| 9 | i_{sgg} (A) | Generation saturation-current of gate-edge l_g . |
| 10 | v_{db} (V) | Diffusion voltage of bottom area a_b . |

Spectre Circuit Simulator Reference Component Statements Part II

- 11 `vds` (V) Diffusion voltage of locos-edge `ls`.
 12 `vdg` (V) Diffusion voltage of gate-edge `lg`.

Operating-Point Parameters

- 1 `v` (V) Diode bias voltage ($v = v_a - v_k$).
 2 `i` (A) Total resistive current from anode to cathode ($i = i_a = -i_k$).
 3 `gm` (S) Total differential conductance.
 4 `q` (Coul) Total junction charge ($q = q_a = -q_k$).
 5 `c` (F) Total capacitance.
 6 `pwr` (W) Power.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

<code>ab</code>	I-1	<code>isdb</code>	O-4	<code>lg</code>	I-3	<code>tr</code>	M-14
<code>bv</code>	M-3	<code>isdg</code>	O-6	<code>ls</code>	I-2	<code>tref</code>	M-15
<code>c</code>	OP-5	<code>isds</code>	O-5	<code>m</code>	I-4	<code>trise</code>	M-13
<code>cjb</code>	O-1	<code>isgb</code>	O-7	<code>nb</code>	M-20	<code>type</code>	M-1
<code>cjbr</code>	M-17	<code>isgg</code>	O-9	<code>ng</code>	M-22	<code>v</code>	OP-1
<code>cjg</code>	O-3	<code>isgs</code>	O-8	<code>ns</code>	M-21	<code>vb</code>	M-2
<code>cjgr</code>	M-19	<code>jsdbr</code>	M-6	<code>pb</code>	M-27	<code>vdb</code>	O-10
<code>cjs</code>	O-2	<code>jsdgr</code>	M-10	<code>pg</code>	M-29	<code>vdbr</code>	M-24
<code>cjsr</code>	M-18	<code>jsdsr</code>	M-8	<code>ps</code>	M-28	<code>vdg</code>	O-12
<code>dta</code>	M-12	<code>jsgbr</code>	M-5	<code>pwr</code>	OP-6	<code>vdgr</code>	M-26
<code>gm</code>	OP-3	<code>jsggr</code>	M-9	<code>q</code>	OP-4	<code>vds</code>	O-11

Spectre Circuit Simulator Reference Component Statements Part II

i	OP-2	jsgsr	M-7	region	I-5	vdsr	M-25
imax	M-11	level	M-4	tnom	M-16	vr	M-23

MISN Field Effect Transistor (misnan)

Description

The MISN model is formulated in terms of solutions for the boundary surface potentials of the channel and has the inherent property of continuous modeling. It is an inhouse MOSFET model of NORTEL. The MISN model requires a model statement.

This device is not supported within altergroup.

This device is dynamically loaded from the shared object /vobs/spectre_dev/tools.sun4v/spectre/lib/cmi/3.0.doc/libnortel_sh.so

Sample Instance Statement

```
mn1 (1 2 0 0) nch w=1.5u l=1u ad=2.6p as=2.6p pd=6.6p ps=6.6p
```

Sample Model Statement

```
model nch misnan type=n cox=4.4e-6 dop=2e17 phi=-0.43 xj=0.23 scrat=1.4 mu=400  
rws=250 is=0.98e-13 cjgo=2e-13 noimdl=1
```

Instance Definition

```
Name d g s b ModelName parameter=value ...
```

Instance Parameters

- | | | |
|---|-------------------------|---------------------------|
| 1 | w=1e-5 m | Channel width. |
| 2 | l=3e-6 m | Channel length. |
| 3 | as=3e-11 m ² | Area of source diffusion. |
| 4 | ad=3e-11 m ² | Area of drain diffusion. |

Spectre Circuit Simulator Reference

Component Statements Part II

- 5 `ps=2.6e-5 m` Perimeter of source diffusion.
- 6 `pd=2.6e-5 m` Perimeter of drain diffusion.
- 7 `m=1` Multiplicity factor (number of MOSFETs in parallel).
- 8 `region=triode` Estimated DC operating region, used as a convergence aid. Possible values are `off`, `triode`, `sat`, or `subthresh`.

Model Definition

`model modelName misnan parameter=value ...`

Model Parameters

Intrinsic MOS parameters

- 1 `type=n` Transistor gender. Possible values are `n` or `p`.
- 2 `cox=4.309e-7 F/cm2` Gate oxide cap per unit area.
- 3 `dop=1.665e17 cm-3` Substrate doping. Default = 2.58×10^{17} for pmos.
- 4 `phi=-0.55 V` Gate Fermi potential.
- 5 `qss=-5.078e-8 Coul/cm2` Effective gate oxide charge per unit area. Default = 1.05×10^{-8} for pmos.
- 6 `doplidd=3.2e17 cm-3` LDD region doping concentration. Default = 3.2×10^{19} for pmos.

Geometry parameters

- 7 `lvar=0 μm` Gate length correction.
- 8 `wvar=0 μm` Gate width correction.

Spectre Circuit Simulator Reference

Component Statements Part II

- 9 $d1s=0.0273 \mu\text{m}$ Sideway diffusion length of source region. Default = 0.037 for pmos.
- 10 $d1d=0.0273 \mu\text{m}$ Sideway diffusion length of drain region. Default = 0.037 for pmos.
- 11 $d1=0.07 \mu\text{m}$ Sideways diffusion length of S/D regions. Default = 0.04 for pmos.
- 12 $d_w=0.032 \mu\text{m}$ Electrical channel width correction. Default = 0.018 for pmos.

Threshold voltage parameters

- 13 $x_j=0.24 \mu\text{m}$ Source/drain-to-substrate junction depth. Default = 0.31 for pmos.
- 14 $scrat=1.5$ Short channel threshold voltage ratio. Default = 0.7 for pmos.
- 15 $scind=1.45$ Short channel threshold voltage index. Default = 1.42 for pmos.
- 16 $ncrat=0.17$ Narrow channel threshold voltage ratio. Default = 0.095 for pmos.
- 17 $athp=7.5$ Factor controlling peak magnitude effect. Default = 3.5 for pmos.
- 18 $athl=2e4 \text{ 1/cm}$ Factor controlling channel length dependence effect. Default = $4e4$ for pmos.
- 19 $athb=-1.7e-3$ Factor controlling substrate bias dependence effect. Default = $-6e-3$.

Mobility parameters

- 20 $\mu=577 \text{ cm}^2/\text{V s}$ Low-field carrier mobility. Default = 120 for pmos.
- 21 $\mu_{txp}=1.72$ Temperature coefficient for the carrier mobility. Default = 1.01 for pmos.
- 22 $k_g=1.4e-7 \text{ cm/V}$ Gate field factor. Default = $1.685e-7$.
- 23 $v_0=3.21e7 \text{ cm/s}$ Scattering limited velocity. Default = $2.45e7$.

Spectre Circuit Simulator Reference

Component Statements Part II

- 24 `v0txp=-6.3` Temp coefficient for scattering limited velocity. Default = -5 for pmos.
- 25 `find=1.25` Field mobility index factor. Default = 1.9 for pmos.
- 26 `gfc=9.1e-10` Gate voltage dependence of enhanced gate-field scattering. Default = 1.05e-10 for pmos.
- 27 `gfc=3e-5` Drain voltage dependence of enhanced gate-field scattering. Default = 2.3e-3 for pmos.
- 28 `gfmb=1.45e-3` Factor controlling substrate bias dependence of enhanced gate-field scattering. Default 3.3e-3 for pmos.
- 29 `csf=1.06e-11` Drain voltage dependence of coulombic scattering. Default = 1.35e-12 for pmos.
- 30 `csfb=1.61e-3` Body voltage dependence of coulombic scattering. Default = 8.5e-3 for pmos.

Saturation parameters

- 31 `dprat=15` Drain region/channel doping ration. Default = 2 for pmos.
- 32 `satpr=0.2` Saturation region shaping factor. Default = 1.0 for pmos.
- 33 `sbd=0.3535534` Primary parameter controlling the onset of saturation.
- 34 `sadr=5` Secondary parameter controlling the onset of saturation.

Capacitance parameters

- 35 `sccf=0.25` Inner fringing factor for the N+ S/D.

Extrinsic parameters

- 36 `rws=480 Ω μm` Source series resistance. Default = 1180 for pmos.
- 37 `rwd=480 Ω μm` Drain series resistance. Default = 1180 for pmos.
- 38 `rsd=-1 Ω μm` Drain/source series resistance. Negative value for asymmetrical devices.

Spectre Circuit Simulator Reference

Component Statements Part II

- 39 $rgsh=0 \ \Omega \ \mu m$ Gate series resistance.
- 40 $wtgf=0.28 \ \mu m$ Width of transition from gate to field oxide under poly.
- 41 $cpts=5.7e-9 \ F/cm^2$
Poly-to-substrate capacitance per unit area.
- 42 $cgfrs=1e-12 \ F/cm$ Gate-source overlap fringing field capacitance.
- 43 $cgfrd=1e-12 \ F/cm$ Gate-drain overlap fringing field capacitance.
- 44 $cgfr=1.36e-12 \ F/cm$
Gate overlap fringing field capacitance.
- Junction parameters**
- 45 $is=1.02e-12 \ A/cm^2$
Sat current per unit area of S/D region-injection component.
Default = $9.21e-13$ for pmos.
- 46 $isg=1e-20 \ A/cm$ Sat current per unit length of gate oxide periphery-injection component. Default = $1.17e-20$.
- 47 $isf=1e-20 \ A/cm$ Sat current per unit length of field oxide periphery-injection component. Default = $1.17e-20$.
- 48 $ig=1.31e-10 \ A/cm^2$
Sat current per unit area of S/D region-generation component.
Default = $8.27e-10$.
- 49 $igg=6.99e-14 \ A/cm$ Sat current per unit length of gate oxide periphery-generation/recombination component. Default = $6.47e-14$.
- 50 $igf=6.99e-14 \ A/cm$ Sat current per unit length of field oxide periphery-generation/recombination component. Default = $6.47e-14$.
- 51 $cjo=9.39e-8 \ F/cm^2$
Zero bias junction capacitance per unit area. Default = $1.273e-7$ for pmos.
- 52 $ena=0.387$ Junction capacitance coefficient for the area component. Default = 0.472 for pmos.

Spectre Circuit Simulator Reference

Component Statements Part II

- 53 `cjgo=2.085e-12 F/cm` Zero bias junction cap per unit length of gate oxide periphery.
Default = 1.864e-12 for pmos.
- 54 `eng=0.322` Junction cap coefficient for gate oxide periphery component.
Default = 0.334 for pmos.
- 55 `cjfo=3.037e-12 F/cm` Zero bias junction cap per unit length of field oxide periphery.
Default = 3.077e-12 for pmos.
- 56 `enf=0.322` Junction cap coefficient for field oxide periphery component.
Default = 0.334 for pmos.

Noise parameters

- 57 `noimdl=1` Noise model selector.
- 58 `nt=1.6e10 cm-2` Surface trap density. Default = 4e9 for pmos.
- 59 `nttx=-4` Surface trap density temperature coefficient.
- 60 `fidx=0.85` Flicker noise frequency coefficient.
- 61 `beta=1` Thermal noise proportional constant.
- 62 `sgma=3e-16` Capture cross section. Default = 3e-15 for pmos.
- 63 `xtau=1e-8` 1/E depth.
- 64 `wbar=1` Barrier height for tunneling. Default = 4 for pmos.
- 65 `dept=3e-7` Depth of trap distribution.

Operating-Point Parameters

- 1 `vgs (V)` Gate-source voltage.
- 2 `vds (V)` Drain-source voltage.
- 3 `vbs (V)` Bulk-source voltage.

Spectre Circuit Simulator Reference

Component Statements Part II

4	<code>id</code> (A)	Drain current.
5	<code>vth</code> (V)	Threshold voltage.
6	<code>vdsat</code> (V)	Drain-source saturation voltage.
7	<code>gm</code> (S)	Common-source transconductance.
8	<code>gd</code> (S)	Common-source output conductance.
9	<code>gs</code> (S)	Body-transconductance.
10	<code>gmb</code> (S)	Body transconductance.
11	<code>gjs</code> (S)	Drain-bulk junction conductance.
12	<code>ibs</code> (A)	Drain-bulk junction current.
13	<code>gjd</code> (S)	Source-bulk junction conductance.
14	<code>ibd</code> (A)	Source-bulk junction current.
15	<code>qgg</code> (Coul)	Gate charge.
16	<code>qss</code> (Coul)	Source charge.
17	<code>qdd</code> (Coul)	Drain charge.
18	<code>qbb</code> (Coul)	Bulk charge.
19	<code>cgg</code> (F)	C _{gg} .
20	<code>cgs</code> (F)	C _{gs} .
21	<code>cgd</code> (F)	C _{gd} .
22	<code>cgb</code> (F)	C _{gb} .
23	<code>csg</code> (F)	C _{sg} .
24	<code>css</code> (F)	C _{ss} .
25	<code>csd</code> (F)	C _{sd} .

Spectre Circuit Simulator Reference

Component Statements Part II

26	csb (F)	Csb.
27	cdg (F)	Cdg.
28	cds (F)	Cds.
29	cdd (F)	Cdd.
30	cdb (F)	Cdb.
31	cbg (F)	Cbg.
32	cbs (F)	Cbs.
33	cbd (F)	Cbd.
34	cbb (F)	Cbb.

Parameter Index

In the following index, I refers to instance parameters, M refers to the model parameters section, O refers to the output parameters section, and OP refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of M-35 means the 35th model parameter.

ad	I-4	csd	OP-25	ibd	OP-14	region	I-8
as	I-3	csf	M-29	ibs	OP-12	rgsh	M-39
athb	M-19	csfb	M-30	id	OP-4	rsd	M-38
athl	M-18	csg	OP-23	ig	M-48	rwd	M-37
athp	M-17	css	OP-24	igf	M-50	rws	M-36
beta	M-61	dept	M-65	igg	M-49	sadr	M-34
cbb	OP-34	dl	M-11	is	M-45	satpr	M-32
cbd	OP-33	dld	M-10	isf	M-47	sbdr	M-33
cbg	OP-31	dls	M-9	isg	M-46	sccf	M-35
cbs	OP-32	dop	M-3	kg	M-22	scind	M-15
cdb	OP-30	dopldd	M-6	l	I-2	scrat	M-14

Spectre Circuit Simulator Reference Component Statements Part II

cdd	OP-29	dprat	M-31	lvar	M-7	sgma	M-62
cdg	OP-27	dw	M-12	m	I-7	type	M-1
cds	OP-28	ena	M-52	mu	M-20	v0	M-23
cgb	OP-22	enf	M-56	mutxp	M-21	v0txp	M-24
cgd	OP-21	eng	M-54	ncrat	M-16	vbs	OP-3
cgfr	M-44	fidx	M-60	noimdl	M-57	vds	OP-2
cgfrd	M-43	find	M-25	nt	M-58	vdsat	OP-6
cgfrs	M-42	gd	OP-8	nttx	M-59	vgs	OP-1
cgg	OP-19	gfc	M-26	pd	I-6	vth	OP-5
cgs	OP-20	gfcm	M-27	phi	M-4	w	I-1
cjfo	M-55	gfmb	M-28	ps	I-5	wbar	M-64
cjgo	M-53	gjd	OP-13	qbb	OP-18	wtgf	M-40
cjo	M-51	gjs	OP-11	qdd	OP-17	wvar	M-8
cox	M-2	gm	OP-7	qgg	OP-15	xj	M-13
cpts	M-41	gmb	OP-10	qss	M-5	xtau	M-63
csb	OP-26	gs	OP-9	qss	OP-16		

MOS Level-0 Transistor (mos0)

Description

The MOS0 model is a simplified MOS level-1 model. The MOS0 DC drain current model is different from the Shichman and Hodges model because body effects are not modeled. The intrinsic MOS gate capacitances are replaced by the following linear overlap capacitances:

Gate to source/drain ($capmod = overlap$)

Gate to bulk ($capmod = bulk$)

Gate, source, and drain to ground ($capmod = gnd$)

MOS0 is usually used as a MOS switch. This model recognizes all the MOS and BSIM instance parameters but only uses l and w , ignoring all other parameters. MOS0 transistors require that you use a model statement.

This device is not supported within altergroup.

Spectre Circuit Simulator Reference

Component Statements Part II

Sample Instance Statement

```
mpl (0 1 2 2) pchmod0 l=2u w=30u ad=120p as=75p pd=36u ps=6u
```

Sample Model Statement

```
model pchmod0 mos0 type=p vto=-0.683 tox=0.21e-7 ld=0.45e-6 tnom=27
```

Instance Definition

```
Name d g s b ModelName parameter=value ...
```

Instance Parameters

- | | | |
|---|-------|--|
| 1 | w (m) | Channel width. |
| 2 | l (m) | Channel length. |
| 3 | m=1 | Multiplicity factor (number of MOSFETs in parallel). |

Model Definition

```
model modelName mos0 parameter=value ...
```

Model Parameters

Device type parameters

- | | | |
|---|--------|---|
| 1 | type=n | Transistor type.
Possible values are n or p. |
|---|--------|---|

Drain current model parameters

- | | | |
|---|-------------------------------|--------------------------------------|
| 2 | vto=0 V | Threshold voltage at zero body bias. |
| 3 | kp=2.0718e-5 A/V ² | Transconductance parameter. |
| 4 | lambda=0.02 1/V | Channel length modulation parameter. |
| 5 | tox=1e-7 m | Gate oxide thickness. |

Spectre Circuit Simulator Reference

Component Statements Part II

6 `ld=0 m` Lateral diffusion.

7 `wd=0 m` Field-oxide encroachment.

Charge model selection parameters

8 `capmod=gnd` Intrinsic charge model.
Possible values are `none`, `overlap`, `bulk`, or `gnd`.

Temperature parameters

9 `tnom (C)` Parameters measurement temperature. Default set by `options`.

10 `trise=0 C` Temperature rise from ambient.

Default device parameters

11 `w=3e-6 m` Default channel width.

12 `l=3e-6 m` Default channel length.

Operating-Point Parameters

1 `type=n` Transistor type.
Possible values are `n` or `p`.

2 `id (A)` Resistive drain current.

3 `vgs (V)` Gate-source voltage.

4 `vds (V)` Drain-source voltage.

5 `vbs (V)` Bulk-source voltage.

6 `vth (V)` Threshold voltage.

7 `vdsat (V)` Drain-source saturation voltage.

8 `gm (S)` Common-source transconductance.

9 `gds (S)` Common-source output conductance.

Spectre Circuit Simulator Reference Component Statements Part II

10	<code>cgs</code> (F)	Gate-source capacitance.
11	<code>cgd</code> (F)	Gate-drain capacitance.
12	<code>cgate</code> (F)	Gate-Ground capacitance.
13	<code>ron</code> (Ω)	On-resistance.
14	<code>pwr</code> (W)	Power at op point.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

<code>capmod</code>	M-8	<code>l</code>	I-2	<code>tox</code>	M-5	<code>vth</code>	OP-6
<code>cgate</code>	OP-12	<code>l</code>	M-12	<code>trise</code>	M-10	<code>vto</code>	M-2
<code>cgd</code>	OP-11	<code>lambda</code>	M-4	<code>type</code>	OP-1	<code>w</code>	M-11
<code>cgs</code>	OP-10	<code>ld</code>	M-6	<code>type</code>	M-1	<code>w</code>	I-1
<code>gds</code>	OP-9	<code>m</code>	I-3	<code>vbs</code>	OP-5	<code>wd</code>	M-7
<code>gm</code>	OP-8	<code>pwr</code>	OP-14	<code>vds</code>	OP-4		
<code>id</code>	OP-2	<code>ron</code>	OP-13	<code>vdsat</code>	OP-7		
<code>kp</code>	M-3	<code>tnom</code>	M-9	<code>vgs</code>	OP-3		

MOS Level-1 Transistor (`mos1`)

Description

The `MOS1` model is derived from the FET model of Shichman and Hodges. The velocity saturation and the mobility variation effects can also be incorporated into `MOS1`. Three charge models are available. `MOS1` transistors require that you use a model statement.

This device is supported within altergroups.

Spectre Circuit Simulator Reference

Component Statements Part II

Sample Instance Statement

```
nchl (1 2 0 0) nchmod1 l=2u w=15u ad=60p as=37.5p pd=23u ps=6u
```

Sample Model Statement

```
model nchmod1 mos1 vto=0.78 gamma=0.56 kp=0.8675e-4 tox=0.21e-7 nsub=0.21e17  
ld=0.55e-6 capmod=yang vmax=4e5 theta=0.19 cbs=11e-15 cbd=10e-15 lambda=0.1
```

Instance Definition

```
Name d g s b ModelName parameter=value ...
```

Instance Parameters

1	w (m)	Channel width.
2	l (m)	Channel length.
3	as (m ²)	Area of source diffusion.
4	ad (m ²)	Area of drain diffusion.
5	ps (m)	Perimeter of source diffusion.
6	pd (m)	Perimeter of drain diffusion.
7	nrd (m/m)	Number of squares of drain diffusion.
8	nrs (m/m)	Number of squares of source diffusion.
9	ld (m)	Length of drain diffusion region.
10	ls (m)	Length of source diffusion region.
11	m=1	Multiplicity factor (number of MOSFETs in parallel).
12	region=triode	Estimated operating region. Spectre outputs number (0-4) in a rawfile. Possible values are off, triode, sat, subth, or breakdown.
13	trise	Temperature rise from ambient.

Spectre Circuit Simulator Reference

Component Statements Part II

14 `degradation=no` Hot-electron degradation flag.
Possible values are `no` or `yes`.

Model Definition

`model modelName mos1 parameter=value ...`

Model Parameters

Device type parameters

1 `type=n` Transistor type.
Possible values are `n` or `p`.

Drain current model parameters

2 `vto=0 V` Threshold voltage at zero body bias.

3 `kp=2.0718e-5 A/V2` Transconductance parameter.

4 `lambda=0 1/V` Channel length modulation parameter.

5 `phi=0.7 V` Surface potential at strong inversion.

6 `gamma=0 \sqrt{V}` Body-effect parameter.

7 `uo=600 cm2/V s` Carrier surface mobility.

8 `vmax= ∞ m/s` Carrier saturation velocity.

9 `theta=0 1/V` Mobility modulation coefficient.

Process parameters

10 `nsub=1.13e16 cm-3` Channel doping concentration.

11 `nss=0 cm-2` Surface state density.

12 `nfs=0 cm-2` Fast surface state density.

Spectre Circuit Simulator Reference

Component Statements Part II

13	$t_{pg}=+1$	Type of gate (+1 = opposite of substate, -1 = same as substate, 0 = aluminum).
14	$ld=0$ m	Lateral diffusion.
15	$wd=0$ m	Field-oxide encroachment.
16	$xw=0$ m	Width variation due to masking and etching.
17	$xl=0$ m	Length variation due to masking and etching.
18	$t_{ox}=1e-7$ m	Gate oxide thickness.

Impact ionization parameters

19	$a_{i0}=0$ 1/V	Impact ionization current coefficient.
20	$l_{ai0}=0$ $\mu\text{m}/\text{V}$	Length sensitivity of a_{i0} .
21	$w_{ai0}=0$ $\mu\text{m}/\text{V}$	Width sensitivity of a_{i0} .
22	$b_{i0}=0$ V	Impact ionization current exponent.
23	$l_{bi0}=0$ μm V	Length sensitivity of b_{i0} .
24	$w_{bi0}=0$ μm V	Width sensitivity of b_{i0} .

Overlap capacitance parameters

25	$c_{gso}=0$ F/m	Gate-source overlap capacitance.
26	$c_{gdo}=0$ F/m	Gate-drain overlap capacitance.
27	$c_{gbo}=0$ F/m	Gate-bulk overlap capacitance.
28	$meto=0$ m	Metal overlap in fringing field.

Charge model selection parameters

29	$capmod=bsim$	Intrinsic charge model. Possible values are none, meyer, yang, or bsim.
----	---------------	--

Spectre Circuit Simulator Reference

Component Statements Part II

- 30 `xpart=1` Drain/source channel charge partition in saturation for BSIM charge model, use 0.0 for 40/60, 0.5 for 50/50, or 1.0 for 0/100.
- 31 `xqc=0` Drain/source channel charge partition in saturation for charge models, e.g. use 0.4 for 40/60, 0.5 for 50/50, 0 for 0/100.

Parasitic resistance parameters

- 32 `rs=0` Ω Source resistance.
- 33 `rd=0` Ω Drain resistance.
- 34 `rss=0` Ω m Scalable source resistance.
- 35 `rdd=0` Ω m Scalable drain resistance.
- 36 `rsh=0` Ω/sqr Source/drain diffusion sheet resistance.
- 37 `rsc=0` Ω Source contact resistance.
- 38 `rdc=0` Ω Drain contact resistance.
- 39 `minr=0.1` Ω Minimum source/drain resistance.
- 40 `ldif=0` m Lateral diffusion beyond the gate.
- 41 `hdif=0` m Length of heavily doped diffusion.
- 42 `lgcs=0` m Gate-to-contact length of source side.
- 43 `lgcd=0` m Gate-to-contact length of drain side.
- 44 `sc= ∞` m Spacing between contacts.

Junction diode model parameters

- 45 `js` (A/m^2) Bulk junction reverse saturation current density.
- 46 `is=1e-14` A Bulk junction reverse saturation current.
- 47 `n=1` Junction emission coefficient.

Spectre Circuit Simulator Reference

Component Statements Part II

- 48 `dskip=yes` Use simple piece-wise linear model for diode currents below $0.1 \cdot i_{abstol}$. Possible values are `no` or `yes`.
- 49 `imelt=`imaxA`` Explosion current, diode is linearized beyond this current to aid convergence.
- 50 `jmelt=`jmaxA/m`2` Explosion current density, diode is linearized beyond this current to aid convergence.

Junction capacitance model parameters

- 51 `cbs=0 F` Bulk-source zero-bias junction capacitance.
- 52 `cbd=0 F` Bulk-drain zero-bias junction capacitance.
- 53 `cj=0 F/m2` Zero-bias junction bottom capacitance density.
- 54 `mj=1/2` Bulk junction bottom grading coefficient.
- 55 `pb=0.8 V` Bulk junction built-in potential.
- 56 `fc=0.5` Forward-bias depletion capacitance threshold.
- 57 `cjsw=0 F/m` Zero-bias junction sidewall capacitance density.
- 58 `mjsw=1/3` Bulk junction sidewall grading coefficient.
- 59 `pbsw=0.8 V` Side-wall junction built-in potential.
- 60 `fcs=0.5` Side-wall forward-bias depletion capacitance threshold.

Operating region warning control parameters

- 61 `alarm=none` Forbidden operating region. Possible values are `none`, `off`, `triode`, `sat`, `subth`, or `rev`.
- 62 `imax=1 A` Maximum current, currents above this limit generate a warning.
- 63 `jmax=1e8 A/m2` Maximum current density, currents above this limit generate a warning.

Spectre Circuit Simulator Reference

Component Statements Part II

64 `bvj= ∞ V` Junction reverse breakdown voltage.

65 `vbox=1e9 tox V` Oxide breakdown voltage.

Temperature effects parameters

66 `tnom (C)` Parameters measurement temperature. Default set by `options`.

67 `trise=0 C` Temperature rise from ambient.

68 `uto=0 C` Mobility temperature offset.

69 `ute=-1.5` Mobility temperature exponent.

70 `tlev=0` DC temperature selector.

71 `tlevc=0` AC temperature selector.

72 `eg=1.12452 V` Energy band gap.

73 `gap1=7.02e-4 V/C` Band gap temperature coefficient.

74 `gap2=1108 C` Band gap temperature offset.

75 `flex=0` Temperature exponent for `ucrit`.

76 `lamex=0 1/C` Temperature parameter for `lambda` and `kappa`.

77 `trs=0 1/C` Temperature parameter for source resistance.

78 `trd=0 1/C` Temperature parameter for drain resistance.

79 `xti=3` Saturation current temperature exponent.

80 `ptc=0 V/C` Surface potential temperature coefficient.

81 `tcv=0 V/C` Threshold voltage temperature coefficient.

82 `pta=0 V/C` Junction potential temperature coefficient.

83 `ptp=0 V/C` Sidewall junction potential temperature coefficient.

84 `cta=0 1/C` Junction capacitance temperature coefficient.

Spectre Circuit Simulator Reference

Component Statements Part II

85 `ctp=0 1/C` Sidewall junction capacitance temperature coefficient.

Default instance parameters

86 `w=3e-6 m` Default channel width.

87 `l=3e-6 m` Default channel length.

88 `as=0 m2` Default area of source diffusion.

89 `ad=0 m2` Default area of drain diffusion.

90 `ps=0 m` Default perimeter of source diffusion.

91 `pd=0 m` Default perimeter of drain diffusion.

92 `nrd=0 m/m` Default number of squares of drain diffusion.

93 `nrs=0 m/m` Default number of squares of source diffusion.

94 `ldd=0 m` Default length of drain diffusion region.

95 `lds=0 m` Default length of source diffusion region.

Noise model parameters

96 `noisemod=1` Noise model selector.

97 `kf=0` Flicker (1/f) noise coefficient.

98 `af=1` Flicker (1/f) noise exponent.

99 `ef=1` Flicker (1/f) noise frequency exponent.

100 `wnoi=1e-5 m` Channel width at which noise parameters were extracted.

Auto Model Selector parameters

101 `wmax=1.0 m` Maximum channel width for which the model is valid.

102 `wmin=0.0 m` Minimum channel width for which the model is valid.

Spectre Circuit Simulator Reference

Component Statements Part II

103 `lmax=1.0 m` Maximum channel length for which the model is valid.

104 `lmin=0.0 m` Minimum channel length for which the model is valid.

Degradation parameters

105 `degramod=spectre` Degradation model selector.
Possible values are `spectre` or `bert`.

106 `degradation=no` Hot-electron degradation flag.
Possible values are `no` or `yes`.

107 `dvthc=1 V` Degradation coefficient for threshold voltage.

108 `dvthe=1` Degradation exponent for threshold voltage.

109 `duoc=1 S` Degradation coefficient for transconductance.

110 `duoe=1` Degradation exponent for transconductance.

111 `crivth=0.1 V` Maximum allowable threshold voltage shift.

112 `criuo=10%` Maximum allowable normalized mobility change.

113 `crigm=10%` Maximum allowable normalized transconductance change.

114 `criids=10%` Maximum allowable normalized drain current change.

115 `wnom=5e-6 m` Nominal device width in degradation calculation.

116 `lnom=1e-6 m` Nominal device length in degradation calculation.

117 `vbsn=0 V` Substrate voltage in degradation calculation.

118 `vdsni=0.1 V` Drain voltage in I_{ds} degradation calculation.

119 `vgsni=5 V` Gate voltage in I_{ds} degradation calculation.

120 `vdsng=0.1 V` Drain voltage in G_m degradation calculation.

121 `vgsng=5 V` Gate voltage in G_m degradation calculation.

Spectre Circuit Simulator Reference

Component Statements Part II

Spectre stress parameters

- 122 $esat=1.1e7$ V/m Critical field in Vdsat calculation.
- 123 $esatg=2.5e6$ 1/m Gate voltage dependence of esat.
- 124 $vpg=-0.25$ Gate voltage modifier.
- 125 $vpb=-0.13$ Gate voltage modifier.
- 126 $subc1=2.24e-5$ Substrate current coefficient.
- 127 $subc2=-0.1e-5$ 1/V Substrate current coefficient.
- 128 $sube=6.4$ Substrate current exponent.
- 129 $strc=1$ Stress coefficient.
- 130 $stre=1$ Stress exponent.

BERT stress parameters

- 131 $h0=1$ Aging coefficient.
- 132 $hgd=0$ 1/V Bias dependence of h0.
- 133 $m0=1$ Aging exponent.
- 134 $mgd=0$ 1/V Bias dependence of m0.
- 135 $ecrit0=1.1e5$ V/cm Critical electric field.
- 136 $lecrit0=0$ μm V/cm Length dependence of ecrit0.
- 137 $wecrit0=0$ μm V/cm Width dependence of ecrit0.
- 138 $ecritg=0$ 1/cm Gate voltage dependence of ecrit0.
- 139 $lecritg=0$ $\mu\text{m}/\text{cm}$ Length dependence of ecritg.
- 140 $wecritg=0$ $\mu\text{m}/\text{cm}$ Width dependence of ecritg.
- 141 $ecritb=0$ 1/cm Substrate voltage dependence of ecrit0.

Spectre Circuit Simulator Reference

Component Statements Part II

142	$l_{critb}=0$	$\mu\text{m}/\text{cm}$	Length dependence of l_{critb} .
143	$w_{critb}=0$	$\mu\text{m}/\text{cm}$	Width dependence of l_{critb} .
144	$lc0=1$		Substrate current coefficient.
145	$llc0=0$	μm	Length dependence of $lc0$.
146	$wlc0=0$	μm	Width dependence of $lc0$.
147	$lc1=1$		Substrate current coefficient.
148	$llc1=0$	μm	Length dependence of $lc1$.
149	$wlc1=0$	μm	Width dependence of $lc1$.
150	$lc2=1$		Substrate current coefficient.
151	$llc2=0$	μm	Length dependence of $lc2$.
152	$wlc2=0$	μm	Width dependence of $lc2$.
153	$lc3=1$		Substrate current coefficient.
154	$llc3=0$	μm	Length dependence of $lc3$.
155	$wlc3=0$	μm	Width dependence of $lc3$.
156	$lc4=1$		Substrate current coefficient.
157	$llc4=0$	μm	Length dependence of $lc4$.
158	$wlc4=0$	μm	Width dependence of $lc4$.
159	$lc5=1$		Substrate current coefficient.
160	$llc5=0$	μm	Length dependence of $lc5$.
161	$wlc5=0$	μm	Width dependence of $lc5$.
162	$lc6=1$		Substrate current coefficient.
163	$llc6=0$	μm	Length dependence of $lc6$.

Spectre Circuit Simulator Reference

Component Statements Part II

164	$wlc6=0 \mu m$	Width dependence of $lc6$.
165	$lc7=1$	Substrate current coefficient.
166	$llc7=0 \mu m$	Length dependence of $lc7$.
167	$wlc7=0 \mu m$	Width dependence of $lc7$.

Imax and Imelt

The $imax$ parameter aids convergence and prevents numerical overflow. The junction characteristics of the device are accurately modeled for current up to $imax$. If $imax$ is exceeded during iterations, the linear model is substituted until the current drops below $imax$ or until convergence is achieved. If convergence is achieved with the current exceeding $imax$, the results are inaccurate, and Spectre prints a warning.

A separate model parameter, $imelt$, is used as a limit warning for the junction current. This parameter can be set to the maximum current rating of the device. When any component of the junction current exceeds $imelt$, note that base and collector currents are composed of many exponential terms, Spectre issues a warning and the results become inaccurate. The junction current is linearized above the value of $imelt$ to prevent arithmetic exception, with the exponential term replaced by a linear equation at $imelt$.

Both of these parameters have current density counterparts, $jmax$ and $jimelt$, that you can specify if you want the absolute current values to depend on the device area.

Auto Model Selection

Many models need to be characterized for different geometries in order to obtain accurate results for model development. The model selector program automatically searches for a model with the length and width range specified in the instance statement and uses this model in the simulations.

For the auto model selector program to find a specific model, the models to be searched should be grouped together within braces. Such a group is called a model group. An opening brace is required at the end of the line defining each model group. Every model in the group is given a name followed by a colon and the list of parameters. Also, the four geometric parameters $lmax$, $lmin$, $wmax$, and $wmin$ should be given. The selection criteria to choose a model is as follows:

$$lmin \leq inst_length < lmax \quad \text{and} \quad wmin \leq inst_width < wmax$$

Example:

Spectre Circuit Simulator Reference

Component Statements Part II

```
model ModelName ModelType {  
    1: <model parameters> lmin=2 lmax=4 wmin=1 wmax=2  
    2: <model parameters> lmin=1 lmax=2 wmin=2 wmax=4  
    3: <model parameters> lmin=2 lmax=4 wmin=4 wmax=6  
}
```

Then for a given instance

```
M1 1 2 3 4 ModelName w=3 l=1.5
```

the program would search all the models in the model group with the name `ModelName` and then pick the first model whose geometric range satisfies the selection criteria. In the preceding example, the auto model selector program would choose `ModelName.2`.

You must specify both length (`l`) and width (`w`) on the device instance line to enable automatic model selection.

Output Parameters

1	<code>w_{eff}</code> (m)	Effective channel width.
2	<code>l_{eff}</code> (m)	Effective channel length.
3	<code>r_{seff}</code> (Ω)	Effective source resistance.
4	<code>r_{deff}</code> (Ω)	Effective drain resistance.
5	<code>a_{seff}</code> (m ²)	Effective area of source diffusion.
6	<code>a_{deff}</code> (m ²)	Effective area of drain diffusion.
7	<code>p_{seff}</code> (m)	Effective perimeter of source diffusion.
8	<code>p_{deff}</code> (m)	Effective perimeter of source diffusion.
9	<code>i_{sseff}</code> (A)	Effective source-bulk junction reverse saturation current.
10	<code>i_{sdeff}</code> (A)	Effective drain-bulk junction reverse saturation current.
11	<code>cb_{seff}</code> (F)	Effective zero-bias source-bulk junction capacitance.

Spectre Circuit Simulator Reference

Component Statements Part II

12 `cbdeff` (F) Effective zero-bias drain-bulk junction capacitance.

Operating-Point Parameters

1 `type=n` Transistor type.
Possible values are `n` or `p`.

2 `region=triode` Estimated operating region. Spectre outputs number (0-4) in a
rawfile.
Possible values are `off`, `triode`, `sat`, `subth`, or `breakdown`.

3 `degradation=no` Hot-electron degradation flag.
Possible values are `no` or `yes`.

4 `reversed` Reverse mode indicator.
Possible values are `no` or `yes`.

5 `ids` (A) Resistive drain-to-source current.

6 `vgs` (V) Gate-source voltage.

7 `vds` (V) Drain-source voltage.

8 `vbs` (V) Bulk-source voltage.

9 `vth` (V) Threshold voltage.

10 `vdsat` (V) Drain-source saturation voltage.

11 `gm` (S) Common-source transconductance.

12 `gds` (S) Common-source output conductance.

13 `gmbs` (S) Body-transconductance.

14 `gameff` (\sqrt{V}) Effective body effect coefficient.

15 `betaeff` (A/V^2) Effective beta.

16 `cbd` (F) Drain-bulk junction capacitance.

17 `cbs` (F) Source-bulk junction capacitance.

Spectre Circuit Simulator Reference

Component Statements Part II

18	<code>cgs</code> (F)	Gate-source capacitance.
19	<code>cgd</code> (F)	Gate-drain capacitance.
20	<code>cgb</code> (F)	Gate-bulk capacitance.
21	<code>ron</code> (Ω)	On-resistance.
22	<code>id</code> (A)	Resistive drain current.
23	<code>ibulk</code> (A)	Resistive bulk current.
24	<code>pwr</code> (W)	Power at op point.
25	<code>gmoverid</code> (1/V)	Gm/Ids.
26	<code>isub</code> (A)	Substrate current.
27	<code>stress</code>	Hot-electron stress.
28	<code>age</code> (s)	Device age.
29	<code>he_vdsat</code> (V)	Hot Electron Vdsat.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

<code>ad</code>	I-4	<code>gap2</code>	M-74	<code>lmax</code>	M-103	<code>tlevc</code>	M-71
<code>ad</code>	M-89	<code>gds</code>	OP-12	<code>lmin</code>	M-104	<code>tnom</code>	M-66
<code>adef</code>	O-6	<code>gm</code>	OP-11	<code>lnom</code>	M-116	<code>tox</code>	M-18
<code>af</code>	M-98	<code>gmbs</code>	OP-13	<code>ls</code>	I-10	<code>tpg</code>	M-13
<code>age</code>	OP-28	<code>gmoverid</code>	OP-25	<code>m</code>	I-11	<code>trd</code>	M-78
<code>ai0</code>	M-19	<code>h0</code>	M-131	<code>m0</code>	M-133	<code>trise</code>	M-67
<code>alarm</code>	M-61	<code>hdif</code>	M-41	<code>meto</code>	M-28	<code>trise</code>	I-13

Spectre Circuit Simulator Reference Component Statements Part II

as	I-3	he_vdsat	OP-29	mgd	M-134	trs	M-77
as	M-88	hgd	M-132	minr	M-39	type	M-1
aseff	O-5	ibulk	OP-23	mj	M-54	type	OP-1
betaeff	OP-15	id	OP-22	mjsw	M-58	uo	M-7
bi0	M-22	ids	OP-5	n	M-47	ute	M-69
bvj	M-64	imax	M-62	nfs	M-12	uto	M-68
capmod	M-29	imelt	M-49	noisemod	M-96	vbox	M-65
cbd	M-52	is	M-46	nrd	M-92	vbs	OP-8
cbd	OP-16	isdeff	O-10	nrd	I-7	vbsn	M-117
cbdeff	O-12	isseff	O-9	nrs	M-93	vds	OP-7
cbs	OP-17	isub	OP-26	nrs	I-8	vdsat	OP-10
cbs	M-51	jmax	M-63	nss	M-11	vdsng	M-120
cbseff	O-11	jmelt	M-50	nsub	M-10	vdsni	M-118
cgb	OP-20	js	M-45	pb	M-55	vgs	OP-6
cgbo	M-27	kf	M-97	pbsw	M-59	vgsng	M-121
cgd	OP-19	kp	M-3	pd	M-91	vgsni	M-119
cgdo	M-26	l	I-2	pd	I-6	vmax	M-8
cgs	OP-18	l	M-87	pdeff	O-8	vpb	M-125
cgso	M-25	lai0	M-20	phi	M-5	vpg	M-124
cj	M-53	lambda	M-4	ps	I-5	vth	OP-9
cjsw	M-57	lamex	M-76	ps	M-90	vto	M-2
crigm	M-113	lbi0	M-23	pseff	O-7	w	I-1
criids	M-114	lc0	M-144	pta	M-82	w	M-86
criuo	M-112	lc1	M-147	ptc	M-80	wai0	M-21
crivth	M-111	lc2	M-150	ptp	M-83	wbi0	M-24
cta	M-84	lc3	M-153	pwr	OP-24	wd	M-15
ctp	M-85	lc4	M-156	rd	M-33	wecrit0	M-137
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degradation	I-14	lc6	M-162	rdd	M-35	wecritg	M-140
degradation	M-106	lc7	M-165	rdeff	O-4	weff	O-1
degramod	M-105	ld	M-14	region	OP-2	wlc0	M-146
dskip	M-48	ld	I-9	region	I-12	wlc1	M-149

Spectre Circuit Simulator Reference Component Statements Part II

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dvthe	M-108	lecrit0	M-136	rsc	M-37	wlc5	M-161
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ecritb	M-141	lecritg	M-139	rsh	M-36	wlc7	M-167
ecritg	M-138	leff	O-2	rss	M-34	wmax	M-101
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gap1	M-73	llc7	M-166	tlev	M-70		

Compact MOS-Transistor Distortion Model (mos1000)

Description

The mos10.00 model is an experimental model based on the thesis of Ronald van Langevelde: "A compact MOSFET Model for Distortion Analysis in Analog Circuit Design", Technische Universiteit Eindhoven, 1998.

Note: In noise analysis, mos10.00 instances will not generate any contribution, since there are no noise sources included (yet) in the mos10.00 model.

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In extension to the description a minimum conductance `gmin` is inserted between the drain and source node, to aid convergence. The value of `gmin` is set by an options statement, default = 1e-12 S.

This device is supported within altergroups.

Spectre Circuit Simulator Reference

Component Statements Part II

This device is dynamically loaded from the shared object /vobs/spectre_dev/tools.sun4v/spectre/lib/cmi/3.0.doc/libphilips_sh.so

Instance Definition

```
Name d g s [b] ModelName parameter=value ...
```

Instance Parameters

- | | | |
|---|---------------|---|
| 1 | w=1.0 scale m | Drawn channel width in the lay-out. Scale set by option scale. |
| 2 | l=1.0 scale m | Drawn channel length in the lay-out. Scale set by option scale. |
| 3 | mult=1 | Number of devices in parallel. |
| 4 | area=1 | Alias of mult. |
| 5 | region=triode | Estimated DC operating region, used as a convergence aid. Possible values are off, triode, sat, or subth. |
| 6 | m=1 | Multiplicity factor. |

Model Definition

```
model modelName mos1000 parameter=value ...
```

Model Parameters

Device type parameters

- | | | |
|---|--------|--|
| 1 | type=n | Transistor gender. Possible values are n or p. |
|---|--------|--|

Geometry parameters

- | | | |
|---|--------------|--|
| 2 | ler=1.0e-6 m | Effective channel length of the reference transistor. |
| 3 | wer=1e-6 m | Effective channel width of the reference transistor. |
| 4 | lvar=0.0 m | Difference between the actual and the programmed poly-silicon gate length. |

Spectre Circuit Simulator Reference

Component Statements Part II

- 5 $lap=45.0e-9$ m Effective channel length reduction per side.
- 6 $wvar=-5.0e-9$ m Difference between the actual and the programmed field-oxide opening.
- 7 $wot=50.0e-9$ m Effective channel width reduction per side.

Threshold-voltage parameters

- 8 $vfbr=-518.9e-03$ V Flat-band voltage for reference transistor.
- 9 $stvfb=-1.2e-03$ V/K
Coefficient of temperature dependence of v_{fb} .
- 10 $slvfb=24.0e-09$ V m
Coefficient of length dependence of v_{fb} .
- 11 $sl2vfb=-1.1e-15$ V m²
Second coefficient of length dependence of v_{fb} .
- 12 $swvfb=4.400e-09$ V m
Coefficient of the width dependency of v_{fb} .
- 13 $kor=368.0e-03$ \sqrt{V}
Body effect coefficient for the reference transistor.
- 14 $slko=-8.240e-09$ \sqrt{V} m
Coefficient of the length dependence of k_o .
- 15 $sl2ko=-2.260e-15$ \sqrt{V} m²
Second coefficient of the length dependence of k_o .
- 16 $swko=5.86e-09$ \sqrt{V} m
Coefficient of the width dependence of k_o .
- 17 $phibr=0.6$ V Surface potential at strong inversion.

Channel-current parameters

- 18 $betsq=370.9e-06$ A/V²
Gain factor for an infinite square transistor.

Spectre Circuit Simulator Reference

Component Statements Part II

- 19 `etabet=1.6` Exponent of the temperature dependence of the gain factor.
- 20 `thesrr=16.10e-3 1/V2` Mobility degradation parameter due to surface roughness scattering.
- 21 `stthesr=0.0 1/(V2 K)` Coefficient of the temperature dependence of `thesr`.
- 22 `swthesr=0.0 1/(V2 m)` Coefficient of the width dependence of `thesr`.
- 23 `thephr=0.055 1/V` Mobility degradation parameter due to phonon scattering.
- 24 `sttheph=0.0 1/(V K)` Coefficient of the temperature dependence of `theph`.
- 25 `swtheph=0.0 1/(V m)` Coefficient of the width dependence of `theph`.
- 26 `etamobr=1.6` Effective field parameter for dependence on depletion charge.
- 27 `swetamob=0.0 1/m` Coefficient of the width dependence of `etamobr`.
- 28 `thersq=0.155 1/V` Coefficient of gate voltage independent part of series resistance.
- 29 `swther=0.0 1/(V m)` Coefficient of the width dependence of `ther`.
- 30 `ther1=0.0 V` Numerator of gate voltage independent part of series resistance.
- 31 `ther2=1.0 V` Denominator of gate voltage independent part of series resistance.
- 32 `thenr=0.480 1/V` Velocity saturation parameter due to optical phonon scattering.
- 33 `stthen=0.0 1/(V K)` Coefficient of the temperature dependence of `then`.
- 34 `swthen=0.0 1/(V m)` Coefficient of the width dependence of `then`.
- 35 `thepr=0.0 1/V` Velocity saturation parameter due to acoustic phonon scattering.

Spectre Circuit Simulator Reference

Component Statements Part II

- 36 $stthep=0.0 \text{ 1/(V K)}$ Coefficient of the temperature dependence of $thep$.
- 37 $swthep=0.0 \text{ 1/(V m)}$ Coefficient of the width dependence of $thep$.
- 38 $gthep=1.0$ Velocity saturation factor due to acoustic phonon scattering.
- 39 $thethr=3.227e-3 \text{ 1/V}^3$ Coefficient of self-heating.
- 40 $sltheth=2.460e-9 \text{ 1/(V}^3 \text{ m)}$ Coefficient of the length dependence of $theth$.
- 41 $swtheth=0.0 \text{ 1/(V}^3 \text{ m)}$ Coefficient of the width dependence of $theth$.

Sub-threshold parameters

- 42 $sdiblo=2.030e-03 \text{ 1/}\sqrt{\text{V}}$ Drain-induced barrier lowering parameter.
- 43 $sdiblexp=1.340$ Exponent of the length dependence of $sdibl$.
- 44 $dphi=0.800 \text{ V}$ Parameter for short-channel subthreshold behaviour.

Saturation parameters

- 45 $ssfsq=6.250e-03 \text{ 1/}\sqrt{\text{V}}$ Static feedback parameter.
- 46 $swssf=0.0 \text{ 1/(\}\sqrt{\text{V}} \text{ m)}$ Coefficient of the width dependence of ssf .
- 47 $alpsq=0.010 \text{ m}$ Characteristic length parameter for channel length modulation.
- 48 $swalp=0.0 \text{ m}$ Coefficient of the width dependence of alp .
- 49 $vp=0.075 \text{ V}$ Characteristic voltage of channel-length modulation.

Spectre Circuit Simulator Reference

Component Statements Part II

Smoothing parameters

- 50 $m_{\text{expo}}=0.093$ Smoothing factor.
- 51 $m_{\text{expl}}=0.065$ Coefficient of the length dependence of m_{exp} .

Weak-avalanche parameters

- 52 $a_{1r}=6$ Factor of the weak-avalanche current.
- 53 $sta1=0.0$ 1/K Coefficient of the temperature dependence of a_1 .
- 54 $sla1=1.30e-6$ m Coefficient of the length dependence of a_1 .
- 55 $swa1=3.0e-06$ m Coefficient of the width dependence of a_1 .
- 56 $a_{2r}=38.0$ V Exponent of the weak-avalanche current.
- 57 $sla2=1.00e-06$ V m Coefficient of the length dependence of a_2 .
- 58 $swa2=2.00e-06$ V m Coefficient of the width dependence of a_2 .
- 59 $a_{3r}=0.650$ Factor of the drain-source voltage above which weak-avalanche occurs.
- 60 $sla3=-550.0e-06$ m Coefficient of the length dependence of a_3 .
- 61 $swa3=0.0$ m Coefficient of the width dependence of a_3 .

Charge parameters

- 62 $t_{\text{ox}}=4.5e-09$ m Thickness of the oxide layer.
- 63 $col=320e-12$ F/m Gate overlap capacitance per unit channel width.

Temperature parameters

- 64 t_r (C) Reference temperature. Default set by option t_{nom} .
- 65 t_{ref} (C) Alias of t_r . Default set by option t_{nom} .
- 66 t_{nom} (C) Alias of t_r . Default set by option t_{nom} .

Spectre Circuit Simulator Reference

Component Statements Part II

67 `dta=0.0 K` Temperature offset of the device.

68 `trise=0.0 K` Alias of `dta`.

Output Parameters

1 `le (m)` Effective channel length.

2 `we (m)` Effective channel width.

3 `vfb (V)` Flat-band voltage.

4 `ko (\sqrt{V})` Body effect coefficient.

5 `phib (V)` Surface potential at strong inversion.

6 `bet (A/V^2)` Gain factor.

7 `thesr ($1/V^2$)` Mobility degradation parameter due to surface roughness scattering.

8 `theph ($1/V$)` Mobility degradation parameter due to phonon scattering.

9 `etamob` Effective field parameter for dependence on depletion charge.

10 `ther ($1/V$)` Coefficient of gate voltage independent part of series resistance.

11 `ther1 (V)` Numerator of gate voltage independent part of series resistance.

12 `ther2 (V)` Denominator of gate voltage independent part of series resistance.

13 `then ($1/V$)` Velocity saturation parameter due to optical phonon scattering.

14 `thep ($1/V$)` Velocity saturation parameter due to acoustic phonon scattering.

15 `gthep` Velocity saturation factor due to acoustic phonon scattering.

16 `theth ($1/V^3$)` Coefficient of self-heating.

17 `sdibl ($1/\sqrt{V}$)` Drain-induced barrier lowering parameter.

18 `dphi (V)` Parameter for short-channel subthreshold behaviour.

Spectre Circuit Simulator Reference

Component Statements Part II

19	ssf (1/ \sqrt{V})	Static feedback parameter.
20	alp (m)	Characteristic length parameter for channel length modulation.
21	vp (V)	Characteristic voltage of channel-length modulation.
22	mexp	Smoothing factor.
23	phit (V)	Thermal voltage.
24	a1	Factor of the weak-avalanche current.
25	a2 (V)	Exponent of the weak-avalanche current.
26	a3	Factor of the drain-source voltage above which weak-avalanche occurs.
27	cox (F)	Gate-to-channel capacitance (* mult).
28	cgdo (F)	Gate-drain overlap capacitance (* mult).
29	cgso (F)	Gate-source overlap capacitance (* mult).

Operating-Point Parameters

1	ide (A)	Resistive drain current.
2	ige (A)	Resistive gate current.
3	ise (A)	Resistive source current.
4	ibe (A)	Resistive bulk current.
5	vds (V)	Drain-source voltage.
6	vgs (V)	Gate-source voltage.
7	vsb (V)	Source-bulk voltage.
8	ids (A)	Resistive drain current.
9	idb (A)	Resistive drain-bulk current.

Spectre Circuit Simulator Reference

Component Statements Part II

10	<code>isb</code> (A)	Resistive source-bulk current.
11	<code>iavl</code> (A)	Substrate current.
12	<code>pwr</code> (W)	Power.
13	<code>vto</code> (V)	Threshold voltage at zero back-bias.
14	<code>vts</code> (V)	v_{ts} .
15	<code>vgt</code> (V)	Effective gate drive including backbias and drain effects.
16	<code>vdss</code> (V)	Saturation voltage at actual bias.
17	<code>vsat</code> (V)	Saturation limit.
18	<code>gm</code> (S)	Transconductance ($d i_{ds} / d v_{gs}$).
19	<code>gmb</code> (S)	Bulk transconductance ($d i_{ds} / d v_{bs}$).
20	<code>gds</code> (S)	Output conductance ($d i_{ds} / d v_{ds}$).
21	<code>cdd</code> (F)	Capacitance ($d q_d / d v_d$).
22	<code>cdg</code> (F)	Capacitance ($- d q_d / d v_g$).
23	<code>cds</code> (F)	Capacitance ($- d q_d / d v_s$).
24	<code>cdb</code> (F)	Capacitance ($- d q_d / d v_b$).
25	<code>cgd</code> (F)	Capacitance ($- d q_g / d v_d$).
26	<code>cgg</code> (F)	Capacitance ($d q_g / d v_g$).
27	<code>cgs</code> (F)	Capacitance ($- d q_g / d v_s$).
28	<code>cgb</code> (F)	Capacitance ($- d q_g / d v_b$).
29	<code>csd</code> (F)	Capacitance ($- d q_s / d v_d$).
30	<code>csg</code> (F)	Capacitance ($- d q_s / d v_g$).
31	<code>css</code> (F)	Capacitance ($d q_s / d v_s$).

Spectre Circuit Simulator Reference

Component Statements Part II

32	<code>csb</code> (F)	Capacitance (- d qs / d vb).
33	<code>cbd</code> (F)	Capacitance (- d qb / d vd).
34	<code>cbg</code> (F)	Capacitance (- d qb / d vg).
35	<code>cbs</code> (F)	Capacitance (- d qb / d vs).
36	<code>cbb</code> (F)	Capacitance (d qb / d vb).
37	<code>u</code>	Transistor gain (gm/gds).
38	<code>rout</code> (Ω)	Small signal output resistance (1/gds).
39	<code>vearly</code> (V)	Equivalent Early voltage ($ id /gds$).
40	<code>keff</code> (\sqrt{V})	Describes body effect at actual bias.
41	<code>beff</code> (S/V)	Effective beta at actual bias in the simple MOS model ($2* ids /vgt^2$).
42	<code>fug</code> (Hz)	Unity gain frequency at actual bias ($gm/(2*pi*cin)$).

Parameter Index

In the following index, I refers to instance parameters, M refers to the model parameters section, O refers to the output parameters section, and OP refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of M-35 means the 35th model parameter.

<code>a1</code>	O-24	<code>etamobr</code>	M-26	<code>sl2ko</code>	M-15	<code>ther1</code>	M-30
<code>a1r</code>	M-52	<code>fug</code>	OP-42	<code>sl2vfb</code>	M-11	<code>ther1</code>	O-11
<code>a2</code>	O-25	<code>gds</code>	OP-20	<code>sla1</code>	M-54	<code>ther2</code>	O-12
<code>a2r</code>	M-56	<code>gm</code>	OP-18	<code>sla2</code>	M-57	<code>ther2</code>	M-31
<code>a3</code>	O-26	<code>gmb</code>	OP-19	<code>sla3</code>	M-60	<code>thersq</code>	M-28
<code>a3r</code>	M-59	<code>gthep</code>	M-38	<code>slko</code>	M-14	<code>thesr</code>	O-7
<code>alp</code>	O-20	<code>gthep</code>	O-15	<code>sltheth</code>	M-40	<code>thesrr</code>	M-20
<code>alpsq</code>	M-47	<code>iavl</code>	OP-11	<code>slvfb</code>	M-10	<code>theth</code>	O-16

Spectre Circuit Simulator Reference Component Statements Part II

area	I-4	ibe	OP-4	ssf	O-19	thethr	M-39
beff	OP-41	idb	OP-9	ssfsq	M-45	tnom	M-66
bet	O-6	ide	OP-1	stal	M-53	tox	M-62
betsq	M-18	ids	OP-8	stthen	M-33	tr	M-64
cbb	OP-36	ige	OP-2	stthep	M-36	tref	M-65
cbd	OP-33	isb	OP-10	sttheph	M-24	trise	M-68
cbg	OP-34	ise	OP-3	stthesr	M-21	type	M-1
cbs	OP-35	keff	OP-40	stvfb	M-9	u	OP-37
cdb	OP-24	ko	O-4	swal	M-55	vds	OP-5
cdd	OP-21	kor	M-13	swa2	M-58	vdss	OP-16
cdg	OP-22	l	I-2	swa3	M-61	vearly	OP-39
cds	OP-23	lap	M-5	swalp	M-48	vfb	O-3
cgb	OP-28	le	O-1	swetamob	M-27	vfbr	M-8
cgd	OP-25	ler	M-2	swko	M-16	vgs	OP-6
cgdo	O-28	lvar	M-4	swssf	M-46	vgt	OP-15
cgg	OP-26	m	I-6	swthen	M-34	vp	O-21
cgs	OP-27	mexp	O-22	swthep	M-37	vp	M-49
cgso	O-29	mexpl	M-51	swtheph	M-25	vsat	OP-17
col	M-63	mexpo	M-50	swther	M-29	vsb	OP-7
cox	O-27	mult	I-3	swthesr	M-22	vto	OP-13
csb	OP-32	phib	O-5	swtheth	M-41	vts	OP-14
csd	OP-29	phibr	M-17	swvfb	M-12	w	I-1
csg	OP-30	phit	O-23	then	O-13	we	O-2
css	OP-31	pwr	OP-12	thenr	M-32	wer	M-3
dphi	O-18	region	I-5	thep	O-14	wot	M-7
dphi	M-44	rout	OP-38	theph	O-8	wvar	M-6
dta	M-67	sdibl	O-17	thephr	M-23		
etabet	M-19	sdiblexp	M-43	thepr	M-35		
etamob	O-9	sdiblo	M-42	ther	O-10		

Compact MOS-Transistor Distortion Model (mos1100)

Description

The mos1100 model is based on the thesis of Ronald van Langevelde: "A compact MOSFET Model for Distortion Analysis in Analog Circuit Design", Technische Universiteit Eindhoven, 1998.

(c) Philips Electronics N.V. 2001

In extension to the description a minimum conductance g_{min} is inserted between the drain and source node, to aid convergence. The value of g_{min} is set by an options statement, default = 1e-12 S.

This device is supported within altergroups.

This device is dynamically loaded from the shared object /vobs/spectre_dev/tools.sun4v/spectre/lib/cmi/3.0.doc/libphilips_sh.so

Instance Definition

```
Name d g s [b] ModelName parameter=value ...
```

Instance Parameters

- | | | |
|---|---------------|--|
| 1 | w=1.0 scale m | Drawn channel width in the layout. Scale set by option scale.. |
| 2 | l=1.0 scale m | Drawn channel length in the layout. Scale set by option scale.. |
| 3 | mult=1 | Number of devices in parallel. |
| 4 | area=1 | Alias of mult. |
| 5 | region=triode | Estimated DC operating region, used as a convergence aid.
Possible values are off, triode, sat, or subth. |
| 6 | m=1 | Multiplicity factor. |

Model Definition

```
model modelName mos1100 parameter=value ...
```

Spectre Circuit Simulator Reference

Component Statements Part II

Model Parameters

Device type parameters

1 `type=n` Transistor gender.
Possible values are n or p.

Geometry parameters

2 `ler=1.0e-6 m` Effective channel length of the reference transistor.

3 `wer=1e-5 m` Effective channel width of the reference transistor.

4 `lvar=0.0 m` Difference between the actual and the programmed poly-silicon gate length.

5 `lap=4.0e-8 m` Effective channel length reduction per side.

6 `wvar=0.0 m` Difference between the actual and the programmed field-oxide opening.

7 `wot=0.0 m` Effective channel width reduction per side.

Threshold-voltage parameters

8 `vfbr=-1.050 V` Flat-band voltage for reference transistor.

9 `stvfb=0.5e-03 V/K` Coefficient of temperature dependence of `vfbr`.

10 `kor=0.5 \sqrt{V}` Body effect coefficient for the reference transistor.

11 `slko=0.0 \sqrt{V} m` Coefficient of the length dependence of `ko`.

12 `sl2ko=0.0 \sqrt{V} m2` Second coefficient of the length dependence of `ko`.

13 `swko=0.0 \sqrt{V} m` Coefficient of the width dependence of `ko`.

14 `phibr=0.95 V` Surface potential at strong inversion.

Spectre Circuit Simulator Reference

Component Statements Part II

15	slphib=0.0 Vm	Coefficient of the length dependence of phib.
16	sl2phib=0.0 Vm ²	Second coefficient of the length dependence of phib.
17	swphib=0.0 Vm ²	Coefficient of the width dependence of phib.
Channel-current parameters		
18	fbet1=0.0	Relative mobility decrease due to first lateral profile.
19	lp1=0.8e-6	Characteristic length of first lateral profile.
20	lp2=0.8e-6	Characteristic length of second lateral profile.
21	fbet2=0.0	Relative mobility decrease due to second lateral profile.
22	betsq=370.9e-06 A/V ²	Gain factor for an infinite square transistor.
23	etabet=1.3	Exponent of the temperature dependence of the gain factor.
24	thesrr=0.4 1/V	Coefficient of the mobility reduction due to surface roughness scattering.
25	swthesr=0.0 m	Coefficient of the width dependence of thesr.
26	thephr=1.29e-2 1/V	Coefficient of the mobility reduction due to phonon scattering.
27	swtheph=0.0 m	Coefficient of the width dependence of theph.
28	etaph=1.75	Exponent of the temperature dependence of theph.
29	etamobr=1.4	Effective field parameter for dependence on depletion/inversion charge.
30	stetamob=0.0 1/K	Coefficient of the temperature dependence of etamob.
31	swetamob=0.0 m	Coefficient of the width dependence of etamob.
32	nur=1.0	Exponent of the field dependence of the mobility model minus 1.
33	nuexp=5.25	Exponent of the temperature dependence of parameter nu.

Spectre Circuit Simulator Reference

Component Statements Part II

34	<code>therr=0.155</code>	$1/V$	Coefficient of the series resistance.
35	<code>swther=0.0</code>	m	Coefficient of the width dependence of ther.
36	<code>etar=0.95</code>		Exponent of the temperature dependence of ther.
37	<code>thesatr=0.5</code>	$1/V$	Velocity saturation parameter due to optical/acoustic phonon scattering.
38	<code>etasat=1.04</code>	$1/V$	Exponent of the temperature dependence of thesat.
39	<code>swthesat=0.0</code>	m	Coefficient of the width dependence of thesat.
40	<code>slthesat=1.0</code>		Coefficient of length dependence of thesat.
41	<code>thesatexp=1.0</code>		Exponent of length dependence of thesat.

Drain-feedback parameters

42	<code>thethr=1e-3</code>	$1/V^3$	Coefficient of self-heating.
43	<code>thethexp=1.0</code>		Exponent of the length dependence of theth.
44	<code>ssfr=6.25e-3</code>	$1/\sqrt{V}$	Static feedback parameter.
45	<code>swssf=0.0</code>	m	Coefficient of the width dependence of ssf.
46	<code>slssf=1.0e-6</code>	m	Coefficient of the length dependence of ssf.
47	<code>alp_r=0.010</code>		Factor of the channel length modulation.
48	<code>swalp=0.0</code>	m	Coefficient of the width dependence of alp.
49	<code>slalp=1.0</code>	m	Coefficient of the length dependence of alp.
50	<code>alp_{exp}=1.0</code>		Exponent of the length dependence of alp.
51	<code>vp=5.0e-2</code>	V	Characteristic voltage of channel-length modulation.

Spectre Circuit Simulator Reference

Component Statements Part II

Sub-threshold parameters

- 52 `sdiblo=2e-03 1/√V` Drain-induced barrier lowering parameter.
- 53 `sdiblexp=1.35` Exponent of the length dependence of `sdibl`.
- 54 `mor=0.0` Parameter for short-channel subthreshold slope.
- 55 `moexp=1.34` Exponent of the length dependence of `mo`.

Smoothing parameter

- 56 `lmin=1.5e-7 m` Minimum effective channel length in technology, used for calculation of smoothing factor `m`.

Weak-avalanche parameters

- 57 `a1r=6` Factor of the weak-avalanche current.
- 58 `sla1=0.0 m` Coefficient of the length dependence of `a1`.
- 59 `swa1=0.0 m` Coefficient of the width dependence of `a1`.
- 60 `sta1=0.0 1/K` Coefficient of the temperature dependence of `a1`.
- 61 `a2r=38.0 V` Exponent of the weak-avalanche current.
- 62 `sla2=0.0 V m` Coefficient of the length dependence of `a2`.
- 63 `swa2=0.0 V m` Coefficient of the width dependence of `a2`.
- 64 `a3r=1.0` Factor of the drain-source voltage above which weak-avalanche occurs.
- 65 `sla3=0.0 m` Coefficient of the length dependence of `a3`.
- 66 `swa3=0.0 m` Coefficient of the width dependence of `a3`.

Gate current parameters

- 67 `iginvr=0.0 A/V2` Gain factor for intrinsic gate tunnelling current in inversion.

Spectre Circuit Simulator Reference

Component Statements Part II

68 $igaccr=0.0 \text{ A/V}^2$ Gain factor for intrinsic gate tunnelling current in accumulation.

69 $igovr=0.0 \text{ A/V}^2$ Gain factor for Source/Drain overlap gate tunnelling current.

Charge parameters

70 $tox=3.2e-09 \text{ m}$ Thickness of the oxide layer.

71 $col=3.2e-10 \text{ F/m}$ Gate overlap capacitance per unit channel width.

Noise parameters

72 $ntr=1.656e-20 \text{ J}$ Coefficient of the thermal noise.

73 $nfar=1.573e22 \text{ 1/(Vm}^4\text{)}$
First coefficient of the flicker noise.

74 $nfbr=4.752e8 \text{ 1/(Vm}^2\text{)}$
Second coefficient of the flicker noise.

75 $nfcr=0.0 \text{ 1/V}$ Third coefficient of the flicker noise.

Temperature parameters

76 $tr \text{ (C)}$ Reference temperature. Default set by option $tnom$.

77 $tref \text{ (C)}$ Alias of tr . Default set by option $tnom$.

78 $tnom \text{ (C)}$ Alias of tr . Default set by option $tnom$.

79 $dta=0.0 \text{ K}$ Temperature offset of the device.

80 $trise=0.0 \text{ K}$ Alias of dta .

Other parameters

81 $kpinv=0.0 \text{ 1}/\sqrt{V}$
Inverse of body-effect factor of the poly-silicon gate.

82 $ther1=0.0 \text{ V}$ Numerator of gate voltage dependent part of series resistance.

Spectre Circuit Simulator Reference

Component Statements Part II

83	<code>ther2=1.0 V</code>	Denominator of gate voltage dependent part of series resistance.
84	<code>vp=5.0e-2 V</code>	Characteristic voltage of channel-length modulation.
85	<code>binv=48.0 V</code>	Probability factor for intrinsic gate tunnelling current in inversion.
86	<code>bacc=48.0 V</code>	Probability factor for intrinsic gate tunnelling current in accumulation.
87	<code>vfbov=0.0 V</code>	Flat-band voltage for the Source/Drain overlap extensions.
88	<code>kov=2.5 \sqrt{V}</code>	Body-effect factor for the Source/Drain overlap extensions.
89	<code>gatenoise=0.0</code>	Flag for in/exclusion of induced gate thermal noise.

Output Parameters

1	<code>vt0 (V)</code>	Zero-bias threshold voltage.
2	<code>le (m)</code>	Effective channel length.
3	<code>we (m)</code>	Effective channel width.
4	<code>ko (\sqrt{V})</code>	Body-effect factor.
5	<code>phib (V)</code>	Surface potential at the onset of strong inversion.
6	<code>bet (A/V^2)</code>	Gain factor.
7	<code>thesr (1/V)</code>	Mobility degradation parameter due to surface roughness scattering.
8	<code>theph (1/V)</code>	Mobility degradation parameter due to phonon scattering.
9	<code>etamob</code>	Effective field parameter for dependence on depletion charge.
10	<code>nu</code>	Exponent of field dependence of mobility model.
11	<code>ther (1/V)</code>	Coefficient of series resistance.
12	<code>thesat (1/V)</code>	Velocity saturation parameter due to optical/acoustic phonon scattering.

Spectre Circuit Simulator Reference

Component Statements Part II

13	theth (1/V ³)	Coefficient of self-heating.
14	sdibl (1/√V)	Drain-induced barrier lowering parameter.
15	mo	Parameter for (short-channel) subthreshold slope.
16	ssf (1/√V)	Static-feedback parameter.
17	alp	Factor of channel length modulation.
18	mexp	Smoothing factor.
19	a1	Factor of the weak-avalanche current.
20	a2 (V)	Exponent of the weak-avalanche current.
21	a3	Factor of the drain-source voltage above which weak-avalanche occurs.
22	iginv (A/V ²)	Gain factor for intrinsic gate tunnelling current in inversion.
23	igacc (A/V ²)	Gain factor for intrinsic gate tunnelling current in accumulation.
24	igov (A/V ²)	Gain factor for Source/Drain overlap tunnelling current.
25	cox (F)	Oxide capacitance for the intrinsic channel (* mult).
26	cgdo (F)	Oxide capacitance for the gate-drain overlap (* mult).
27	cgso (F)	Oxide capacitance for the gate-source overlap (* mult).
28	nt (J)	Thermal noise coefficient.
29	nfa (1/(Vm ⁴))	First coefficient of the flicker noise.
30	nfb (1/(Vm ²))	Second coefficient of the flicker noise.
31	nfc (1/V)	Third coefficient of the flicker noise.
32	tox (m)	Thickness of gate oxide layer.

Spectre Circuit Simulator Reference

Component Statements Part II

Operating-Point Parameters

1	<code>nfc_r=0.0</code>	1/V	Third coefficient of the flicker noise.
2	<code>ide</code>	(A)	Resistive drain current.
3	<code>ige</code>	(A)	Resistive gate current.
4	<code>ise</code>	(A)	Resistive source current.
5	<code>ibe</code>	(A)	Resistive bulk current.
6	<code>isb</code>	(A)	Resistive source-bulk current.
7	<code>idb</code>	(A)	Resistive drain-bulk current.
8	<code>pwr</code>	(W)	Power.
9	<code>ids</code>	(A)	Drain current, excl. avalanche and tunnel currents.
10	<code>iavl</code>	(A)	Substrate current due to weak-avalanche.
11	<code>igs</code>	(A)	Gate-to-source current due to direct tunnelling.
12	<code>igd</code>	(A)	Gate-to-drain current due to direct tunnelling.
13	<code>igb</code>	(A)	Gate-to-bulk current due to direct tunnelling.
14	<code>vds</code>	(V)	Drain-source voltage.
15	<code>vgs</code>	(V)	Gate-source voltage.
16	<code>vsb</code>	(V)	Source-bulk voltage.
17	<code>vts</code>	(V)	Threshold voltage including back-bias effects.
18	<code>vth</code>	(V)	Threshold voltage including back-bias and drain-bias effects.
19	<code>vgt</code>	(V)	Effective gate drive voltage including back-bias and drain voltage effects.
20	<code>vdss</code>	(V)	Drain saturation voltage at actual bias.

Spectre Circuit Simulator Reference

Component Statements Part II

21	v_{sat} (V)	Saturation limit.
22	g_m (S)	Transconductance ($d i_{ds} / d v_{gs}$).
23	g_{mb} (S)	Substrate-transconductance ($d i_{ds} / d v_{bs}$).
24	g_{ds} (S)	Output conductance ($d i_{ds} / d v_{ds}$).
25	c_{dd} (F)	Capacitance ($d q_d / d v_d$).
26	c_{dg} (F)	Capacitance ($- d q_d / d v_g$).
27	c_{ds} (F)	Capacitance ($- d q_d / d v_s$).
28	c_{db} (F)	Capacitance ($- d q_d / d v_b$).
29	c_{gd} (F)	Capacitance ($- d q_g / d v_d$).
30	c_{gg} (F)	Capacitance ($d q_g / d v_g$).
31	c_{gs} (F)	Capacitance ($- d q_g / d v_s$).
32	c_{gb} (F)	Capacitance ($- d q_g / d v_b$).
33	c_{sd} (F)	Capacitance ($- d q_s / d v_d$).
34	c_{sg} (F)	Capacitance ($- d q_s / d v_g$).
35	c_{ss} (F)	Capacitance ($d q_s / d v_s$).
36	c_{sb} (F)	Capacitance ($- d q_s / d v_b$).
37	c_{bd} (F)	Capacitance ($- d q_b / d v_d$).
38	c_{bg} (F)	Capacitance ($- d q_b / d v_g$).
39	c_{bs} (F)	Capacitance ($- d q_b / d v_s$).
40	c_{bb} (F)	Capacitance ($d q_b / d v_b$).
41	u	Transistor gain (g_m/g_{ds}).
42	r_{out} (Ω)	Small-signal output resistance ($1/g_{ds}$).

Spectre Circuit Simulator Reference

Component Statements Part II

43	vearly (V)	Equivalent Early voltage ($ id /gds$).
44	keff (\sqrt{V})	Body effect parameter.
45	beff (A/V^2)	Gain factor.
46	fug (Hz)	Unity gain frequency at actual bias ($gm/(2*\pi*cin)$).
47	sqrtsfw (V/\sqrt{Hz})	Input-referred RMS white noise voltage density.
48	sqrtsff (V/\sqrt{Hz})	Input-referred RMS white noise voltage density at 1 kHz.
49	fknee (Hz)	Cross-over frequency above which white noise is dominant.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

a1	O-19	fknee	OP-49	nfb	O-30	swther	M-35
alr	M-57	fug	OP-46	nibr	M-74	swthesat	M-39
a2	O-20	gatenoise	M-89	nfc	O-31	swthesr	M-25
a2r	M-61	gds	OP-24	nibr	M-75	theph	O-8
a3	O-21	gm	OP-22	nibr	OP-1	theph	M-26
a3r	M-64	gmb	OP-23	nt	O-28	ther	O-11
alp	O-17	iavl	OP-10	ntr	M-72	ther1	M-82
alpexp	M-50	ibe	OP-5	nu	O-10	ther2	M-83
alpr	M-47	idb	OP-7	nuexp	M-33	therr	M-34
area	I-4	ide	OP-2	nur	M-32	thesat	O-12
bacc	M-86	ids	OP-9	phib	O-5	thesatexp	M-41
beff	OP-45	igacc	O-23	phibr	M-14	thesatr	M-37
bet	O-6	igaccr	M-68	pwr	OP-8	thesr	O-7

Spectre Circuit Simulator Reference Component Statements Part II

betsq M-22	igb OP-13	region I-5	thesrr M-24
binv M-85	igd OP-12	rout OP-42	theth O-13
cbb OP-40	ige OP-3	sdibl O-14	thethexp M-43
cbd OP-37	iginv O-22	sdiblexp M-53	thethr M-42
cbg OP-38	iginvr M-67	sdiblo M-52	tnom M-78
cbs OP-39	igov O-24	sl2ko M-12	tox O-32
cdb OP-28	igovr M-69	sl2phib M-16	tox M-70
cdd OP-25	igs OP-11	sla1 M-58	tr M-76
cdg OP-26	isb OP-6	sla2 M-62	tref M-77
cds OP-27	ise OP-4	sla3 M-65	trise M-80
cgb OP-32	keff OP-44	slalp M-49	type M-1
cgd OP-29	ko O-4	slko M-11	u OP-41
cgdo O-26	kor M-10	slphib M-15	vds OP-14
cgg OP-30	kov M-88	slssf M-46	vdss OP-20
cgs OP-31	kpinv M-81	slthesat M-40	vearly OP-43
cgso O-27	l I-2	sqrtsff OP-48	vfbov M-87
col M-71	lap M-5	sqrtsfw OP-47	vfbr M-8
cox O-25	le O-2	ssf O-16	vgs OP-15
csb OP-36	ler M-2	ssfr M-44	vgt OP-19
csd OP-33	lmin M-56	sta1 M-60	vp M-84
csg OP-34	lp1 M-19	stetamob M-30	vp M-51
css OP-35	lp2 M-20	stvfb M-9	vsat OP-21
dta M-79	lvar M-4	swa1 M-59	vsb OP-16
etabet M-23	m I-6	swa2 M-63	vth OP-18
etamob O-9	mexp O-18	swa3 M-66	vto O-1
etamobr M-29	mo O-15	swalp M-48	vts OP-17
etaph M-28	moexp M-55	swetamob M-31	w I-1
etar M-36	mor M-54	swko M-13	we O-3
etasat M-38	mult I-3	swphib M-17	wer M-3
fbet1 M-18	nfa O-29	swssf M-45	wot M-7
fbet2 M-21	nfarr M-73	swtheph M-27	wvar M-6

Compact MOS-Transistor Distortion Model (mos11010)

Description

The mos1101 model is based on the thesis of Ronald van Langevelde: "A compact MOSFET Model for Distortion Analysis in Analog Circuit Design", Technische Universiteit Eindhoven, 1998.

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In extension to the description a minimum conductance g_{min} is inserted between the drain and source node, to aid convergence. The value of g_{min} is set by an options statement, default = 1e-12 S.

This device is supported within altergroups.

This device is dynamically loaded from the shared object /vobs/spectre_dev/tools.sun4v/spectre/lib/cmi/3.0.doc/libphilips_sh.so

Instance Definition

```
Name d g s [b] ModelName parameter=value ...
```

Instance Parameters

- | | | |
|---|---------------|--|
| 1 | w=1.0 scale m | Drawn channel width in the layout. Scale set by option scale.. |
| 2 | l=1.0 scale m | Drawn channel length in the layout. Scale set by option scale.. |
| 3 | mult=1 | Number of devices in parallel. |
| 4 | area=1 | Alias of mult. |
| 5 | region=triode | Estimated DC operating region, used as a convergence aid.
Possible values are off, triode, sat, or subth. |
| 6 | m=1 | Multiplicity factor. |

Model Definition

```
model modelName mos11010 parameter=value ...
```

Spectre Circuit Simulator Reference

Component Statements Part II

Model Parameters

Geometry parameters

- | | | |
|---|---------------------------|--|
| 1 | <code>lvar=0.0 m</code> | Difference between the actual and the programmed poly-silicon gate length. |
| 2 | <code>lap=4.0e-8 m</code> | Effective channel length reduction per side. |
| 3 | <code>wvar=0.0 m</code> | Difference between the actual and the programmed field-oxide opening. |
| 4 | <code>wot=0.0 m</code> | Effective channel width reduction per side. |

Device type parameters

- | | | |
|----|-----------------------------|--|
| 5 | <code>type=n</code> | Transistor gender.
Possible values are <code>n</code> or <code>p</code> . |
| 6 | <code>level=1.0</code> | Transistor Level. Possible value is 1 (Philips MOSFET Modelbook June 2002). |
| 7 | <code>vfb=-1.050 V</code> | Flat-band voltage at reference temperature. |
| 8 | <code>ther1=0.0 V</code> | Numerator of gate voltage dependent part of series resistance. |
| 9 | <code>ther2=1.0 V</code> | Denominator of gate voltage dependent part of series resistance. |
| 10 | <code>vp=5.0e-2 V</code> | Characteristic voltage of channel-length modulation. |
| 11 | <code>dta=0.0 K</code> | Temperature offset of the device. |
| 12 | <code>trise=0.0 K</code> | Alias of <code>dta</code> . |
| 13 | <code>nt=1.656e-20 J</code> | Coefficient of the thermal noise. |
| 14 | <code>tox=3.2e-09 m</code> | Thickness of the oxide layer. |

Threshold-voltage parameters

- | | | |
|----|--------------------------------|---|
| 15 | <code>stvfb=0.5e-03 V/K</code> | Coefficient of temperature dependence of <code>vfb</code> . |
|----|--------------------------------|---|

Spectre Circuit Simulator Reference

Component Statements Part II

- 16 $k_{or}=0.5 \sqrt{V}$ Body effect coefficient for the reference transistor.
- 17 $slk_o=0.0 \sqrt{V} \text{ m}$ Coefficient of the length dependence of k_o .
- 18 $sl2k_o=0.0 \sqrt{V} \text{ m}^2$ Second coefficient of the length dependence of k_o .
- 19 $swk_o=0.0 \sqrt{V} \text{ m}$ Coefficient of the width dependence of k_o .
- 20 $phibr=0.95 \text{ V}$ Surface potential at strong inversion.
- 21 $slphib=0.0 \text{ Vm}$ Coefficient of the length dependence of $phib$.
- 22 $sl2phib=0.0 \text{ Vm}^2$ Second coefficient of the length dependence of $phib$.
- 23 $swphib=0.0 \text{ Vm}^2$ Coefficient of the width dependence of $phib$.
- 24 $stphib=-8.5e-4 \text{ VK}^{-1}$ Coefficient of the temperature dependency of $PhiB$.

Channel-current parameters

- 25 $fbet1=0.0$ Relative mobility decrease due to first lateral profile.
- 26 $lp1=0.8e-6$ Characteristic length of first lateral profile.
- 27 $lp2=0.8e-6$ Characteristic length of second lateral profile.
- 28 $fbet2=0.0$ Relative mobility decrease due to second lateral profile.
- 29 $betsq=370.9e-06 \text{ A/V}^2$ Gain factor for an infinite square transistor.
- 30 $etabetr=1.3$ Exponent of the temperature dependence of the gain factor.
- 31 $sletabet=0.0$ Coefficient of length dependence of nbr .
- 32 $etasr=0.65$ Exponent of the temperature dependence of $thetaSR$.
- 33 $thesrr=0.4 \text{ 1/V}$ Coefficient of the mobility reduction due to surface roughness scattering.

Spectre Circuit Simulator Reference

Component Statements Part II

- 34 `swthesr=0.0 m` Coefficient of the width dependence of `thesr`.
- 35 `thephr=1.29e-2 1/V` Coefficient of the mobility reduction due to phonon scattering.
- 36 `swtheeph=0.0 m` Coefficient of the width dependence of `theeph`.
- 37 `etaph=1.35` Exponent of the temperature dependence of `theeph`.
- 38 `etamobr=1.4` Effective field parameter for dependence on depletion/inversion charge.
- 39 `stetamob=0.0 1/K` Coefficient of the temperature dependence of `etamob`.
- 40 `swetamob=0.0 m` Coefficient of the width dependence of `etamob`.
- 41 `nu=1.0` Exponent of the field dependence of the mobility model minus 1.
- 42 `nuexp=5.25` Exponent of the temperature dependence of parameter `nu`.
- 43 `therr=0.155 1/V` Coefficient of the series resistance.
- 44 `swther=0.0 m` Coefficient of the width dependence of `ther`.
- 45 `etar=0.95` Exponent of the temperature dependence of `ther`.
- 46 `thesatr=0.5 1/V` Velocity saturation parameter due to optical/acoustic phonon scattering.
- 47 `etasat=1.04 1/V` Exponent of the temperature dependence of `thesat`.
- 48 `swthesat=0.0 m` Coefficient of the width dependence of `thesat`.
- 49 `slthesat=1.0` Coefficient of length dependence of `thesat`.
- 50 `thesatexp=1.0` Exponent of length dependence of `thesat`.

Drain-feedback parameters

- 51 `thethr=1e-3 1/V3` Coefficient of self-heating.
- 52 `thethexp=1.0` Exponent of the length dependence of `theth`.

Spectre Circuit Simulator Reference

Component Statements Part II

- 53 `ssfr=6.25e-3 1/√V` Static feedback parameter.
- 54 `swssf=0.0 m` Coefficient of the width dependence of `ssf`.
- 55 `slssf=1.0e-6 m` Coefficient of the length dependence of `ssf`.
- 56 `alpr=0.010` Factor of the channel length modulation.
- 57 `swalp=0.0 m` Coefficient of the width dependence of `alp`.
- 58 `slalp=1.0 m` Coefficient of the length dependence of `alp`.
- 59 `alpexp=1.0` Exponent of the length dependence of `alp`.

Sub-threshold parameters

- 60 `sdiblo=1.0e-4 1/√V` Drain-induced barrier lowering parameter.
- 61 `sdiblexp=1.35` Exponent of the length dependence of `sdibl`.
- 62 `mor=0.0` Parameter for short-channel subthreshold slope.
- 63 `moexp=1.34` Exponent of the length dependence of `mo`.
- 64 `mo0=0.0` Offset parameter for (short-channel) subthreshold slope.

Smoothing parameter

- 65 `lmin=1.5e-7 m` Minimum effective channel length in technology, used for calculation of smoothing factor `m`.

Weak-avalanche parameters

- 66 `alr=6` Factor of the weak-avalanche current.
- 67 `sla1=0.0 m` Coefficient of the length dependence of `a1`.
- 68 `swa1=0.0 m` Coefficient of the width dependence of `a1`.
- 69 `sta1=0.0 1/K` Coefficient of the temperature dependence of `a1`.

Spectre Circuit Simulator Reference

Component Statements Part II

70	$a2r=38.0$	V	Exponent of the weak-avalanche current.
71	$sla2=0.0$	V m	Coefficient of the length dependence of $a2$.
72	$swa2=0.0$	V m	Coefficient of the width dependence of $a2$.
73	$a3r=1.0$		Factor of the drain-source voltage above which weak-avalanche occurs.
74	$sla3=0.0$	m	Coefficient of the length dependence of $a3$.
75	$swa3=0.0$	m	Coefficient of the width dependence of $a3$.

Gate current parameters

76	$iginvr=0.0$	A/V ²	Gain factor for intrinsic gate tunnelling current in inversion.
77	$igaccr=0.0$	A/V ²	Gain factor for intrinsic gate tunnelling current in accumulation.
78	$igovr=0.0$	A/V ²	Gain factor for Source/Drain overlap gate tunnelling current.

Charge parameters

79	$col=3.2e-10$	F/m	Gate overlap capacitance per unit channel width.
----	---------------	-----	--

Noise parameters

80	$nfar=1.573e23$	1/(V ^m ⁴)	First coefficient of the flicker noise.
81	$nfbr=4.752e9$	1/(V ^m ²)	Second coefficient of the flicker noise.
82	$nfcr=0.0$	1/V	Third coefficient of the flicker noise.

Temperature parameters

83	tr	(C)	Reference temperature. Default set by option t_{nom} .
84	$tref$	(C)	Alias of tr . Default set by option t_{nom} .

Spectre Circuit Simulator Reference

Component Statements Part II

85 `tnom` (C) Alias of `tr`. Default set by option.

Other parameters

86 `kpinv`=0.0 $1/\sqrt{V}$ Inverse of body-effect factor of the poly-silicon gate.

87 `binv`=48.0 V Probability factor for intrinsic gate tunnelling current in inversion.

88 `bacc`=48.0 V Probability factor for intrinsic gate tunnelling current in accumulation.

89 `vfbov`=0.0 V Flat-band voltage for the Source/Drain overlap extensions.

90 `kov`=2.5 \sqrt{V} Body-effect factor for the Source/Drain overlap extensions.

91 `gatenoise`=0.0 Flag for in/exclusion of induced gate thermal noise.

Output Parameters

1 `vto` (V) Zero-bias threshold voltage.

2 `le` (m) Effective channel length.

3 `we` (m) Effective channel width.

4 `ko` (\sqrt{V}) Body-effect factor.

5 `phib` (V) Surface potential at the onset of strong inversion.

6 `bet` (A/V^2) Gain factor.

7 `thesr` ($1/V$) Mobility degradation parameter due to surface roughness scattering.

8 `theph` ($1/V$) Mobility degradation parameter due to phonon scattering.

9 `etamob` Effective field parameter for dependence on depletion charge.

10 `nu` Exponent of field dependence of mobility model.

11 `ther` ($1/V$) Coefficient of series resistance.

Spectre Circuit Simulator Reference

Component Statements Part II

12	thesat (1/V)	Velocity saturation parameter due to optical/acoustic phonon scattering.
13	theth (1/V ³)	Coefficient of self-heating.
14	sdibl (1/√V)	Drain-induced barrier lowering parameter.
15	mo	Parameter for (short-channel) subthreshold slope.
16	ssf (1/√V)	Static-feedback parameter.
17	alp	Factor of channel length modulation.
18	mexp	Smoothing factor.
19	a1	Factor of the weak-avalanche current.
20	a2 (V)	Exponent of the weak-avalanche current.
21	a3	Factor of the drain-source voltage above which weak-avalanche occurs.
22	iginv (A/V ²)	Gain factor for intrinsic gate tunnelling current in inversion.
23	igacc (A/V ²)	Gain factor for intrinsic gate tunnelling current in accumulation.
24	igov (A/V ²)	Gain factor for Source/Drain overlap tunnelling current.
25	cox (F)	Oxide capacitance for the intrinsic channel (* mult).
26	cgdo (F)	Oxide capacitance for the gate-drain overlap (* mult).
27	cgso (F)	Oxide capacitance for the gate-source overlap (* mult).
28	nt (J)	Thermal noise coefficient.
29	nfa (1/(Vm ⁴))	First coefficient of the flicker noise.
30	nfb (1/(Vm ²))	Second coefficient of the flicker noise.
31	nfc (1/V)	Third coefficient of the flicker noise.
32	tox (m)	Thickness of gate oxide layer.

Spectre Circuit Simulator Reference

Component Statements Part II

Operating-Point Parameters

1	<code>ide</code> (A)	Resistive drain current.
2	<code>ige</code> (A)	Resistive gate current.
3	<code>ise</code> (A)	Resistive source current.
4	<code>ibe</code> (A)	Resistive bulk current.
5	<code>isb</code> (A)	Resistive source-bulk current.
6	<code>idb</code> (A)	Resistive drain-bulk current.
7	<code>pwr</code> (W)	Power.
8	<code>ids</code> (A)	Drain current, excl. avalanche and tunnel currents.
9	<code>iavl</code> (A)	Substrate current due to weak-avalanche.
10	<code>igs</code> (A)	Gate-to-source current due to direct tunnelling.
11	<code>igd</code> (A)	Gate-to-drain current due to direct tunnelling.
12	<code>igb</code> (A)	Gate-to-bulk current due to direct tunnelling.
13	<code>vds</code> (V)	Drain-source voltage.
14	<code>vgs</code> (V)	Gate-source voltage.
15	<code>vsb</code> (V)	Source-bulk voltage.
16	<code>vts</code> (V)	Threshold voltage including back-bias effects.
17	<code>vth</code> (V)	Threshold voltage including back-bias and drain-bias effects.
18	<code>vgt</code> (V)	Effective gate drive voltage including back-bias and drain voltage effects.
19	<code>vdss</code> (V)	Drain saturation voltage at actual bias.
20	<code>vsat</code> (V)	Saturation limit.

Spectre Circuit Simulator Reference

Component Statements Part II

21	g_m (S)	Transconductance ($d i_{ds} / d v_{gs}$).
22	g_{mb} (S)	Substrate-transconductance ($d i_{ds} / d v_{bs}$).
23	g_{ds} (S)	Output conductance ($d i_{ds} / d v_{ds}$).
24	c_{dd} (F)	Capacitance ($d q_d / d v_d$).
25	c_{dg} (F)	Capacitance ($- d q_d / d v_g$).
26	c_{ds} (F)	Capacitance ($- d q_d / d v_s$).
27	c_{db} (F)	Capacitance ($- d q_d / d v_b$).
28	c_{gd} (F)	Capacitance ($- d q_g / d v_d$).
29	c_{gg} (F)	Capacitance ($d q_g / d v_g$).
30	c_{gs} (F)	Capacitance ($- d q_g / d v_s$).
31	c_{gb} (F)	Capacitance ($- d q_g / d v_b$).
32	c_{sd} (F)	Capacitance ($- d q_s / d v_d$).
33	c_{sg} (F)	Capacitance ($- d q_s / d v_g$).
34	c_{ss} (F)	Capacitance ($d q_s / d v_s$).
35	c_{sb} (F)	Capacitance ($- d q_s / d v_b$).
36	c_{bd} (F)	Capacitance ($- d q_b / d v_d$).
37	c_{bg} (F)	Capacitance ($- d q_b / d v_g$).
38	c_{bs} (F)	Capacitance ($- d q_b / d v_s$).
39	c_{bb} (F)	Capacitance ($d q_b / d v_b$).
40	u	Transistor gain (g_m/g_{ds}).
41	r_{out} (Ω)	Small-signal output resistance ($1/g_{ds}$).
42	v_{early} (V)	Equivalent Early voltage ($ i_d /g_{ds}$).

Spectre Circuit Simulator Reference

Component Statements Part II

43	k_{eff} (\sqrt{V})	Body effect parameter.
44	b_{eff} (A/V^2)	Gain factor.
45	f_{ug} (Hz)	Unity gain frequency at actual bias ($gm/(2*\pi*cin)$).
46	$sqrtsfw$ (V/\sqrt{Hz})	Input-referred RMS white noise voltage density.
47	$sqrtsff$ (V/\sqrt{Hz})	Input-referred RMS white noise voltage density at 1 kHz.
48	f_{knee} (Hz)	Cross-over frequency above which white noise is dominant.

Parameter Index

In the following index, I refers to instance parameters, M refers to the model parameters section, O refers to the output parameters section, and OP refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of M-35 means the 35th model parameter.

a1	O-19	fknee	OP-48	nfb	O-30	swther	M-44
a1r	M-66	fug	OP-45	nfbr	M-81	swthesat	M-48
a2	O-20	gatenoise	M-91	nfc	O-31	swthesr	M-34
a2r	M-70	gds	OP-23	nfcrr	M-82	theph	O-8
a3	O-21	gm	OP-21	nt	M-13	thephr	M-35
a3r	M-73	gmb	OP-22	nt	O-28	ther	O-11
alp	O-17	iavl	OP-9	nu	M-41	ther1	M-8
alpexp	M-59	ibe	OP-4	nu	O-10	ther2	M-9
alpr	M-56	idb	OP-6	nuexp	M-42	therr	M-43
area	I-4	ide	OP-1	phib	O-5	thesat	O-12
bacc	M-88	ids	OP-8	phibr	M-20	thesatexp	M-50
beff	OP-44	igacc	O-23	pwr	OP-7	thesatr	M-46
bet	O-6	igaccr	M-77	region	I-5	thesr	O-7
betsq	M-29	igb	OP-12	rout	OP-41	thesrr	M-33

Spectre Circuit Simulator Reference Component Statements Part II

binv	M-87	igd	OP-11	sdibl	O-14	theth	O-13
cbb	OP-39	ige	OP-2	sdiblexp	M-61	thethexp	M-52
cbd	OP-36	iginv	O-22	sdiblo	M-60	thethr	M-51
cbg	OP-37	iginvr	M-76	sl2ko	M-18	tnom	M-85
cbs	OP-38	igov	O-24	sl2phib	M-22	tox	M-14
cdb	OP-27	igovr	M-78	sla1	M-67	tox	O-32
cdd	OP-24	igs	OP-10	sla2	M-71	tr	M-83
cdg	OP-25	isb	OP-5	sla3	M-74	tref	M-84
cds	OP-26	ise	OP-3	slalp	M-58	trise	M-12
cgb	OP-31	keff	OP-43	sletabet	M-31	type	M-5
cgd	OP-28	ko	O-4	slko	M-17	u	OP-40
cgdo	O-26	kor	M-16	slphib	M-21	vds	OP-13
cgg	OP-29	kov	M-90	slssf	M-55	vdss	OP-19
cgs	OP-30	kpinv	M-86	slthesat	M-49	vearly	OP-42
cgso	O-27	l	I-2	sqrtsff	OP-47	vfb	M-7
col	M-79	lap	M-2	sqrtsfw	OP-46	vfbov	M-89
cox	O-25	le	O-2	ssf	O-16	vgs	OP-14
csb	OP-35	level	M-6	ssfr	M-53	vgt	OP-18
csd	OP-32	lmin	M-65	sta1	M-69	vp	M-10
csg	OP-33	lp1	M-26	stetamob	M-39	vsat	OP-20
css	OP-34	lp2	M-27	stphib	M-24	vsb	OP-15
dta	M-11	lvar	M-1	stvfb	M-15	vth	OP-17
etabetr	M-30	m	I-6	swa1	M-68	vto	O-1
etamob	O-9	mexp	O-18	swa2	M-72	vts	OP-16
etamobr	M-38	mo	O-15	swa3	M-75	w	I-1
etaph	M-37	moexp	M-63	swalp	M-57	we	O-3
etar	M-45	moo	M-64	swetamob	M-40	wot	M-4
etasat	M-47	mor	M-62	swko	M-19	wvar	M-3
etasr	M-32	mult	I-3	swphib	M-23		
fbet1	M-25	nfa	O-29	swssf	M-54		
fbet2	M-28	nfars	M-80	swtheph	M-36		

Compact MOS-Transistor Distortion Model (mos11011)

Description

The mos1101 model is based on the thesis of Ronald van Langevelde: "A compact MOSFET Model for Distortion Analysis in Analog Circuit Design", Technische Universiteit Eindhoven, 1998.

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In extension to the description a minimum conductance g_{min} is inserted between the drain and source node, to aid convergence. The value of g_{min} is set by an options statement, default = 1e-12 S.

This device is supported within altergroups.

This device is dynamically loaded from the shared object /vobs/spectre_dev/tools.sun4v/spectre/lib/cmi/3.0.doc/libphilips_sh.so

Instance Definition

```
Name d g s [b] ModelName parameter=value ...
```

Instance Parameters

- | | | |
|---|---------------|--|
| 1 | w=1.0 scale m | Drawn channel width in the layout. Scale set by option scale.. |
| 2 | l=1.0 scale m | Drawn channel length in the layout. Scale set by option scale.. |
| 3 | mult=1 | Number of devices in parallel. |
| 4 | area=1 | Alias of mult. |
| 5 | region=triode | Estimated DC operating region, used as a convergence aid.
Possible values are off, triode, sat, or subth. |
| 6 | m=1 | Multiplicity factor. |

Model Definition

```
model modelName mos11011 parameter=value ...
```

Spectre Circuit Simulator Reference

Component Statements Part II

Model Parameters

Geometry parameters

- | | | |
|---|---------------------------|--|
| 1 | <code>lvar=0.0 m</code> | Difference between the actual and the programmed poly-silicon gate length. |
| 2 | <code>lap=4.0e-8 m</code> | Effective channel length reduction per side. |
| 3 | <code>wvar=0.0 m</code> | Difference between the actual and the programmed field-oxide opening. |
| 4 | <code>wot=0.0 m</code> | Effective channel width reduction per side. |

Device type parameters

- | | | |
|----|-----------------------------|--|
| 5 | <code>type=n</code> | Transistor gender.
Possible values are <code>n</code> or <code>p</code> . |
| 6 | <code>level=1.0</code> | Transistor Level. Possible value is 1 (Philips MOSFET Modelbook June 2002). |
| 7 | <code>vfb=-1.050 V</code> | Flat-band voltage at reference temperature. |
| 8 | <code>ther1=0.0 V</code> | Numerator of gate voltage dependent part of series resistance. |
| 9 | <code>ther2=1.0 V</code> | Denominator of gate voltage dependent part of series resistance. |
| 10 | <code>vp=5.0e-2 V</code> | Characteristic voltage of channel-length modulation. |
| 11 | <code>dta=0.0 K</code> | Temperature offset of the device. |
| 12 | <code>trise=0.0 K</code> | Alias of <code>dta</code> . |
| 13 | <code>nt=1.656e-20 J</code> | Coefficient of the thermal noise. |
| 14 | <code>tox=3.2e-09 m</code> | Thickness of the oxide layer. |
| 15 | <code>nu=1.0</code> | Exponent of the field dependence of the mobility model minus 1. |
| 16 | <code>vfbov=0.0 V</code> | Flat-band voltage for the Source/Drain overlap extensions. |

Spectre Circuit Simulator Reference

Component Statements Part II

17	<code>kov=2.5</code>	\sqrt{V}	Body-effect factor for the Source/Drain overlap extensions.
18	<code>gatenoise=0.0</code>		Flag for in/exclusion of induced gate thermal noise.
19	<code>poko=0.5</code>	K	Coefficient for the geometry independent part of k_o .
20	<code>plko=0.0</code>	K	Coefficient for the length independent part of k_o .
21	<code>pwko=0.0</code>	K	Coefficient for the width independent part of k_o .
22	<code>plwko=0.0</code>	K	Coefficient for the length times width independent part of k_o .
23	<code>kpinv=0.0</code>	K	inverse of body-effect factor of the poly-silicon gate.
24	<code>pophib=0.950</code>	K	Coefficient for the geometric independent part of Phib .
25	<code>plphib=0.0</code>	K	Coefficient for the length independent part of Phib .
26	<code>pwphib=0.0</code>	K	Coefficient for the width independent part of Phib .
27	<code>plwphib=0.0</code>	K	Coefficient for the length times width independent part of Phib .
28	<code>pobet=1.922e-3</code>	K	Coefficient for the geometry independent part of bet .
29	<code>plbet=0.0</code>	K	Coefficient for the length independent part of bet .
30	<code>pwbet=0.0</code>	K	Coefficient for the width independent part of bet .
31	<code>plwbet=0.0</code>	K	Coefficient for the width over length independent part of bet .
32	<code>pothesr=3.562e-1</code>	K	Coefficient of the geometry independent part of Thetasr .
33	<code>plthesr=0.0</code>	K	Coefficient of the length independent part of Thetasr .
34	<code>pwthesr=0.0</code>	K	Coefficient of the width independent part of Thetasr .
35	<code>plwthesr=0.0</code>	K	Coefficient of the length times width independent part of Thetasr .
36	<code>potheph=1.290e-2</code>	K	Coefficient of the geometry independent part of Thetaph .
37	<code>pltheph=0.0</code>	K	Coefficient of the length independent part of Thetaph .

Spectre Circuit Simulator Reference

Component Statements Part II

38	<code>pwtheph=0.0 K</code>	Coefficient of the width independent part of Thetaph.
39	<code>plwtheph=0.0 K</code>	Coefficient of the length times width independent part of Thetaph.
40	<code>poetamob=1.400 K</code>	Coefficient of the geometry independent part of etamob.
41	<code>pletamob=0.0 K</code>	Coefficient of the length independent part of etamob.
42	<code>pwetamob=0.0 K</code>	Coefficient of the width independent part of etamob.
43	<code>plwetamob=0.0 K</code>	Coefficient of the length times width independent part of etamob.
44	<code>pother=8.120e-2 K</code>	Coefficient of the geometry independent part of ther.
45	<code>plther=0.0 K</code>	Coefficient of the length independent part of ther.
46	<code>pwther=0.0 K</code>	Coefficient of the width independent part of ther.
47	<code>plwther=0.0 K</code>	Coefficient of the length times width independent part of ther.
48	<code>pothesat=2.513e-1 K</code>	Coefficient for the geometry independent part of thesat.
49	<code>plthesat=0.0 K</code>	Coefficient for the length dependence of thesat.
50	<code>pwthesat=0.0 K</code>	Coefficient for the width dependence of thesat.
51	<code>plwthesat=0.0 K</code>	Coefficient for the length times width dependence of thesat.
52	<code>potheth=1.0e-5 K</code>	Coefficient for the geometry independent part of theth.
53	<code>pltheth=0.0 K</code>	Coefficient for the length dependence of theth.
54	<code>pwtheth=0.0 K</code>	Coefficient for the width dependence of theth.
55	<code>plwtheth=0.0 K</code>	Coefficient for the length times width dependence of theth.
56	<code>posdibl=8.530e-4 K</code>	Coefficient for the geometry independent part of sdibl.
57	<code>plsdibl=0.0 K</code>	Coefficient for the length dependence of sdibl.

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58	<code>pwsdibl=0.0 K</code>	Coefficient for the width dependence of <code>sdibl</code> .
59	<code>plwsdibl=0.0 K</code>	Coefficient for the length times width dependence of <code>sdibl</code> .
60	<code>pomo=0.0 K</code>	Coefficient for the geometry independent part of <code>mo</code> .
61	<code>plmo=0.0 K</code>	Coefficient for the length dependence of <code>mo</code> .
62	<code>pwmomo=0.0 K</code>	Coefficient for the width dependence of <code>mo</code> .
63	<code>plwmo=0.0 K</code>	Coefficient for the length times width dependence of <code>mo</code> .
64	<code>possf=1.200e-2 K</code>	Coefficient for the geometry independent part of <code>ssf</code> .
65	<code>plssf=0.0 K</code>	Coefficient for the length dependence of <code>ssf</code> .
66	<code>pwssf=0.0 K</code>	Coefficient for the width dependence of <code>ssf</code> .
67	<code>plwssf=0.0 K</code>	Coefficient for the length times width dependence of <code>ssf</code> .
68	<code>poalp=2.500e-2 K</code>	Coefficient for the geometry independent part of <code>alp</code> .
69	<code>plalp=0.0 K</code>	Coefficient for the length dependence of <code>alp</code> .
70	<code>pwalp=0.0 K</code>	Coefficient for the width dependence of <code>alp</code> .
71	<code>plwalp=0.0 K</code>	Coefficient for the length times width dependence of <code>alp</code> .
72	<code>pomexp=0.200 K</code>	Coefficient for the geometry independent part of <code>mexp</code> .
73	<code>plmexp=0.0 K</code>	Coefficient for the length dependence of <code>mexp</code> .
74	<code>pwmexp=0.0 K</code>	Coefficient for the width dependence of <code>mexp</code> .
75	<code>plwmexp=0.0 K</code>	Coefficient for the length times width dependence of <code>mexp</code> .
76	<code>poa1=6.022 K</code>	Coefficient for the geometry independent part of <code>a1</code> .
77	<code>pla1=0.0 K</code>	Coefficient for the length dependence of <code>a1</code> .
78	<code>pwa1=0.0 K</code>	Coefficient for the width dependence of <code>a1</code> .
79	<code>plwa1=0.0 K</code>	Coefficient for the length times width dependence of <code>a1</code> .

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80	<code>poa2=3.802e+1 K</code>	Coefficient for the geometry independent part of a2.
81	<code>pla2=0.0 K</code>	Coefficient for the length dependence of a2.
82	<code>pwa2=0.0 K</code>	Coefficient for the width dependence of a2.
83	<code>plwa2=0.0 K</code>	Coefficient for the length times width dependence of a2.
84	<code>poa3=6.407e-1 K</code>	Coefficient for the geometry independent part of a3.
85	<code>pla3=0.0 K</code>	Coefficient for the length dependence of a3.
86	<code>pwa3=0.0 K</code>	Coefficient for the width dependence of a3.
87	<code>plwa3=0.0 K</code>	Coefficient for the length times width dependence of a3.
88	<code>poiginv=0.0 K</code>	Coefficient for the geometry independent part of iginv.
89	<code>pliginv=0.0 K</code>	Coefficient for the length dependence of iginv.
90	<code>pwiginv=0.0 K</code>	Coefficient for the width dependence of iginv.
91	<code>plwiginv=0.0 K</code>	Coefficient for the length times width dependence of iginv.
92	<code>pobinv=4.800e+1 K</code>	Coefficient for the geometry independent part of binv.
93	<code>plbinv=0.0 K</code>	Coefficient for the length dependence of binv.
94	<code>pwbinv=0.0 K</code>	Coefficient for the width dependence of binv.
95	<code>plwbinv=0.0 K</code>	Coefficient for the length times width dependence of binv.
96	<code>poigacc=0.0 K</code>	Coefficient for the geometry independent part of igacc.
97	<code>pligacc=0.0 K</code>	Coefficient for the length dependence of igacc.
98	<code>pwigacc=0.0 K</code>	Coefficient for the width dependence of igacc.
99	<code>plwigacc=0.0 K</code>	Coefficient for the length times width dependence of igacc.
100	<code>pobacc=4.800e+1 K</code>	Coefficient for the geometry independent part of bacc.
101	<code>plbacc=0.0 K</code>	Coefficient for the length dependence of bacc.

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Component Statements Part II

102	<code>pwbacc=0.0 K</code>	Coefficient for the width dependence of bacc.
103	<code>plwbacc=0.0 K</code>	Coefficient for the length times width dependence of bacc.
104	<code>poigov=0.0 K</code>	Coefficient for the geometry independent part of igov.
105	<code>pligov=0.0 K</code>	Coefficient for the length dependence of igov.
106	<code>pwigov=0.0 K</code>	Coefficient for the width dependence of igov.
107	<code>plwigov=0.0 K</code>	Coefficient for the length times width dependence of igov.
108	<code>pocox=2.980e-14 K</code>	Coefficient for the geometry independent part of cox.
109	<code>plcox=0.0 K</code>	Coefficient for the length dependence of cox.
110	<code>pwc Cox=0.0 K</code>	Coefficient for the width dependence of cox.
111	<code>plwcox=0.0 K</code>	Coefficient for the length times width dependence of cox.
112	<code>pocgdo=6.392e-15 K</code>	Coefficient for the geometry independent part of cgdo.
113	<code>plcgdo=0.0 K</code>	Coefficient for the length dependence of cgdo.
114	<code>pwcgdo=0.0 K</code>	Coefficient for the width dependence of cgdo.
115	<code>plwcgdo=0.0 K</code>	Coefficient for the width over length dependence of cgdo.
116	<code>pocgso=6.392e-15 K</code>	Coefficient for the geometry independent part of cgso.
117	<code>plcgso=0.0 K</code>	Coefficient for the length dependence of cgso.
118	<code>pwcgso=0.0 K</code>	Coefficient for the width dependence of cgso.
119	<code>plwcgso=0.0 K</code>	Coefficient for the width over length dependence of cgso.
120	<code>ponfa=8.323e+22 K</code>	Coefficient for the geometry independent part of nfa.
121	<code>plnfa=0.0 K</code>	Coefficient for the length dependence of nfa.
122	<code>pwnfa=0.0 K</code>	Coefficient for the width dependence of nfa.

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Component Statements Part II

123	<code>plwnfa=0.0 K</code>	Coefficient for the length times width dependence of nfa.
124	<code>ponfb=2.514e+7 K</code>	Coefficient for the geometry independent part of nfb.
125	<code>plnfb=0.0 K</code>	Coefficient for the length dependence of nfb.
126	<code>pwnfb=0.0 K</code>	Coefficient for the width dependence of nfb.
127	<code>plwnfb=0.0 K</code>	Coefficient for the length times width dependence of nfb.
128	<code>ponfc=0.0 K</code>	Coefficient for the geometry independent part of nfc.
129	<code>plnfc=0.0 K</code>	Coefficient for the length dependence of nfc.
130	<code>pwnfc=0.0 K</code>	Coefficient for the width dependence of nfc.
131	<code>plwnfc=0.0 K</code>	Coefficient for the length times width dependence of nfc.
132	<code>potvfb=5.000e-4 K</code>	Coefficient for the geometry independent part of tvfb.
133	<code>pltvfb=0.0 K</code>	Coefficient for the length dependence of tvfb.
134	<code>pwtvfb=0.0 K</code>	Coefficient for the width dependence of tvfb.
135	<code>plwtvfb=0.0 K</code>	Coefficient for the length times width dependence of tvfb.
136	<code>potphib=-8.500e-4 K</code>	Coefficient for the geometry independent part of tphib.
137	<code>pltphib=0.0 K</code>	Coefficient for the length dependence of tphib.
138	<code>pwtphib=0.0 K</code>	Coefficient for the width dependence of tphib.
139	<code>plwtphib=0.0 K</code>	Coefficient for the length times width dependence of tphib.
140	<code>potetabet=1.300 K</code>	Coefficient for the geometry independent part of tetabet.
141	<code>pltetabet=0.0 K</code>	Coefficient for the length dependence of tetabet.
142	<code>pwtetabet=0.0 K</code>	Coefficient for the width dependence of tetabet.
143	<code>plwtetabet=0.0 K</code>	Coefficient for the length times width dependence of tetabet.

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Component Statements Part II

- 144 potetasr=0.650 K Coefficient for the geometry independent part of tetasr.
- 145 pltetasr=0.0 K Coefficient for the length dependence of tetasr.
- 146 pwtetasr=0.0 K Coefficient for the width dependence of tetasr.
- 147 plwtetasr=0.0 K Coefficient for the length times width dependence of tetasr.
- 148 potetaph=1.350 K Coefficient for the geometry independent part of tetaph.
- 149 pltetaph=0.0 K Coefficient for the length dependence of tetaph.
- 150 pwtetaph=0.0 K Coefficient for the width dependence of tetaph.
- 151 plwtetaph=0.0 K Coefficient for the length times width dependence of tetaph.
- 152 potetamob=0.0 K Coefficient for the geometry independent part of tetamob.
- 153 pltetamob=0.0 K Coefficient for the length dependence of tetamob.
- 154 pwtetamob=0.0 K Coefficient for the width dependence of tetamob.
- 155 plwtetamob=0.0 K Coefficient for the length times width dependence of tetamob.
- 156 potnuexp=5.250 K Coefficient for the geometry independent part of tnuexp.
- 157 pltnuexp=0.0 K Coefficient for the length dependence of tnuexp.
- 158 pwtnuexp=0.0 K Coefficient for the width dependence of tnuexp.
- 159 plwtnuexp=0.0 K Coefficient for the length times width dependence of tnuexp.
- 160 potetar=0.950 K Coefficient for the geometry independent part of tetar.
- 161 pltetar=0.0 K Coefficient for the length dependence of tetar.
- 162 pwtetar=0.0 K Coefficient for the width dependence of tetar.
- 163 plwtetar=0.0 K Coefficient for the length times width dependence of tetar.
- 164 potetasat=1.040 K Coefficient for the geometry independent part of tetasat.
- 165 pltetasat=0.0 K Coefficient for the length dependence of tetasat.

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Component Statements Part II

166	<code>pwtetasat=0.0 K</code>	Coefficient for the width dependence of <code>tetasat</code> .
167	<code>plwtetasat=0.0 K</code>	Coefficient for the length times width dependence of <code>tetasat</code> .
168	<code>pota1=0.0 K</code>	Coefficient for the geometry independent part of <code>ta1</code> .
169	<code>plta1=0.0 K</code>	Coefficient for the length dependence of <code>ta1</code> .
170	<code>pwtal=0.0 K</code>	Coefficient for the width dependence of <code>ta1</code> .
171	<code>plwta1=0.0 K</code>	Coefficient for the length times width dependence of <code>ta1</code> .
172	<code>tr (C)</code>	Reference temperature. Default set by option <code>tnom</code> .
173	<code>tref (C)</code>	Alias of <code>tr</code> . Default set by option <code>tnom</code> .
174	<code>tnom (C)</code>	Alias of <code>tr</code> . Default set by option.

Output Parameters

1	<code>vto (V)</code>	Zero-bias threshold voltage.
2	<code>le (m)</code>	Effective channel length.
3	<code>we (m)</code>	Effective channel width.
4	<code>ko (\sqrt{V})</code>	Body-effect factor.
5	<code>phib (V)</code>	Surface potential at the onset of strong inversion.
6	<code>bet (A/V^2)</code>	Gain factor.
7	<code>thesr ($1/V$)</code>	Mobility degradation parameter due to surface roughness scattering.
8	<code>theph ($1/V$)</code>	Mobility degradation parameter due to phonon scattering.
9	<code>etamob</code>	Effective field parameter for dependence on depletion charge.
10	<code>nu</code>	Exponent of field dependence of mobility model.
11	<code>ther ($1/V$)</code>	Coefficient of series resistance.

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Component Statements Part II

12	thesat (1/V)	Velocity saturation parameter due to optical/acoustic phonon scattering.
13	theth (1/V ³)	Coefficient of self-heating.
14	sdibl (1/ \sqrt{V})	Drain-induced barrier lowering parameter.
15	mo	Parameter for (short-channel) subthreshold slope.
16	ssf (1/ \sqrt{V})	Static-feedback parameter.
17	alp	Factor of channel length modulation.
18	mexp	Smoothing factor.
19	a1	Factor of the weak-avalanche current.
20	a2 (V)	Exponent of the weak-avalanche current.
21	a3	Factor of the drain-source voltage above which weak-avalanche occurs.
22	iginv (A/V ²)	Gain factor for intrinsic gate tunnelling current in inversion.
23	igacc (A/V ²)	Gain factor for intrinsic gate tunnelling current in accumulation.
24	igov (A/V ²)	Gain factor for Source/Drain overlap tunnelling current.
25	cox (F)	Oxide capacitance for the intrinsic channel (* mult).
26	cgdo (F)	Oxide capacitance for the gate-drain overlap (* mult).
27	cgso (F)	Oxide capacitance for the gate-source overlap (* mult).
28	nt (J)	Thermal noise coefficient.
29	nfa (1/(Vm ⁴))	First coefficient of the flicker noise.
30	nfb (1/(Vm ²))	Second coefficient of the flicker noise.
31	nfc (1/V)	Third coefficient of the flicker noise.
32	tox (m)	Thickness of gate oxide layer.

Spectre Circuit Simulator Reference

Component Statements Part II

Operating-Point Parameters

1	<code>ide</code> (A)	Resistive drain current.
2	<code>ige</code> (A)	Resistive gate current.
3	<code>ise</code> (A)	Resistive source current.
4	<code>ibe</code> (A)	Resistive bulk current.
5	<code>isb</code> (A)	Resistive source-bulk current.
6	<code>idb</code> (A)	Resistive drain-bulk current.
7	<code>pwr</code> (W)	Power.
8	<code>ids</code> (A)	Drain current, excl. avalanche and tunnel currents.
9	<code>iavl</code> (A)	Substrate current due to weak-avalanche.
10	<code>igs</code> (A)	Gate-to-source current due to direct tunnelling.
11	<code>igd</code> (A)	Gate-to-drain current due to direct tunnelling.
12	<code>igb</code> (A)	Gate-to-bulk current due to direct tunnelling.
13	<code>vds</code> (V)	Drain-source voltage.
14	<code>vgs</code> (V)	Gate-source voltage.
15	<code>vsb</code> (V)	Source-bulk voltage.
16	<code>vts</code> (V)	Threshold voltage including back-bias effects.
17	<code>vth</code> (V)	Threshold voltage including back-bias and drain-bias effects.
18	<code>vgt</code> (V)	Effective gate drive voltage including back-bias and drain voltage effects.
19	<code>vdss</code> (V)	Drain saturation voltage at actual bias.
20	<code>vsat</code> (V)	Saturation limit.

Spectre Circuit Simulator Reference

Component Statements Part II

21	g_m (S)	Transconductance ($d i_{ds} / d v_{gs}$).
22	g_{mb} (S)	Substrate-transconductance ($d i_{ds} / d v_{bs}$).
23	g_{ds} (S)	Output conductance ($d i_{ds} / d v_{ds}$).
24	c_{dd} (F)	Capacitance ($d q_d / d v_d$).
25	c_{dg} (F)	Capacitance ($- d q_d / d v_g$).
26	c_{ds} (F)	Capacitance ($- d q_d / d v_s$).
27	c_{db} (F)	Capacitance ($- d q_d / d v_b$).
28	c_{gd} (F)	Capacitance ($- d q_g / d v_d$).
29	c_{gg} (F)	Capacitance ($d q_g / d v_g$).
30	c_{gs} (F)	Capacitance ($- d q_g / d v_s$).
31	c_{gb} (F)	Capacitance ($- d q_g / d v_b$).
32	c_{sd} (F)	Capacitance ($- d q_s / d v_d$).
33	c_{sg} (F)	Capacitance ($- d q_s / d v_g$).
34	c_{ss} (F)	Capacitance ($d q_s / d v_s$).
35	c_{sb} (F)	Capacitance ($- d q_s / d v_b$).
36	c_{bd} (F)	Capacitance ($- d q_b / d v_d$).
37	c_{bg} (F)	Capacitance ($- d q_b / d v_g$).
38	c_{bs} (F)	Capacitance ($- d q_b / d v_s$).
39	c_{bb} (F)	Capacitance ($d q_b / d v_b$).
40	u	Transistor gain (g_m/g_{ds}).
41	r_{out} (Ω)	Small-signal output resistance ($1/g_{ds}$).
42	v_{early} (V)	Equivalent Early voltage ($ i_d /g_{ds}$).

Spectre Circuit Simulator Reference

Component Statements Part II

43	k_{eff} (\sqrt{V})	Body effect parameter.
44	b_{eff} (A/V^2)	Gain factor.
45	f_{ug} (Hz)	Unity gain frequency at actual bias ($gm/(2*\pi*cin)$).
46	$sqrtsfw$ (V/\sqrt{Hz})	Input-referred RMS white noise voltage density.
47	$sqrtsff$ (V/\sqrt{Hz})	Input-referred RMS white noise voltage density at 1 kHz.
48	f_{knee} (Hz)	Cross-over frequency above which white noise is dominant.

Parameter Index

In the following index, I refers to instance parameters, M refers to the model parameters section, O refers to the output parameters section, and OP refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of M-35 means the 35th model parameter.

a1	O-19	nt	O-28	plwta1	M-171	pwiginv	M-90
a2	O-20	nu	M-15	plwtetabet	M-143	pwigov	M-106
a3	O-21	nu	O-10	plwtetamob	M-155	pwko	M-21
alp	O-17	phib	O-5	plwtetaph	M-151	pwmexp	M-74
area	I-4	pla1	M-77	plwtetar	M-163	pwm0	M-62
beff	OP-44	pla2	M-81	plwtetasat	M-167	pwnfa	M-122
bet	O-6	pla3	M-85	plwtetasr	M-147	pwnfb	M-126
cbb	OP-39	plalp	M-69	plwtheph	M-39	pwnfc	M-130
cbd	OP-36	plbacc	M-101	plwther	M-47	pwphib	M-26
cbg	OP-37	plbet	M-29	plwthesat	M-51	pwr	OP-7
cbs	OP-38	plbinv	M-93	plwthesr	M-35	pwsdibl	M-58
cdb	OP-27	plcgdo	M-113	plwtheth	M-55	pwssf	M-66
cdd	OP-24	plcgso	M-117	plwtnuexp	M-159	pwtal	M-170
cdg	OP-25	plcox	M-109	plwtphib	M-139	pwtetabet	M-142

Spectre Circuit Simulator Reference Component Statements Part II

cds	OP-26	pletamob	M-41	plwtvfb	M-135	pwtetamob	M-154
cgb	OP-31	pligacc	M-97	poa1	M-76	pwtetaph	M-150
cgd	OP-28	pliginv	M-89	poa2	M-80	pwtetar	M-162
cgdo	O-26	pligov	M-105	poa3	M-84	pwtetasat	M-166
cgg	OP-29	plko	M-20	poalp	M-68	pwtetasr	M-146
cgs	OP-30	plmexp	M-73	pobacc	M-100	pwtheph	M-38
cgso	O-27	plmo	M-61	pobet	M-28	pwther	M-46
cox	O-25	plnfa	M-121	pobinv	M-92	pwthesat	M-50
csb	OP-35	plnfb	M-125	pocgdo	M-112	pwthesr	M-34
csd	OP-32	plnfc	M-129	pocgso	M-116	pwtheth	M-54
csq	OP-33	plphib	M-25	pocox	M-108	pwtnuexp	M-158
css	OP-34	plsdibl	M-57	poetamob	M-40	pwtphib	M-138
dta	M-11	plssf	M-65	poigacc	M-96	pwtvfb	M-134
etamob	O-9	pltal	M-169	poiginv	M-88	region	I-5
fknee	OP-48	pltetabet	M-141	poigov	M-104	rout	OP-41
fug	OP-45	pltetamob	M-153	poko	M-19	sdibl	O-14
gatenoise	M-18	pltetaph	M-149	pomexp	M-72	sqrtsff	OP-47
gds	OP-23	pltetar	M-161	pomo	M-60	sqrtsfw	OP-46
gm	OP-21	pltetasat	M-165	ponfa	M-120	ssf	O-16
gmb	OP-22	pltetasr	M-145	ponfb	M-124	theph	O-8
iavl	OP-9	pltheph	M-37	ponfc	M-128	ther	O-11
ibe	OP-4	plther	M-45	pophib	M-24	ther1	M-8
idb	OP-6	plthesat	M-49	posdibl	M-56	ther2	M-9
ide	OP-1	plthesr	M-33	possf	M-64	thesat	O-12
ids	OP-8	pltheth	M-53	potal	M-168	thesr	O-7
igacc	O-23	pltnuexp	M-157	potetabet	M-140	theth	O-13
igb	OP-12	pltphib	M-137	potetamob	M-152	tnom	M-174
igd	OP-11	pltvfb	M-133	potetaph	M-148	tox	O-32
ige	OP-2	plwal	M-79	potetar	M-160	tox	M-14
iginv	O-22	plwa2	M-83	potetasat	M-164	tr	M-172
igov	O-24	plwa3	M-87	potetasr	M-144	tref	M-173
igs	OP-10	plwalp	M-71	potheph	M-36	trise	M-12

Spectre Circuit Simulator Reference Component Statements Part II

isb	OP-5	plwbacc	M-103	pother	M-44	type	M-5
ise	OP-3	plwbet	M-31	pothesat	M-48	u	OP-40
keff	OP-43	plwbinv	M-95	pothesr	M-32	vds	OP-13
ko	O-4	plwcgdo	M-115	potheth	M-52	vdss	OP-19
kov	M-17	plwcgso	M-119	potnuexp	M-156	vearly	OP-42
kpinv	M-23	plwcox	M-111	potphib	M-136	vfb	M-7
l	I-2	plwetamob	M-43	potvfb	M-132	vfbv	M-16
lap	M-2	plwigacc	M-99	pwa1	M-78	vgs	OP-14
le	O-2	plwiginv	M-91	pwa2	M-82	vgt	OP-18
level	M-6	plwigov	M-107	pwa3	M-86	vp	M-10
lvar	M-1	plwko	M-22	pwalp	M-70	vsat	OP-20
m	I-6	plwmexp	M-75	pwbacc	M-102	vsb	OP-15
mexp	O-18	plwmo	M-63	pwbet	M-30	vth	OP-17
mo	O-15	plwnfa	M-123	pwbinv	M-94	vto	O-1
mult	I-3	plwnfb	M-127	pwcgdo	M-114	vts	OP-16
nfa	O-29	plwnfc	M-131	pwcgso	M-118	w	I-1
nfb	O-30	plwphib	M-27	pwcox	M-110	we	O-3
nfc	O-31	plwsdibl	M-59	pwetamob	M-42	wot	M-4
nt	M-13	plwssf	M-67	pwigacc	M-98	wvar	M-3

MOS Level-15 Transistor (mos15)

Description

The MOS15 model is the AMS level 15 model which is the modified Berkeley SPICE level-2 model with the DC model replaced by that of AMS. It is an analytical one-dimensional model that incorporates most of the second-order small-size effects. A smoother version of the level-15 model (with continuous Gds at Vdsat) was also developed. Three charge models are available. MOS15 transistors require the use of a model statement.

This device is not supported within altergroup.

This device is dynamically loaded from the shared object /vobs/spectre_dev/tools.sun4v/spectre/lib/cmi/3.0.doc/libstmodels_sh.so

Spectre Circuit Simulator Reference

Component Statements Part II

Instance Definition

Name d g s b ModelName parameter=value ...

Instance Parameters

1	w (m)	Channel width.
2	l (m)	Channel length.
3	as (m ²)	Area of source diffusion.
4	ad (m ²)	Area of drain diffusion.
5	ps (m)	Perimeter of source diffusion.
6	pd (m)	Perimeter of drain diffusion.
7	nrd (m/m)	Number of squares of drain diffusion.
8	nrs (m/m)	Number of squares of source diffusion.
9	ld (m)	Length of drain diffusion region.
10	ls (m)	Length of source diffusion region.
11	m=1	Multiplicity factor (number of MOSFETs in parallel).
12	region=triode	Estimated operating region. Spectre outputs number (0-3) in a rawfile. Possible values are off, triode, sat, or subth.
13	trise	Temperature rise from ambient.

Model Definition

model modelName mos15 parameter=value ...

Spectre Circuit Simulator Reference

Component Statements Part II

Model Parameters

Device type parameters

1 `type=n` Transistor type.
Possible values are n or p.

Drain current model parameters

2 `vto=0 V` Threshold voltage at zero body bias.

3 `kp=2.0718e-5 A/V2` Transconductance parameter.

4 `lambda=0 1/V` Channel length modulation parameter.

5 `phi=0.7 V` Surface potential at strong inversion.

6 `gamma=0 \sqrt{V}` Body-effect parameter.

7 `uo=600 cm2/V s` Carrier surface mobility.

8 `vmax= ∞ m/s` Carrier saturation velocity.

9 `ucrit=0 V/cm` Critical field for mobility degradation.

10 `uexp=0` Critical field exponent for mobility degradation.

11 `utra=0 1/V` Transverse field for mobility.

12 `neff=1` Total channel charge coefficient.

13 `delta=0` Width effect on threshold voltage.

Process parameters

14 `nsub=1.13e16 cm-3` Channel doping concentration.

15 `nss=0 cm-2` Surface state density.

16 `nfs=0 cm-2` Fast surface state density.

Spectre Circuit Simulator Reference

Component Statements Part II

17	$t_{pg}=+1$	Type of gate (+1 = opposite of substate, -1 = same as substate, 0 = aluminum).
18	$t_{ox}=1e-7$ m	Gate oxide thickness.
19	$l_d=0$ m	Lateral diffusion.
20	$w_d=0$ m	Field-oxide encroachment.
21	$x_w=0$ m	Width variation due to masking and etching.
22	$x_l=0$ m	Length variation due to masking and etching.
23	$x_j=0$ m	Source/drain junction depth.

Impact ionization parameters

24	$a_{i0}=0$ 1/V	Impact ionization current coefficient.
25	$l_{a_{i0}}=0$ $\mu\text{m}/\text{V}$	Length sensitivity of a_{i0} .
26	$w_{a_{i0}}=0$ $\mu\text{m}/\text{V}$	Width sensitivity of a_{i0} .
27	$b_{i0}=0$ V	Impact ionization current exponent.
28	$l_{b_{i0}}=0$ μm V	Length sensitivity of b_{i0} .
29	$w_{b_{i0}}=0$ μm V	Width sensitivity of b_{i0} .

Overlap capacitance parameters

30	$c_{gso}=0$ F/m	Gate-source overlap capacitance.
31	$c_{gdo}=0$ F/m	Gate-drain overlap capacitance.
32	$c_{gbo}=0$ F/m	Gate-bulk overlap capacitance.
33	$meto=0$ m	Metal overlap in fringing field.

Spectre Circuit Simulator Reference

Component Statements Part II

Charge model selection parameters

- 34 `capmod=bsim` Intrinsic charge model.
Possible values are `none`, `meyer`, `yang`, or `bsim`.
- 35 `xpart=1` Drain/source channel charge partition in saturation for BSIM charge model, use 0.0 for 40/60, 0.5 for 50/50, or 1.0 for 0/100.
- 36 `xqc=0` Drain/source channel charge partition in saturation for charge models, e.g. use 0.4 for 40/60, 0.5 for 50/50, 0 for 0/100.

Parasitic resistance parameters

- 37 `rs=0 Ω` Source resistance.
- 38 `rd=0 Ω` Drain resistance.
- 39 `rsh=0 Ω/sqr` Source/drain diffusion sheet resistance.
- 40 `rss=0 Ω m` Scalable source resistance.
- 41 `rdd=0 Ω m` Scalable drain resistance.
- 42 `rsc=0 Ω` Source contact resistance.
- 43 `rdc=0 Ω` Drain contact resistance.
- 44 `minr=0.1 Ω` Minimum source/drain resistance.
- 45 `ldif=0 m` Lateral diffusion beyond the gate.
- 46 `hdif=0 m` Length of heavily doped diffusion.
- 47 `lgcs=0 m` Gate-to-contact length of source side.
- 48 `lgcd=0 m` Gate-to-contact length of drain side.
- 49 `sc=∞ m` Spacing between contacts.

Junction diode model parameters

- 50 `js (A/m2)` Bulk junction reverse saturation current density.

Spectre Circuit Simulator Reference

Component Statements Part II

51	<code>is=1e-14 A</code>	Bulk junction reverse saturation current.
52	<code>n=1</code>	Junction emission coefficient.
53	<code>dskip=yes</code>	Use simple piece-wise linear model for diode currents below $0.1 \cdot i_{abstol}$. Possible values are <code>no</code> or <code>yes</code> .
54	<code>imax=1 A</code>	Explosion current.
55	<code>jmax=1e8 A/m²</code>	Explosion current density.

Junction capacitance model parameters

56	<code>cbs=0 F</code>	Bulk-source zero-bias junction capacitance.
57	<code>cbd=0 F</code>	Bulk-drain zero-bias junction capacitance.
58	<code>cj=0 F/m²</code>	Zero-bias junction bottom capacitance density.
59	<code>mj=1/2</code>	Bulk junction bottom grading coefficient.
60	<code>pb=0.8 V</code>	Bulk junction built-in potential.
61	<code>fc=0.5</code>	Forward-bias depletion capacitance threshold.
62	<code>cjsw=0 F/m</code>	Zero-bias junction sidewall capacitance density.
63	<code>mjsw=1/3</code>	Bulk junction sidewall grading coefficient.
64	<code>pbsw=0.8 V</code>	Side-wall junction built-in potential.
65	<code>fcsw=0.5</code>	Side-wall forward-bias depletion capacitance threshold.

Operating region warning control parameters

66	<code>alarm=none</code>	Forbidden operating region. Possible values are <code>none</code> , <code>off</code> , <code>triode</code> , <code>sat</code> , <code>subth</code> , or <code>rev</code> .
67	<code>bvj=∞ V</code>	Junction reverse breakdown voltage.

Spectre Circuit Simulator Reference

Component Statements Part II

Temperature effects parameters

68	<code>tnom (C)</code>	Parameters measurement temperature. Default set by options.
69	<code>trise=0 C</code>	Temperature rise from ambient.
70	<code>uto=0 C</code>	Mobility temperature offset.
71	<code>ute=-1.5</code>	Mobility temperature exponent.
72	<code>tlev=0</code>	DC temperature selector.
73	<code>tlevc=0</code>	AC temperature selector.
74	<code>eg=1.12452 V</code>	Energy band gap.
75	<code>gap1=7.02e-4 V/C</code>	Band gap temperature coefficient.
76	<code>gap2=1108 C</code>	Band gap temperature offset.
77	<code>flex=0</code>	Temperature exponent for <code>ucrit</code> .
78	<code>lamex=0 1/C</code>	Temperature parameter for <code>lambda</code> and <code>kappa</code> .
79	<code>trs=0 1/C</code>	Temperature parameter for source resistance.
80	<code>trd=0 1/C</code>	Temperature parameter for drain resistance.
81	<code>x_{ti}=3</code>	Saturation current temperature exponent.
82	<code>ptc=0 V/C</code>	Surface potential temperature coefficient.
83	<code>tcv=0 V/C</code>	Threshold voltage temperature coefficient.
84	<code>pta=0 V/C</code>	Junction potential temperature coefficient.
85	<code>ptp=0 V/C</code>	Sidewall junction potential temperature coefficient.
86	<code>cta=0 1/C</code>	Junction capacitance temperature coefficient.
87	<code>ctp=0 1/C</code>	Sidewall junction capacitance temperature coefficient.

Spectre Circuit Simulator Reference

Component Statements Part II

Default instance parameters

88	$w=3e-6$ m	Default channel width.
89	$l=3e-6$ m	Default channel length.
90	$as=0$ m ²	Default area of source diffusion.
91	$ad=0$ m ²	Default area of drain diffusion.
92	$ps=0$ m	Default perimeter of source diffusion.
93	$pd=0$ m	Default perimeter of drain diffusion.
94	$nrd=0$ m/m	Default number of squares of drain diffusion.
95	$nrs=0$ m/m	Default number of squares of source diffusion.
96	$ldd=0$ m	Default length of drain diffusion region.
97	$lds=0$ m	Default length of source diffusion region.

Noise model parameters

98	$kf=0$	Flicker (1/f) noise coefficient.
99	$af=1$	Flicker (1/f) noise exponent.
100	$ef=1$	Flicker (1/f) noise frequency exponent.
101	$noisemod=1$	Noise model selector.

The i_{max} (j_{max}) parameter is used to aid convergence and prevent numerical overflow. The junction characteristics of the FET are accurately modeled for current (density) up to i_{max} (j_{max}). For currents (density) above i_{max} (j_{max}), the junction is modeled as a linear resistor and a warning is printed.

Output Parameters

1	w_{eff} (m)	Effective channel width.
2	l_{eff} (m)	Effective channel length.

Spectre Circuit Simulator Reference

Component Statements Part II

3 `rseff` (Ω) Effective source resistance.

4 `rdeff` (Ω) Effective drain resistance.

Operating-Point Parameters

1 `type=n` Transistor type.
Possible values are `n` or `p`.

2 `region=triode` Estimated operating region. Spectre outputs number (0-3) in a rawfile.
Possible values are `off`, `triode`, `sat`, or `subth`.

3 `reversed` Reverse mode indicator.
Possible values are `no` or `yes`.

4 `id` (A) Resistive drain current.

5 `vgs` (V) Gate-source voltage.

6 `vds` (V) Drain-source voltage.

7 `vbs` (V) Bulk-source voltage.

8 `vth` (V) Threshold voltage.

9 `vdsat` (V) Drain-source saturation voltage.

10 `gm` (S) Common-source transconductance.

11 `gds` (S) Common-source output conductance.

12 `gmbs` (S) Body-transconductance.

13 `gameff` (\sqrt{V}) Effective body effect coefficient.

14 `betaeff` (A/V^2) Effective beta.

15 `cbd` (F) Drain-bulk junction capacitance.

16 `cbs` (F) Source-bulk junction capacitance.

17 `cgs` (F) Gate-source capacitance.

Spectre Circuit Simulator Reference

Component Statements Part II

18	cgd (F)	Gate-drain capacitance.
19	cgb (F)	Gate-bulk capacitance.
20	ron (Ω)	On-resistance.
21	ib (A)	Resistive bulk current.
22	pwr (W)	Power at op point.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

ad	M-91	gap2	M-76	nfs	M-16	tlevc	M-73
ad	I-4	gds	OP-11	noisemod	M-101	tnom	M-68
af	M-99	gm	OP-10	nrd	M-94	tox	M-18
ai0	M-24	gmbs	OP-12	nrd	I-7	tpg	M-17
alarm	M-66	hdif	M-46	nrs	M-95	trd	M-80
as	I-3	ib	OP-21	nrs	I-8	trise	I-13
as	M-90	id	OP-4	nss	M-15	trise	M-69
betaeff	OP-14	imax	M-54	nsub	M-14	trs	M-79
bi0	M-27	is	M-51	pb	M-60	type	M-1
bvj	M-67	jmax	M-55	pbsw	M-64	type	OP-1
capmod	M-34	js	M-50	pd	I-6	ucrit	M-9
cbd	M-57	kf	M-98	pd	M-93	uexp	M-10
cbd	OP-15	kp	M-3	phi	M-5	uo	M-7
cbs	OP-16	l	I-2	ps	I-5	ute	M-71
cbs	M-56	l	M-89	ps	M-92	uto	M-70
cgb	OP-19	lai0	M-25	pta	M-84	utra	M-11
cgbo	M-32	lambda	M-4	ptc	M-82	vbs	OP-7

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cgd	OP-18	lamex	M-78	ptp	M-85	vds	OP-6
cgdo	M-31	lbi0	M-28	pwr	OP-22	vdsat	OP-9
cgs	OP-17	ld	I-9	rd	M-38	vgs	OP-5
cgso	M-30	ld	M-19	rdc	M-43	vmax	M-8
cj	M-58	ldd	M-96	rdd	M-41	vth	OP-8
cjsw	M-62	ldif	M-45	rdeff	O-4	vto	M-2
cta	M-86	lds	M-97	region	I-12	w	M-88
ctp	M-87	leff	O-2	region	OP-2	w	I-1
delta	M-13	lgcd	M-48	reversed	OP-3	wai0	M-26
dskip	M-53	lgcs	M-47	ron	OP-20	wbi0	M-29
ef	M-100	ls	I-10	rs	M-37	wd	M-20
eg	M-74	m	I-11	rsc	M-42	weff	O-1
flex	M-77	meto	M-33	rseff	O-3	xj	M-23
fc	M-61	minr	M-44	rsh	M-39	xl	M-22
fcsw	M-65	mj	M-59	rss	M-40	xpart	M-35
gameff	OP-13	mjsw	M-63	sc	M-49	xqc	M-36
gamma	M-6	n	M-52	tcv	M-83	xti	M-81
gap1	M-75	neff	M-12	tlev	M-72	xw	M-21

Component Statements Part III

This chapter discusses the following topics:

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- [MOS Level-3 Transistor \(mos3\)](#) on page 503
- [Long Channel JFET/MOSFET Model \(mos30\)](#) on page 520
- [Long Channel JFET/MOSFET Model \(mos3002\)](#) on page 524
- [Long Channel JFET/MOSFET Model \(mos3100\)](#) on page 528
- [Silicon On Isolator JFET Model \(mos40\)](#) on page 532
- [Compact MOS-Transistor Model \(mos705\)](#) on page 536
- [Compact MOS-Transistor Model \(mos902\)](#) on page 546
- [Compact MOS-Transistor Model \(mos903\)](#) on page 557
- [Microstrip Line \(msli432'1'ne\)](#) on page 570
- [Multi-Conductor Transmission Line \(mtline\)](#) on page 570
- [Mutual Inductor \(mutual inductor\)](#) on page 577
- [Node Capacitance \(nodcap\)](#) on page 578
- [Set Node Quantities \(node\)](#) on page 582
- [Linear N Port \(nport\)](#) on page 583
- [Parameter Value Tester \(paramtest\)](#) on page 589
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- [Polynomial Current Controlled Voltage Source \(pccvs\)](#) on page 592
- [Physical Resistor \(phy_res\)](#) on page 594
- [Independent Resistive Source \(port\)](#) on page 602

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Component Statements Part III

- [Poly-Si TFT \(psitft\)](#) on page 608
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- [s-Domain Current Controlled Voltage Source \(sccvs\)](#) on page 637
- [s-Domain Linear Voltage Controlled Current Source \(svccs\)](#) on page 639
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- [VBIC Bipolar Transistor \(vbic\)](#) on page 665
- [Linear Voltage Controlled Current Source \(vccs\)](#) on page 678
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- [Winding for Magnetic Core \(winding\)](#) on page 688
- [z-Domain Linear Current Controlled Current Source \(zcccs\)](#) on page 689
- [z-Domain Current Controlled Voltage Source \(zccvs\)](#) on page 691
- [z-Domain Linear Voltage Controlled Current Source \(zvccs\)](#) on page 694
- [z-Domain Voltage Controlled Voltage Source \(zvcvs\)](#) on page 696

MOS Level-2 Transistor (mos2)

Description

The MOS2 model is the level-2 model from Berkeley SPICE. The MOS2 model is an analytical, one-dimensional model that incorporates most of the second-order small-size effects. A smoother version of the level-2 model (with continuous G_{ds} at V_{dsat}) is also available. Three charge models are available. MOS2 transistors require that you use a model statement.

This device is supported within altergroups.

Sample Instance Statement

```
mn2 (1 2 0 0) nch2 w=10u ad=20p as=20p ps=24u pd=24u
```

Sample Model Statement:

```
model nch2 mos2 type=n vto=0.66 lambda=0.018 gamma=0.6 nsub=0.213e16 kp=0.978e-4  
tpg=-1 vmax=6e4 ucrit=1e7 utra=0.1 uexp=0.2 is=0
```

Instance Definition

```
Name d g s b ModelName parameter=value ...
```

Instance Parameters

- | | | |
|---|----------------------|--|
| 1 | w (m) | Channel width. |
| 2 | l (m) | Channel length. |
| 3 | as (m ²) | Area of source diffusion. |
| 4 | ad (m ²) | Area of drain diffusion. |
| 5 | ps (m) | Perimeter of source diffusion. |
| 6 | pd (m) | Perimeter of drain diffusion. |
| 7 | nrd (m/m) | Number of squares of drain diffusion. |
| 8 | nrs (m/m) | Number of squares of source diffusion. |

Spectre Circuit Simulator Reference

Component Statements Part III

9	<code>ld (m)</code>	Length of drain diffusion region.
10	<code>ls (m)</code>	Length of source diffusion region.
11	<code>m=1</code>	Multiplicity factor (number of MOSFETs in parallel).
12	<code>region=triode</code>	Estimated operating region. Spectre outputs number (0-4) in a rawfile. Possible values are <code>off</code> , <code>triode</code> , <code>sat</code> , <code>subth</code> , or <code>breakdown</code> .
13	<code>trise</code>	Temperature rise from ambient.
14	<code>degradation=no</code>	Hot-electron degradation flag. Possible values are <code>no</code> or <code>yes</code> .

Model Definition

`model modelName mos2 parameter=value ...`

Model Parameters

Device type parameters

1	<code>type=n</code>	Transistor type. Possible values are <code>n</code> or <code>p</code> .
---	---------------------	--

Drain current model parameters

2	<code>vto=0 V</code>	Threshold voltage at zero body bias.
3	<code>kp=2.0718e-5 A/V²</code>	Transconductance parameter.
4	<code>lambda=0 1/V</code>	Channel length modulation parameter.
5	<code>phi=0.7 V</code>	Surface potential at strong inversion.
6	<code>gamma=0 \sqrt{V}</code>	Body-effect parameter.
7	<code>uo=600 cm²/V s</code>	Carrier surface mobility.

Spectre Circuit Simulator Reference

Component Statements Part III

8	$v_{max}=\infty$ m/s	Carrier saturation velocity.
9	$u_{crit}=0$ V/cm	Critical field for mobility degradation.
10	$u_{exp}=0$	Critical field exponent for mobility degradation.
11	$u_{tra}=0$ 1/V	Transverse field for mobility.
12	$n_{eff}=1$	Total channel charge coefficient.
13	$\delta=0$	Width effect on threshold voltage.
14	$smooth=yes$	Drain current smoothing flag. Possible values are <code>no</code> or <code>yes</code> .

Process parameters

15	$n_{sub}=1.13e16$ cm ⁻³	Channel doping concentration.
16	$n_{ss}=0$ cm ⁻²	Surface state density.
17	$n_{fs}=0$ cm ⁻²	Fast surface state density.
18	$t_{pg}=+1$	Type of gate (+1 = opposite of substate, -1 = same as substate, 0 = aluminum).
19	$t_{ox}=1e-7$ m	Gate oxide thickness.
20	$l_d=0$ m	Lateral diffusion.
21	$w_d=0$ m	Field-oxide encroachment.
22	$x_w=0$ m	Width variation due to masking and etching.
23	$x_l=0$ m	Length variation due to masking and etching.
24	$x_j=0$ m	Source/drain junction depth.

Spectre Circuit Simulator Reference

Component Statements Part III

Impact ionization parameters

- 25 `ai0=0` 1/V Impact ionization current coefficient.
- 26 `lai0=0` $\mu\text{m}/\text{V}$ Length sensitivity of `ai0`.
- 27 `wai0=0` $\mu\text{m}/\text{V}$ Width sensitivity of `ai0`.
- 28 `bi0=0` V Impact ionization current exponent.
- 29 `lbi0=0` μm V Length sensitivity of `bi0`.
- 30 `wbi0=0` μm V Width sensitivity of `bi0`.

Overlap capacitance parameters

- 31 `cgso=0` F/m Gate-source overlap capacitance.
- 32 `cgdo=0` F/m Gate-drain overlap capacitance.
- 33 `cgbo=0` F/m Gate-bulk overlap capacitance.
- 34 `meto=0` m Metal overlap in fringing field.

Charge model selection parameters

- 35 `capmod=bsim` Intrinsic charge model.
Possible values are `none`, `meyer`, `yang`, or `bsim`.
- 36 `xpart=1` Drain/source channel charge partition in saturation for BSIM charge model, use 0.0 for 40/60, 0.5 for 50/50, or 1.0 for 0/100.
- 37 `xqc=0` Drain/source channel charge partition in saturation for charge models, e.g. use 0.4 for 40/60, 0.5 for 50/50, 0 for 0/100.

Parasitic resistance parameters

- 38 `rs=0` Ω Source resistance.
- 39 `rd=0` Ω Drain resistance.

Spectre Circuit Simulator Reference

Component Statements Part III

40	<code>rsh=0</code>	Ω/sqr	Source/drain diffusion sheet resistance.
41	<code>rss=0</code>	Ω m	Scalable source resistance.
42	<code>rdd=0</code>	Ω m	Scalable drain resistance.
43	<code>rsc=0</code>	Ω	Source contact resistance.
44	<code>rdc=0</code>	Ω	Drain contact resistance.
45	<code>minr=0.1</code>	Ω	Minimum source/drain resistance.
46	<code>ldif=0</code>	m	Lateral diffusion beyond the gate.
47	<code>hdif=0</code>	m	Length of heavily doped diffusion.
48	<code>lgcs=0</code>	m	Gate-to-contact length of source side.
49	<code>lgcd=0</code>	m	Gate-to-contact length of drain side.
50	<code>sc=∞</code>	m	Spacing between contacts.

Junction diode model parameters

51	<code>js</code>	(A/m ²)	Bulk junction reverse saturation current density.
52	<code>is=1e-14</code>	A	Bulk junction reverse saturation current.
53	<code>n=1</code>		Junction emission coefficient.
54	<code>dskip=yes</code>		Use simple piece-wise linear model for diode currents below 0.1*iabstol. Possible values are <code>no</code> or <code>yes</code> .
55	<code>imelt=<code>`imaxA`</code></code>		Explosion current, diode is linearized beyond this current to aid convergence.
56	<code>jmelt=<code>`jmaxA/m`</code>²</code>		Explosion current density, diode is linearized beyond this current to aid convergence.

Spectre Circuit Simulator Reference

Component Statements Part III

Junction capacitance model parameters

57	<code>cbs=0 F</code>	Bulk-source zero-bias junction capacitance.
58	<code>cbd=0 F</code>	Bulk-drain zero-bias junction capacitance.
59	<code>cj=0 F/m²</code>	Zero-bias junction bottom capacitance density.
60	<code>mj=1/2</code>	Bulk junction bottom grading coefficient.
61	<code>pb=0.8 V</code>	Bulk junction built-in potential.
62	<code>fc=0.5</code>	Forward-bias depletion capacitance threshold.
63	<code>cjsw=0 F/m</code>	Zero-bias junction sidewall capacitance density.
64	<code>mjsw=1/3</code>	Bulk junction sidewall grading coefficient.
65	<code>pbsw=0.8 V</code>	Side-wall junction built-in potential.
66	<code>fcsw=0.5</code>	Side-wall forward-bias depletion capacitance threshold.

Operating region warning control parameters

67	<code>alarm=none</code>	Forbidden operating region. Possible values are <code>none</code> , <code>off</code> , <code>triode</code> , <code>sat</code> , <code>subth</code> , or <code>rev</code> .
68	<code>imax=1 A</code>	Maximum current, currents above this limit generate a warning.
69	<code>jmax=1e8 A/m²</code>	Maximum current density, currents above this limit generate a warning.
70	<code>bvj=∞ V</code>	Junction reverse breakdown voltage.
71	<code>vbox=1e9 tox V</code>	Oxide breakdown voltage.

Temperature effects parameters

72	<code>tnom (C)</code>	Parameters measurement temperature. Default set by <code>options</code> .
73	<code>trise=0 C</code>	Temperature rise from ambient.

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Component Statements Part III

74	<code>uto=0 C</code>	Mobility temperature offset.
75	<code>ute=-1.5</code>	Mobility temperature exponent.
76	<code>tlev=0</code>	DC temperature selector.
77	<code>tlevc=0</code>	AC temperature selector.
78	<code>eg=1.12452 V</code>	Energy band gap.
79	<code>gap1=7.02e-4 V/C</code>	Band gap temperature coefficient.
80	<code>gap2=1108 C</code>	Band gap temperature offset.
81	<code>flex=0</code>	Temperature exponent for <code>ucrit</code> .
82	<code>lamex=0 1/C</code>	Temperature parameter for <code>lambda</code> and <code>kappa</code> .
83	<code>trs=0 1/C</code>	Temperature parameter for source resistance.
84	<code>trd=0 1/C</code>	Temperature parameter for drain resistance.
85	<code>x_{ti}=3</code>	Saturation current temperature exponent.
86	<code>ptc=0 V/C</code>	Surface potential temperature coefficient.
87	<code>tcv=0 V/C</code>	Threshold voltage temperature coefficient.
88	<code>pta=0 V/C</code>	Junction potential temperature coefficient.
89	<code>ptp=0 V/C</code>	Sidewall junction potential temperature coefficient.
90	<code>cta=0 1/C</code>	Junction capacitance temperature coefficient.
91	<code>ctp=0 1/C</code>	Sidewall junction capacitance temperature coefficient.

Default instance parameters

92	<code>w=3e-6 m</code>	Default channel width.
93	<code>l=3e-6 m</code>	Default channel length.

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Component Statements Part III

94	<code>as=0</code>	m^2	Default area of source diffusion.
95	<code>ad=0</code>	m^2	Default area of drain diffusion.
96	<code>ps=0</code>	m	Default perimeter of source diffusion.
97	<code>pd=0</code>	m	Default perimeter of drain diffusion.
98	<code>nrd=0</code>	m/m	Default number of squares of drain diffusion.
99	<code>nrs=0</code>	m/m	Default number of squares of source diffusion.
100	<code>ldd=0</code>	m	Default length of drain diffusion region.
101	<code>lds=0</code>	m	Default length of source diffusion region.

Noise model parameters

102	<code>noisemod=1</code>		Noise model selector.
103	<code>kf=0</code>		Flicker (1/f) noise coefficient.
104	<code>af=1</code>		Flicker (1/f) noise exponent.
105	<code>ef=1</code>		Flicker (1/f) noise frequency exponent.
106	<code>wnoi=1e-5</code>	m	Channel width at which noise parameters were extracted.

Auto Model Selector parameters

107	<code>wmax=1.0</code>	m	Maximum channel width for which the model is valid.
108	<code>wmin=0.0</code>	m	Minimum channel width for which the model is valid.
109	<code>lmax=1.0</code>	m	Maximum channel length for which the model is valid.
110	<code>lmin=0.0</code>	m	Minimum channel length for which the model is valid.

Spectre Circuit Simulator Reference

Component Statements Part III

Degradation parameters

111	<code>degramod=spectre</code>	Degradation model selector. Possible values are <code>spectre</code> or <code>bert</code> .
112	<code>degradation=no</code>	Hot-electron degradation flag. Possible values are <code>no</code> or <code>yes</code> .
113	<code>dvthc=1</code> V	Degradation coefficient for threshold voltage.
114	<code>dvthe=1</code>	Degradation exponent for threshold voltage.
115	<code>duoc=1</code> S	Degradation coefficient for transconductance.
116	<code>duoe=1</code>	Degradation exponent for transconductance.
117	<code>crivth=0.1</code> V	Maximum allowable threshold voltage shift.
118	<code>criuo=10%</code>	Maximum allowable normalized mobility change.
119	<code>crigm=10%</code>	Maximum allowable normalized transconductance change.
120	<code>criids=10%</code>	Maximum allowable normalized drain current change.
121	<code>wnom=5e-6</code> m	Nominal device width in degradation calculation.
122	<code>lnom=1e-6</code> m	Nominal device length in degradation calculation.
123	<code>vbsn=0</code> V	Substrate voltage in degradation calculation.
124	<code>vdsni=0.1</code> V	Drain voltage in I_{ds} degradation calculation.
125	<code>vgsni=5</code> V	Gate voltage in I_{ds} degradation calculation.
126	<code>vdsng=0.1</code> V	Drain voltage in G_m degradation calculation.
127	<code>vgsng=5</code> V	Gate voltage in G_m degradation calculation.

Spectre stress parameters

128	<code>esat=1.1e7</code> V/m	Critical field in V_{dsat} calculation.
-----	-----------------------------	---

Spectre Circuit Simulator Reference

Component Statements Part III

129 $esatg=2.5e6 \text{ 1/m}$ Gate voltage dependence of $esat$.

130 $vpg=-0.25$ Gate voltage modifier.

131 $vpb=-0.13$ Gate voltage modifier.

132 $subc1=2.24e-5$ Substrate current coefficient.

133 $subc2=-0.1e-5 \text{ 1/V}$ Substrate current coefficient.

134 $sube=6.4$ Substrate current exponent.

135 $strc=1$ Stress coefficient.

136 $stre=1$ Stress exponent.

BERT stress parameters

137 $h0=1$ Aging coefficient.

138 $hgd=0 \text{ 1/V}$ Bias dependence of $h0$.

139 $m0=1$ Aging exponent.

140 $mgd=0 \text{ 1/V}$ Bias dependence of $m0$.

141 $ecrit0=1.1e5 \text{ V/cm}$ Critical electric field.

142 $lecrit0=0 \text{ }\mu\text{m V/cm}$ Length dependence of $ecrit0$.

143 $wecrit0=0 \text{ }\mu\text{m V/cm}$ Width dependence of $ecrit0$.

144 $ecritg=0 \text{ 1/cm}$ Gate voltage dependence of $ecrit0$.

145 $lecritg=0 \text{ }\mu\text{m/cm}$ Length dependence of $ecritg$.

146 $wecritg=0 \text{ }\mu\text{m/cm}$ Width dependence of $ecritg$.

147 $ecritb=0 \text{ 1/cm}$ Substrate voltage dependence of $ecrit0$.

148 $lecritb=0 \text{ }\mu\text{m/cm}$ Length dependence of $ecritb$.

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Component Statements Part III

149	wecritb=0 $\mu\text{m}/\text{cm}$	Width dependence of ecritb.
150	lc0=1	Substrate current coefficient.
151	llc0=0 μm	Length dependence of lc0.
152	wlc0=0 μm	Width dependence of lc0.
153	lc1=1	Substrate current coefficient.
154	llc1=0 μm	Length dependence of lc1.
155	wlc1=0 μm	Width dependence of lc1.
156	lc2=1	Substrate current coefficient.
157	llc2=0 μm	Length dependence of lc2.
158	wlc2=0 μm	Width dependence of lc2.
159	lc3=1	Substrate current coefficient.
160	llc3=0 μm	Length dependence of lc3.
161	wlc3=0 μm	Width dependence of lc3.
162	lc4=1	Substrate current coefficient.
163	llc4=0 μm	Length dependence of lc4.
164	wlc4=0 μm	Width dependence of lc4.
165	lc5=1	Substrate current coefficient.
166	llc5=0 μm	Length dependence of lc5.
167	wlc5=0 μm	Width dependence of lc5.
168	lc6=1	Substrate current coefficient.
169	llc6=0 μm	Length dependence of lc6.
170	wlc6=0 μm	Width dependence of lc6.

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Component Statements Part III

171 $lc7=1$	Substrate current coefficient.
172 $llc7=0 \mu\text{m}$	Length dependence of $lc7$.
173 $wlc7=0 \mu\text{m}$	Width dependence of $lc7$.

Imax and Imelt

The $imax$ parameter aids convergence and prevents numerical overflow. The junction characteristics of the device are accurately modeled for current up to $imax$. If $imax$ is exceeded during iterations, the linear model is substituted until the current drops below $imax$ or until convergence is achieved. If convergence is achieved with the current exceeding $imax$, the results are inaccurate, and Spectre prints a warning.

A separate model parameter, $imelt$, is used as a limit warning for the junction current. This parameter can be set to the maximum current rating of the device. When any component of the junction current exceeds $imelt$, note that base and collector currents are composed of many exponential terms, Spectre issues a warning and the results become inaccurate. The junction current is linearized above the value of $imelt$ to prevent arithmetic exception, with the exponential term replaced by a linear equation at $imelt$.

Both of these parameters have current density counterparts, $jmax$ and $jmelt$, that you can specify if you want the absolute current values to depend on the device area

Auto Model Selection

Many models need to be characterized for different geometries in order to obtain accurate results for model development. The model selector program automatically searches for a model with the length and width range specified in the instance statement and uses this model in the simulations.

For the auto model selector program to find a specific model, the models to be searched should be grouped together within braces. Such a group is called a model group. An opening brace is required at the end of the line defining each model group. Every model in the group is given a name followed by a colon and the list of parameters. Also, the four geometric parameters $lmax$, $lmin$, $wmax$, and $wmin$ should be given. The selection criteria to choose a model is as follows:

$$lmin \leq inst_length < lmax \quad \text{and} \quad wmin \leq inst_width < wmax$$

Example:

```
model ModelName ModelType {
```

Spectre Circuit Simulator Reference

Component Statements Part III

- 1: <model parameters> lmin=2 lmax=4 wmin=1 wmax=2
- 2: <model parameters> lmin=1 lmax=2 wmin=2 wmax=4
- 3: <model parameters> lmin=2 lmax=4 wmin=4 wmax=6

}

Then for a given instance

```
M1 1 2 3 4 ModelName w=3 l=1.5
```

the program would search all the models in the model group with the name `ModelName` and then pick the first model whose geometric range satisfies the selection criteria. In the preceding example, the auto model selector program would choose `ModelName.2`.

You must specify both length (l) and width (w) on the device instance line to enable automatic model selection.

Output Parameters

- | | | |
|----|---|--|
| 1 | <code>w_{eff}</code> (m) | Effective channel width. |
| 2 | <code>l_{eff}</code> (m) | Effective channel length. |
| 3 | <code>r_{seff}</code> (Ω) | Effective source resistance. |
| 4 | <code>r_{deff}</code> (Ω) | Effective drain resistance. |
| 5 | <code>a_{seff}</code> (m ²) | Effective area of source diffusion. |
| 6 | <code>a_{deff}</code> (m ²) | Effective area of drain diffusion. |
| 7 | <code>p_{seff}</code> (m) | Effective perimeter of source diffusion. |
| 8 | <code>p_{deff}</code> (m) | Effective perimeter of drain diffusion. |
| 9 | <code>i_{sseff}</code> (A) | Effective source-bulk junction reverse saturation current. |
| 10 | <code>i_{sdeff}</code> (A) | Effective drain-bulk junction reverse saturation current. |
| 11 | <code>cb_{seff}</code> (F) | Effective zero-bias source-bulk junction capacitance. |
| 12 | <code>cb_{deff}</code> (F) | Effective zero-bias drain-bulk junction capacitance. |

Spectre Circuit Simulator Reference

Component Statements Part III

Operating-Point Parameters

1	<code>type=n</code>	Transistor type. Possible values are <code>n</code> or <code>p</code> .
2	<code>region=triode</code>	Estimated operating region. Spectre outputs number (0-4) in a rawfile. Possible values are <code>off</code> , <code>triode</code> , <code>sat</code> , <code>subth</code> , or <code>breakdown</code> .
3	<code>degradation=no</code>	Hot-electron degradation flag. Possible values are <code>no</code> or <code>yes</code> .
4	<code>reversed</code>	Reverse mode indicator. Possible values are <code>no</code> or <code>yes</code> .
5	<code>ids (A)</code>	Resistive drain-to-source current.
6	<code>vgs (V)</code>	Gate-source voltage.
7	<code>vds (V)</code>	Drain-source voltage.
8	<code>vbs (V)</code>	Bulk-source voltage.
9	<code>vth (V)</code>	Threshold voltage.
10	<code>vdsat (V)</code>	Drain-source saturation voltage.
11	<code>gm (S)</code>	Common-source transconductance.
12	<code>gds (S)</code>	Common-source output conductance.
13	<code>gmbs (S)</code>	Body-transconductance.
14	<code>gameff (\sqrt{V})</code>	Effective body effect coefficient.
15	<code>betaeff (A/V^2)</code>	Effective beta.
16	<code>cbd (F)</code>	Drain-bulk junction capacitance.
17	<code>cbs (F)</code>	Source-bulk junction capacitance.
18	<code>cgs (F)</code>	Gate-source capacitance.

Spectre Circuit Simulator Reference

Component Statements Part III

19	<code>cgd</code> (F)	Gate-drain capacitance.
20	<code>cgb</code> (F)	Gate-bulk capacitance.
21	<code>ron</code> (Ω)	On-resistance.
22	<code>id</code> (A)	Resistive drain current.
23	<code>ibulk</code> (A)	Resistive bulk current.
24	<code>pwr</code> (W)	Power at op point.
25	<code>gmoverid</code> (1/V)	Gm/Ids.
26	<code>isub</code> (A)	Substrate current.
27	<code>stress</code>	Hot-electron stress.
28	<code>age</code> (s)	Device age.
29	<code>he_vdsat</code> (V)	Hot Electron Vdsat.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

<code>ad</code>	I-4	<code>gap2</code>	M-80	<code>lmin</code>	M-110	<code>tnom</code>	M-72
<code>ad</code>	M-95	<code>gds</code>	OP-12	<code>lnom</code>	M-122	<code>tox</code>	M-19
<code>adefl</code>	O-6	<code>gm</code>	OP-11	<code>ls</code>	I-10	<code>tpg</code>	M-18
<code>af</code>	M-104	<code>gmbs</code>	OP-13	<code>m</code>	I-11	<code>trd</code>	M-84
<code>age</code>	OP-28	<code>gmoverid</code>	OP-25	<code>m0</code>	M-139	<code>trise</code>	I-13
<code>ai0</code>	M-25	<code>h0</code>	M-137	<code>meto</code>	M-34	<code>trise</code>	M-73
<code>alarm</code>	M-67	<code>hdif</code>	M-47	<code>mgd</code>	M-140	<code>trs</code>	M-83
<code>as</code>	I-3	<code>he_vdsat</code>	OP-29	<code>minr</code>	M-45	<code>type</code>	M-1

Spectre Circuit Simulator Reference Component Statements Part III

as	M-94	hgd	M-138	mj	M-60	type	OP-1
aseff	O-5	ibulk	OP-23	mjsw	M-64	ucrit	M-9
betaeff	OP-15	id	OP-22	n	M-53	uexp	M-10
bi0	M-28	ids	OP-5	neff	M-12	uo	M-7
bvj	M-70	imax	M-68	nfs	M-17	ute	M-75
capmod	M-35	imelt	M-55	noisemod	M-102	uto	M-74
cbd	OP-16	is	M-52	nrd	M-98	utra	M-11
cbd	M-58	isdeff	O-10	nrd	I-7	vbox	M-71
cbdeff	O-12	isseff	O-9	nrs	I-8	vbs	OP-8
cbs	OP-17	isub	OP-26	nrs	M-99	vbsn	M-123
cbs	M-57	jmax	M-69	nss	M-16	vds	OP-7
cbseff	O-11	jmelt	M-56	nsub	M-15	vdsat	OP-10
cgb	OP-20	js	M-51	pb	M-61	vdsng	M-126
cgbo	M-33	kf	M-103	pbsw	M-65	vdsni	M-124
cgd	OP-19	kp	M-3	pd	I-6	vgs	OP-6
cgdo	M-32	l	I-2	pd	M-97	vgsng	M-127
cgs	OP-18	l	M-93	pdeff	O-8	vgsni	M-125
cgso	M-31	lai0	M-26	phi	M-5	vmax	M-8
cj	M-59	lambda	M-4	ps	I-5	vpb	M-131
cjsw	M-63	lamex	M-82	ps	M-96	vpg	M-130
crigm	M-119	lbi0	M-29	pseff	O-7	vth	OP-9
criids	M-120	lc0	M-150	pta	M-88	vto	M-2
criuo	M-118	lc1	M-153	ptc	M-86	w	M-92
crivth	M-117	lc2	M-156	ptp	M-89	w	I-1
cta	M-90	lc3	M-159	pwr	OP-24	wai0	M-27
ctp	M-91	lc4	M-162	rd	M-39	wbi0	M-30
degradation	M-112	lc5	M-165	rdc	M-44	wd	M-21
degradation	OP-3	lc6	M-168	rdd	M-42	wecrit0	M-143
degradation	I-14	lc7	M-171	rdeff	O-4	wecritb	M-149
degramod	M-111	ld	M-20	region	OP-2	wecritg	M-146
delta	M-13	ld	I-9	region	I-12	weff	O-1
dskip	M-54	ldd	M-100	reversed	OP-4	wlc0	M-152

Spectre Circuit Simulator Reference Component Statements Part III

duoc	M-115	ldif	M-46	ron	OP-21	wlc1	M-155
duoe	M-116	lds	M-101	rs	M-38	wlc2	M-158
dvthc	M-113	lecrit0	M-142	rsc	M-43	wlc3	M-161
dvthe	M-114	lecritb	M-148	rseff	O-3	wlc4	M-164
ecrit0	M-141	lecritg	M-145	rsh	M-40	wlc5	M-167
ecritb	M-147	leff	O-2	rss	M-41	wlc6	M-170
ecritg	M-144	lgcd	M-49	sc	M-50	wlc7	M-173
ef	M-105	lgcs	M-48	smooth	M-14	wmax	M-107
eg	M-78	llc0	M-151	strc	M-135	wmin	M-108
esat	M-128	llc1	M-154	stre	M-136	wnoi	M-106
esatg	M-129	llc2	M-157	stress	OP-27	wnom	M-121
flex	M-81	llc3	M-160	subc1	M-132	xj	M-24
fc	M-62	llc4	M-163	subc2	M-133	xl	M-23
fcsw	M-66	llc5	M-166	sube	M-134	xpart	M-36
gameff	OP-14	llc6	M-169	tcv	M-87	xqc	M-37
gamma	M-6	llc7	M-172	tlev	M-76	xti	M-85
gap1	M-79	lmax	M-109	tlevc	M-77	xw	M-22

MOS Level-3 Transistor (mos3)

Description

The MOS3 model is the level-3 model from Berkeley SPICE, and is a semi-empirical model. Three charge models are available. MOS3 transistors require that you use a model statement.

This device is supported within altergroups.

Sample Instance Statement

```
mp3 (0 1 2 2) pchmos3 l=2u w=30u ad=120p as=75p pd=36u ps=6u
```

Spectre Circuit Simulator Reference

Component Statements Part III

Sample Model Statement

```
model pchmos3 mos3 type=p vto=-0.83 gamma=0.4511 kappa=5 ld=0.45e-6 kp=0.334e-4  
tox=0.3e-7 nsub=0.2e17 capmod=yang vmax=4.5e5 theta=0.25 cbs=10e-15 cbd=10e-15
```

Instance Definition

```
Name d g s b ModelName parameter=value ...
```

Instance Parameters

1	w (m)	Channel width.
2	l (m)	Channel length.
3	as (m ²)	Area of source diffusion.
4	ad (m ²)	Area of drain diffusion.
5	ps (m)	Perimeter of source diffusion.
6	pd (m)	Perimeter of drain diffusion.
7	nrd (m/m)	Number of squares of drain diffusion.
8	nrs (m/m)	Number of squares of source diffusion.
9	ld (m)	Length of drain diffusion region.
10	ls (m)	Length of source diffusion region.
11	m=1	Multiplicity factor (number of MOSFETs in parallel).
12	region=triode	Estimated operating region. Spectre outputs number (0-4) in a rawfile. Possible values are off, triode, sat, subth, or breakdown.
13	trise	Temperature rise from ambient.
14	degradation=no	Hot-electron degradation flag. Possible values are no or yes.

Spectre Circuit Simulator Reference

Component Statements Part III

Model Definition

model modelName mos3 parameter=value ...

Model Parameters

Device type parameters

1 type=n Transistor type.
Possible values are n or p.

Drain current model parameters

2 vto=0 V Threshold voltage at zero body bias.

3 kp=2.0718e-5 A/V² Transconductance parameter.

4 theta=0 1/V Mobility modulation coefficient.

5 phi=0.7 V Surface potential at strong inversion.

6 gamma=0 \sqrt{V} Body-effect parameter.

7 uo=600 cm²/V s Carrier surface mobility.

8 vmax= ∞ m/s Carrier saturation velocity.

9 eta=0 1/V Static feedback coefficient.

10 kappa=0.2 Saturation field factor.

11 delta=0 Width effect on threshold voltage.

Process parameters

12 nsub=1.13e16 cm⁻³ Channel doping concentration.

13 nss=0 cm⁻² Surface state density.

Spectre Circuit Simulator Reference

Component Statements Part III

14	$nfs=0 \text{ cm}^{-2}$	Fast surface state density.
15	$tpg=+1$	Type of gate (+1 = opposite of substate, -1 = same as substate, 0 = aluminum).
16	$tox=1e-7 \text{ m}$	Gate oxide thickness.
17	$ld=0 \text{ m}$	Lateral diffusion.
18	$wd=0 \text{ m}$	Field-oxide encroachment.
19	$xw=0 \text{ m}$	Width variation due to masking and etching.
20	$xl=0 \text{ m}$	Length variation due to masking and etching.
21	$xj=0 \text{ m}$	Source/drain junction depth.

Impact ionization parameters

22	$ai0=0 \text{ 1/V}$	Impact ionization current coefficient.
23	$lai0=0 \text{ }\mu\text{m/V}$	Length sensitivity of $ai0$.
24	$wai0=0 \text{ }\mu\text{m/V}$	Width sensitivity of $ai0$.
25	$bi0=0 \text{ V}$	Impact ionization current exponent.
26	$lbi0=0 \text{ }\mu\text{m V}$	Length sensitivity of $bi0$.
27	$wbi0=0 \text{ }\mu\text{m V}$	Width sensitivity of $bi0$.

Overlap capacitance parameters

28	$cgs0=0 \text{ F/m}$	Gate-source overlap capacitance.
29	$cgd0=0 \text{ F/m}$	Gate-drain overlap capacitance.
30	$cgbo=0 \text{ F/m}$	Gate-bulk overlap capacitance.
31	$meto=0 \text{ m}$	Metal overlap in fringing field.

Spectre Circuit Simulator Reference

Component Statements Part III

Charge model selection parameters

- 32 `capmod=bsim` Intrinsic charge model.
Possible values are `none`, `meyer`, `yang`, or `bsim`.
- 33 `xpart=1` Drain/source channel charge partition in saturation for BSIM charge model, use 0.0 for 40/60, 0.5 for 50/50, or 1.0 for 0/100.
- 34 `xqc=0` Drain/source channel charge partition in saturation for charge models, e.g. use 0.4 for 40/60, 0.5 for 50/50, 0 for 0/100.

Parasitic resistance parameters

- 35 `rs=0` Ω Source resistance.
- 36 `rd=0` Ω Drain resistance.
- 37 `rsh=0` Ω/sqr Source/drain diffusion sheet resistance.
- 38 `rss=0` Ω m Scalable source resistance.
- 39 `rdd=0` Ω m Scalable drain resistance.
- 40 `rsc=0` Ω Source contact resistance.
- 41 `rdc=0` Ω Drain contact resistance.
- 42 `minr=0.1` Ω Minimum source/drain resistance.
- 43 `ldif=0` m Lateral diffusion beyond the gate.
- 44 `hdif=0` m Length of heavily doped diffusion.
- 45 `lgcs=0` m Gate-to-contact length of source side.
- 46 `lgcd=0` m Gate-to-contact length of drain side.
- 47 `sc= ∞` m Spacing between contacts.

Spectre Circuit Simulator Reference

Component Statements Part III

Junction diode model parameters

48	<code>js</code> (A/m ²)	Bulk junction reverse saturation current density.
49	<code>is=1e-14</code> A	Bulk junction reverse saturation current.
50	<code>n=1</code>	Junction emission coefficient.
51	<code>dskip=yes</code>	Use simple piece-wise linear model for diode currents below 0.1* <code>iabstol</code> . Possible values are <code>no</code> or <code>yes</code> .
52	<code>imelt=`imaxA`</code>	Explosion current, diode is linearized beyond this current to aid convergence.
53	<code>jmelt=`jmaxA/m`²</code>	Explosion current density, diode is linearized beyond this current to aid convergence.

Junction capacitance model parameters

54	<code>cbs=0</code> F	Bulk-source zero-bias junction capacitance.
55	<code>cbd=0</code> F	Bulk-drain zero-bias junction capacitance.
56	<code>cj=0</code> F/m ²	Zero-bias junction bottom capacitance density.
57	<code>mj=1/2</code>	Bulk junction bottom grading coefficient.
58	<code>pb=0.8</code> V	Bulk junction built-in potential.
59	<code>fc=0.5</code>	Forward-bias depletion capacitance threshold.
60	<code>cjsw=0</code> F/m	Zero-bias junction sidewall capacitance density.
61	<code>mjsw=1/3</code>	Bulk junction sidewall grading coefficient.
62	<code>pbsw=0.8</code> V	Side-wall junction built-in potential.
63	<code>fcsw=0.5</code>	Side-wall forward-bias depletion capacitance threshold.

Spectre Circuit Simulator Reference

Component Statements Part III

Operating region warning control parameters

64	alarm=none	Forbidden operating region. Possible values are none, off, triode, sat, subth, or rev.
65	imax=1 A	Maximum current, currents above this limit generate a warning.
66	jmax=1e8 A/m ²	Maximum current density, currents above this limit generate a warning.
67	bvj= ∞ V	Junction reverse breakdown voltage.
68	vbox=1e9 tox V	Oxide breakdown voltage.

Temperature effects parameters

69	tnom (C)	Parameters measurement temperature. Default set by options.
70	trise=0 C	Temperature rise from ambient.
71	uto=0 C	Mobility temperature offset.
72	ute=-1.5	Mobility temperature exponent.
73	tlev=0	DC temperature selector.
74	tlevc=0	AC temperature selector.
75	eg=1.12452 V	Energy band gap.
76	gap1=7.02e-4 V/C	Band gap temperature coefficient.
77	gap2=1108 C	Band gap temperature offset.
78	flex=0	Temperature exponent for ucrit.
79	lamex=0 1/C	Temperature parameter for lambda and kappa.
80	trs=0 1/C	Temperature parameter for source resistance.
81	trd=0 1/C	Temperature parameter for drain resistance.

Spectre Circuit Simulator Reference

Component Statements Part III

82	$x_{ti}=3$	Saturation current temperature exponent.
83	$p_{tc}=0$ V/C	Surface potential temperature coefficient.
84	$t_{cv}=0$ V/C	Threshold voltage temperature coefficient.
85	$p_{ta}=0$ V/C	Junction potential temperature coefficient.
86	$p_{tp}=0$ V/C	Sidewall junction potential temperature coefficient.
87	$c_{ta}=0$ 1/C	Junction capacitance temperature coefficient.
88	$c_{tp}=0$ 1/C	Sidewall junction capacitance temperature coefficient.

Default instance parameters

89	$w=3e-6$ m	Default channel width.
90	$l=3e-6$ m	Default channel length.
91	$a_s=0$ m ²	Default area of source diffusion.
92	$a_d=0$ m ²	Default area of drain diffusion.
93	$p_s=0$ m	Default perimeter of source diffusion.
94	$p_d=0$ m	Default perimeter of drain diffusion.
95	$n_{rd}=0$ m/m	Default number of squares of drain diffusion.
96	$n_{rs}=0$ m/m	Default number of squares of source diffusion.
97	$l_{dd}=0$ m	Default length of drain diffusion region.
98	$l_{ds}=0$ m	Default length of source diffusion region.

Noise model parameters

99	$noisemod=1$	Noise model selector.
100	$k_f=0$	Flicker (1/f) noise coefficient.

Spectre Circuit Simulator Reference

Component Statements Part III

101	<code>af=1</code>	Flicker (1/f) noise exponent.
102	<code>ef=1</code>	Flicker (1/f) noise frequency exponent.
103	<code>wnoi=1e-5 m</code>	Channel width at which noise parameters were extracted.

Auto Model Selector parameters

104	<code>wmax=1.0 m</code>	Maximum channel width for which the model is valid.
105	<code>wmin=0.0 m</code>	Minimum channel width for which the model is valid.
106	<code>lmax=1.0 m</code>	Maximum channel length for which the model is valid.
107	<code>lmin=0.0 m</code>	Minimum channel length for which the model is valid.

Degradation parameters

108	<code>degramod=spectre</code>	Degradation model selector. Possible values are <code>spectre</code> or <code>bert</code> .
109	<code>degradation=no</code>	Hot-electron degradation flag. Possible values are <code>no</code> or <code>yes</code> .
110	<code>dvthc=1 V</code>	Degradation coefficient for threshold voltage.
111	<code>dvthe=1</code>	Degradation exponent for threshold voltage.
112	<code>duoc=1 S</code>	Degradation coefficient for transconductance.
113	<code>duoe=1</code>	Degradation exponent for transconductance.
114	<code>crivth=0.1 V</code>	Maximum allowable threshold voltage shift.
115	<code>criuo=10%</code>	Maximum allowable normalized mobility change.
116	<code>crigm=10%</code>	Maximum allowable normalized transconductance change.
117	<code>criids=10%</code>	Maximum allowable normalized drain current change.
118	<code>wnom=5e-6 m</code>	Nominal device width in degradation calculation.

Spectre Circuit Simulator Reference

Component Statements Part III

119	$l_{nom}=1e-6$ m	Nominal device length in degradation calculation.
120	$v_{bsn}=0$ V	Substrate voltage in degradation calculation.
121	$v_{dsni}=0.1$ V	Drain voltage in I_{ds} degradation calculation.
122	$v_{gsni}=5$ V	Gate voltage in I_{ds} degradation calculation.
123	$v_{dsng}=0.1$ V	Drain voltage in G_m degradation calculation.
124	$v_{gsng}=5$ V	Gate voltage in G_m degradation calculation.

Spectre stress parameters

125	$esat=1.1e7$ V/m	Critical field in V_{dsat} calculation.
126	$esatg=2.5e6$ 1/m	Gate voltage dependence of $esat$.
127	$v_{pg}=-0.25$	Gate voltage modifier.
128	$v_{pb}=-0.13$	Gate voltage modifier.
129	$subc1=2.24e-5$	Substrate current coefficient.
130	$subc2=-0.1e-5$ 1/V	Substrate current coefficient.
131	$sube=6.4$	Substrate current exponent.
132	$strc=1$	Stress coefficient.
133	$stre=1$	Stress exponent.

BERT stress parameters

134	$h0=1$	Aging coefficient.
135	$hgd=0$ 1/V	Bias dependence of $h0$.
136	$m0=1$	Aging exponent.
137	$mgd=0$ 1/V	Bias dependence of $m0$.

Spectre Circuit Simulator Reference Component Statements Part III

- 138 `ecrit0=1.1e5 V/cm` Critical electric field.
- 139 `lecrit0=0 μm V/cm` Length dependence of `ecrit0`.
- 140 `wecrit0=0 μm V/cm` Width dependence of `ecrit0`.
- 141 `ecritg=0 1/cm` Gate voltage dependence of `ecrit0`.
- 142 `lecritg=0 μm/cm` Length dependence of `ecritg`.
- 143 `wecritg=0 μm/cm` Width dependence of `ecritg`.
- 144 `ecritb=0 1/cm` Substrate voltage dependence of `ecrit0`.
- 145 `lecritb=0 μm/cm` Length dependence of `ecritb`.
- 146 `wecritb=0 μm/cm` Width dependence of `ecritb`.
- 147 `lc0=1` Substrate current coefficient.
- 148 `llc0=0 μm` Length dependence of `lc0`.
- 149 `wlc0=0 μm` Width dependence of `lc0`.
- 150 `lc1=1` Substrate current coefficient.
- 151 `llc1=0 μm` Length dependence of `lc1`.
- 152 `wlc1=0 μm` Width dependence of `lc1`.
- 153 `lc2=1` Substrate current coefficient.
- 154 `llc2=0 μm` Length dependence of `lc2`.
- 155 `wlc2=0 μm` Width dependence of `lc2`.
- 156 `lc3=1` Substrate current coefficient.
- 157 `llc3=0 μm` Length dependence of `lc3`.
- 158 `wlc3=0 μm` Width dependence of `lc3`.
- 159 `lc4=1` Substrate current coefficient.

Spectre Circuit Simulator Reference

Component Statements Part III

160	$llc4=0 \mu\text{m}$	Length dependence of $lc4$.
161	$wlc4=0 \mu\text{m}$	Width dependence of $lc4$.
162	$lc5=1$	Substrate current coefficient.
163	$llc5=0 \mu\text{m}$	Length dependence of $lc5$.
164	$wlc5=0 \mu\text{m}$	Width dependence of $lc5$.
165	$lc6=1$	Substrate current coefficient.
166	$llc6=0 \mu\text{m}$	Length dependence of $lc6$.
167	$wlc6=0 \mu\text{m}$	Width dependence of $lc6$.
168	$lc7=1$	Substrate current coefficient.
169	$llc7=0 \mu\text{m}$	Length dependence of $lc7$.
170	$wlc7=0 \mu\text{m}$	Width dependence of $lc7$.

i_{max} and i_{melt}

The i_{max} parameter aids convergence and prevents numerical overflow. The junction characteristics of the device are accurately modeled for current up to i_{max} . If i_{max} is exceeded during iterations, the linear model is substituted until the current drops below i_{max} or until convergence is achieved. If convergence is achieved with the current exceeding i_{max} , the results are inaccurate, and Spectre prints a warning.

A separate model parameter, i_{melt} , is used as a limit warning for the junction current. This parameter can be set to the maximum current rating of the device. When any component of the junction current exceeds i_{melt} , note that base and collector currents are composed of many exponential terms, Spectre issues a warning and the results become inaccurate. The junction current is linearized above the value of i_{melt} to prevent arithmetic exception, with the exponential term replaced by a linear equation at i_{melt} .

Both of these parameters have current density counterparts, j_{max} and j_{melt} , that you can specify if you want the absolute current values to depend on the device area.

Spectre Circuit Simulator Reference

Component Statements Part III

Auto Model Selection

Many models need to be characterized for different geometries in order to obtain accurate results for model development. The model selector program automatically searches for a model with the length and width range specified in the instance statement and uses this model in the simulations.

For the auto model selector program to find a specific model, the models to be searched should be grouped together within braces. Such a group is called a model group. An opening brace is required at the end of the line defining each model group. Every model in the group is given a name followed by a colon and the list of parameters. Also, the four geometric parameters `lmax`, `lmin`, `wmax`, and `wmin` should be given. The selection criteria to choose a model is as follows:

$$l_{min} \leq inst_length < l_{max} \text{ and } w_{min} \leq inst_width < w_{max}$$

Example:

```
model ModelName ModelType {  
    1: <model parameters> lmin=2 lmax=4 wmin=1 wmax=2  
    2: <model parameters> lmin=1 lmax=2 wmin=2 wmax=4  
    3: <model parameters> lmin=2 lmax=4 wmin=4 wmax=6  
}
```

Then for a given instance

```
M1 1 2 3 4 ModelName w=3 l=1.5
```

the program would search all the models in the model group with the name `ModelName` and then pick the first model whose geometric range satisfies the selection criteria. In the preceding example, the auto model selector program would choose `ModelName.2`.

You must specify both length (`l`) and width (`w`) on the device instance line to enable automatic model selection.

Output Parameters

- | | | |
|---|----------------------------------|---------------------------|
| 1 | <code>w_{eff}</code> (m) | Effective channel width. |
| 2 | <code>l_{eff}</code> (m) | Effective channel length. |

Spectre Circuit Simulator Reference

Component Statements Part III

3	<code>rseff</code> (Ω)	Effective source resistance.
4	<code>rdeff</code> (Ω)	Effective drain resistance.
5	<code>aseff</code> (m^2)	Effective area of source diffusion.
6	<code>adeff</code> (m^2)	Effective area of drain diffusion.
7	<code>pseff</code> (m)	Effective perimeter of source diffusion.
8	<code>pdeff</code> (m)	Effective perimeter of drain diffusion.
9	<code>isseff</code> (A)	Effective source-bulk junction reverse saturation current.
10	<code>isdeff</code> (A)	Effective drain-bulk junction reverse saturation current.
11	<code>cbseff</code> (F)	Effective zero-bias source-bulk junction capacitance.
12	<code>cbdeff</code> (F)	Effective zero-bias drain-bulk junction capacitance.

Operating-Point Parameters

1	<code>type=n</code>	Transistor type. Possible values are <code>n</code> or <code>p</code> .
2	<code>region=triode</code>	Estimated operating region. Spectre outputs number (0-4) in a rawfile. Possible values are <code>off</code> , <code>triode</code> , <code>sat</code> , <code>subth</code> , or <code>breakdown</code> .
3	<code>degradation=no</code>	Hot-electron degradation flag. Possible values are <code>no</code> or <code>yes</code> .
4	<code>reversed</code>	Reverse mode indicator. Possible values are <code>no</code> or <code>yes</code> .
5	<code>ids</code> (A)	Resistive drain-to-source current.
6	<code>vgs</code> (V)	Gate-source voltage.
7	<code>vds</code> (V)	Drain-source voltage.
8	<code>vbs</code> (V)	Bulk-source voltage.

Spectre Circuit Simulator Reference

Component Statements Part III

9	vth (V)	Threshold voltage.
10	vdsat (V)	Drain-source saturation voltage.
11	gm (S)	Common-source transconductance.
12	gds (S)	Common-source output conductance.
13	gmbs (S)	Body-transconductance.
14	gameff (\sqrt{V})	Effective body effect coefficient.
15	betaeff (A/V^2)	Effective beta.
16	cbd (F)	Drain-bulk junction capacitance.
17	cbs (F)	Source-bulk junction capacitance.
18	cgs (F)	Gate-source capacitance.
19	cgd (F)	Gate-drain capacitance.
20	cgb (F)	Gate-bulk capacitance.
21	ron (Ω)	On-resistance.
22	id (A)	Resistive drain current.
23	ibulk (A)	Resistive bulk current.
24	pwr (W)	Power at op point.
25	gmoverid (1/V)	Gm/Ids.
26	isub (A)	Substrate current.
27	stress	Hot-electron stress.
28	age (s)	Device age.
29	he_vdsat (V)	Hot Electron Vdsat.

Spectre Circuit Simulator Reference

Component Statements Part III

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

ad	M-92	gap1	M-76	lmax	M-106	tnom	M-69
ad	I-4	gap2	M-77	lmin	M-107	tox	M-16
adefeff	O-6	gds	OP-12	lnom	M-119	tpg	M-15
af	M-101	gm	OP-11	ls	I-10	trd	M-81
age	OP-28	gmbs	OP-13	m	I-11	trise	I-13
ai0	M-22	gmoverid	OP-25	m0	M-136	trise	M-70
alarm	M-64	h0	M-134	meto	M-31	trs	M-80
as	I-3	hdif	M-44	mgd	M-137	type	OP-1
as	M-91	he_vdsat	OP-29	minr	M-42	type	M-1
aseff	O-5	hgd	M-135	mj	M-57	uo	M-7
betaeff	OP-15	ibulk	OP-23	mjsw	M-61	ute	M-72
bi0	M-25	id	OP-22	n	M-50	uto	M-71
bvj	M-67	ids	OP-5	nfs	M-14	vbox	M-68
capmod	M-32	imax	M-65	noisemod	M-99	vbs	OP-8
cbd	M-55	imelt	M-52	nrd	M-95	vbsn	M-120
cbd	OP-16	is	M-49	nrd	I-7	vds	OP-7
cbdefeff	O-12	isdefeff	O-10	nrs	I-8	vdsat	OP-10
cbs	OP-17	isseff	O-9	nrs	M-96	vdsng	M-123
cbs	M-54	isub	OP-26	nss	M-13	vdsni	M-121
cbseff	O-11	jmax	M-66	nsub	M-12	vgs	OP-6
cgb	OP-20	jmelt	M-53	pb	M-58	vgsng	M-124
cgbo	M-30	js	M-48	pbsw	M-62	vgsni	M-122
cgd	OP-19	kappa	M-10	pd	I-6	vmax	M-8
cgdo	M-29	kf	M-100	pd	M-94	vpb	M-128
cgs	OP-18	kp	M-3	pdefeff	O-8	vpg	M-127

Spectre Circuit Simulator Reference Component Statements Part III

cgso	M-28	l	M-90	phi	M-5	vth	OP-9
cj	M-56	l	I-2	ps	I-5	vto	M-2
cjsw	M-60	lai0	M-23	ps	M-93	w	M-89
crigm	M-116	lamex	M-79	pseff	O-7	w	I-1
criids	M-117	lbi0	M-26	pta	M-85	wai0	M-24
criuo	M-115	lc0	M-147	ptc	M-83	wbi0	M-27
crivth	M-114	lc1	M-150	ptp	M-86	wd	M-18
cta	M-87	lc2	M-153	pwr	OP-24	wecrit0	M-140
ctp	M-88	lc3	M-156	rd	M-36	wecritb	M-146
degradation	M-109	lc4	M-159	rdc	M-41	wecritg	M-143
degradation	OP-3	lc5	M-162	rdd	M-39	weff	O-1
degradation	I-14	lc6	M-165	rdeff	O-4	wlc0	M-149
degramod	M-108	lc7	M-168	region	I-12	wlc1	M-152
delta	M-11	ld	M-17	region	OP-2	wlc2	M-155
dskip	M-51	ld	I-9	reversed	OP-4	wlc3	M-158
duoc	M-112	ldd	M-97	ron	OP-21	wlc4	M-161
duoe	M-113	ldif	M-43	rs	M-35	wlc5	M-164
dvthc	M-110	lds	M-98	rsc	M-40	wlc6	M-167
dvthe	M-111	lecrit0	M-139	rseff	O-3	wlc7	M-170
ecrit0	M-138	lecritb	M-145	rsh	M-37	wmax	M-104
ecritb	M-144	lecritg	M-142	rss	M-38	wmin	M-105
ecritg	M-141	leff	O-2	sc	M-47	wnoi	M-103
ef	M-102	lgcd	M-46	strc	M-132	wnom	M-118
eg	M-75	lgcs	M-45	stre	M-133	xj	M-21
esat	M-125	llc0	M-148	stress	OP-27	xl	M-20
esatg	M-126	llc1	M-151	subc1	M-129	xpart	M-33
eta	M-9	llc2	M-154	subc2	M-130	xqc	M-34
flex	M-78	llc3	M-157	sube	M-131	xti	M-82
fc	M-59	llc4	M-160	tcv	M-84	xw	M-19
fcsw	M-63	llc5	M-163	theta	M-4		
gameff	OP-14	llc6	M-166	tlev	M-73		
gamma	M-6	llc7	M-169	tlevc	M-74		

Long Channel JFET/MOSFET Model (mos30)

Description

This long channel JFET/MOSFET model is specially developed to describe the drift region of LDMOS, EPMOS and VDMOS devices. It is described in the Philips MOST Modelbook (Dec.95) as MOS model, level 30 (Used for DMOS). Information on how to obtain this document can be found on Source Link by searching for Philips.

Note: In noise analysis, mos30 instances will not generate any contribution, since there are no noise sources included in the mos30 model.

Warning: Dont use this model. It is obsolete.

Mos30 will be removed from spectre in the next release.

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This device is supported within altergroups.

This device is dynamically loaded from the shared object /vobs/spectre_dev/tools.sun4v/spectre/lib/cmi/3.0.doc/libphilips_sh.so

Sample Instance Statement

```
mn30 (1 2 0 0) nchmod area=2 mult=1
```

Sample Model Statement

```
model nchmod mos30 type=n tox=1.1e-5 ron=150 rsat=500 psat=2 vsat=1 vsub=0.59  
cgate=1.65e-12 csub=1.1e-9 tref=25
```

Instance Definition

```
Name d g s [b] ModelName parameter=value ...
```

Instance Parameters

- | | | |
|---|--------|--------------------------------|
| 1 | mult=1 | Number of devices in parallel. |
| 2 | area=1 | Alias of mult. |

Spectre Circuit Simulator Reference

Component Statements Part III

- 3 `region=triode` Estimated DC operating region, used as a convergence aid. Possible values are `off`, `triode`, `sat`, or `subth`.
- 4 `m=1` Multiplicity factor.

Model Definition

```
model modelName mos30 parameter=value ...
```

Model Parameters

- 1 `type=n` Transistor gender. Possible values are `n` or `p`.
- 2 `ron=1.0 Ω` Ohmic resistance at zero bias.
- 3 `rsat=1.0 Ω` Space charge resistance at zero bias.
- 4 `vsat=10.0 V` Critical drain-source voltage for hot carriers.
- 5 `psat=1.0` Velocity saturation coefficient.
- 6 `vp=-1.0 V` Pinch off voltage at zero gate and substrate voltages.
- 7 `tox=-1.0 cm` Gate oxide thickness.
- 8 `dch=1.0e15 cm-3` Doping level channel.
- 9 `dsub=1.0e15 cm-3` Doping level substrate.
- 10 `vsub=0.6 V` Substrate diffusion voltage.
- 11 `vgap=1.2 V` Bandgap voltage channel.
- 12 `cgate=0.0 F` Gate capacitance at zero bias.
- 13 `csub=0.0 F` Substrate capacitance at zero bias.
- 14 `tausc=0.0 s` Space charge transit time of the channel.
- 15 `ach=0.0` Temperature coefficient resistivity of the channel.
- 16 `kf=0.0` Flickernoise coefficient.

Spectre Circuit Simulator Reference

Component Statements Part III

17	af=1.0	Flickernoise exponent.
18	tr (C)	Reference temperature. Default set by option tnom.
19	tref (C)	Alias of tr. Default set by option tnom.
20	tnom (C)	Alias of tr. Default set by option tnom.
21	dta=0.0 K	Temperature offset of the device.
22	trise=0.0 K	Alias of dta.

Output Parameters

1	ront (Ω)	Ohmic resistance at zero bias.
2	rsat (Ω)	Space charge resistance at zero bias.
3	vsatt (V)	Critical drain-source voltage for hot carriers.
4	vsubt (V)	Substrate diffusion voltage.
5	cgate (F)	Gate capacitance at zero bias.
6	csubt (F)	Substrate capacitance at zero bias.

Operating-Point Parameters

1	pwr (W)	Power.
2	ids (A)	Total current including velocity saturation.
3	qb (Coul)	Substrate charge.
4	qg (Coul)	Gate charge.
5	qds (Coul)	Space charge in the channel.
6	gdsc (S)	Conductance (d ids / d vd).
7	gdsg (S)	Conductance (d ids / d vg).

Spectre Circuit Simulator Reference

Component Statements Part III

8	gdss (S)	Conductance (d ids / d vs).
9	gdsb (S)	Conductance (d ids / d vb).
10	cbd (F)	Capacitance (d qb / d vd).
11	cbg (F)	Capacitance (d qb / d vg).
12	cbs (F)	Capacitance (d qb / d vs).
13	cbb (F)	Capacitance (d qb / d vb).
14	cgd (F)	Capacitance (d qg / d vd).
15	cgg (F)	Capacitance (d qg / d vg).
16	cgs (F)	Capacitance (d qg / d vs).
17	cgb (F)	Capacitance (d qg / d vb).
18	cdsd (F)	Capacitance (d qds / d vd).
19	cdsg (F)	Capacitance (d qds / d vg).
20	cdss (F)	Capacitance (d qds / d vs).
21	cdsb (F)	Capacitance (d qds / d vb).

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

ach	M-15	cgd	OP-14	m	I-4	tox	M-7
af	M-17	cgg	OP-15	mult	I-1	tr	M-18
area	I-2	cgs	OP-16	psat	M-5	tref	M-19
cbb	OP-13	csub	M-13	pwr	OP-1	trise	M-22

Spectre Circuit Simulator Reference Component Statements Part III

cbd	OP-10	csubt	O-6	qb	OP-3	type	M-1
cbg	OP-11	dch	M-8	qds	OP-5	vgap	M-11
cbs	OP-12	dsub	M-9	qg	OP-4	vp	M-6
cdsb	OP-21	dta	M-21	region	I-3	vsat	M-4
cdsd	OP-18	gdsb	OP-9	ron	M-2	vsatt	O-3
cdsg	OP-19	gdsd	OP-6	ront	O-1	vsub	M-10
cdss	OP-20	gdsg	OP-7	rsat	O-2	vsubt	O-4
cgate	O-5	gdss	OP-8	rsat	M-3		
cgate	M-12	ids	OP-2	tausc	M-14		
cgb	OP-17	kf	M-16	tnom	M-20		

Long Channel JFET/MOSFET Model (mos3002)

Description

This long channel JFET/MOSFET model is specially developed to describe the drift region of LDMOS, EP MOS and VDMOS devices. It is described in the Philips MOST Modelbook (Dec.98) as MOS model, level 3002 (Used for DMOS). Information on how to obtain this document can be found on Source Link by searching for Philips.

Note: In noise analysis, mos3002 instances will not generate any contribution, since there are no noise sources included in the mos3002 model.

(c) Philips Electronics N.V. 1993, 1994, 1996, 1998

This device is supported within altergroups.

This device is dynamically loaded from the shared object /vobs/spectre_dev/tools.sun4v/spectre/lib/cmi/3.0.doc/libphilips_sh.so

Sample Instance Statement

```
mn3 (1 2 0 0) nch3002 area=1 m=2
```

Sample Model Statement

```
model nch3002 mos3002 ron=20 rsat=150 vsat=1 tox=1.23e-5 dch=1.1e16 vsub=0.58  
csub=5.43e-13 tausc=1.2e-12 kf=1 tref=27 psat=1 dta=0
```

Spectre Circuit Simulator Reference

Component Statements Part III

Instance Definition

Name d g s [b] ModelName parameter=value ...

Instance Parameters

- | | | |
|---|---------------|--|
| 1 | mult=1 | Number of devices in parallel. |
| 2 | area=1 | Alias of mult. |
| 3 | region=triode | Estimated DC operating region, used as a convergence aid.
Possible values are off, triode, sat, or subth. |
| 4 | m=1 | Multiplicity factor. |

Model Definition

model modelName mos3002 parameter=value ...

Model Parameters

- | | | |
|----|------------------------------|--|
| 1 | type=n | Transistor gender.
Possible values are n or p. |
| 2 | ron=1.0 Ω | Ohmic resistance at zero bias. |
| 3 | rsat=1.0 Ω | Space charge resistance at zero bias. |
| 4 | vsat=10.0 V | Critical drain-source voltage for hot carriers. |
| 5 | psat=1.0 | Velocity saturation coefficient. |
| 6 | vp=-1.0 V | Pinch off voltage at zero gate and substrate voltages. |
| 7 | tox=-1.0 cm | Gate oxide thickness. |
| 8 | dch=1.0e15 cm^{-3} | Doping level channel. |
| 9 | dsub=1.0e15 cm^{-3} | Doping level substrate. |
| 10 | vsub=0.6 V | Substrate diffusion voltage. |
| 11 | vgap=1.2 V | Bandgap voltage channel. |

Spectre Circuit Simulator Reference

Component Statements Part III

12	<code>cgate=0.0 F</code>	Gate capacitance at zero bias.
13	<code>csub=0.0 F</code>	Substrate capacitance at zero bias.
14	<code>tausc=0.0 s</code>	Space charge transit time of the channel.
15	<code>ach=0.0</code>	Temperature coefficient resistivity of the channel.
16	<code>kf=0.0</code>	Flickernoise coefficient.
17	<code>af=1.0</code>	Flickernoise exponent.
18	<code>tr (C)</code>	Reference temperature. Default set by option <code>tnom</code> .
19	<code>trref (C)</code>	Alias of <code>tr</code> . Default set by option <code>tnom</code> .
20	<code>tnom (C)</code>	Alias of <code>tr</code> . Default set by option <code>tnom</code> .
21	<code>dta=0.0 K</code>	Temperature offset of the device.
22	<code>trise=0.0 K</code>	Alias of <code>dta</code> .

Output Parameters

1	<code>ront (Ω)</code>	Ohmic resistance at zero bias.
2	<code>rsat (Ω)</code>	Space charge resistance at zero bias.
3	<code>vsatt (V)</code>	Critical drain-source voltage for hot carriers.
4	<code>vsubt (V)</code>	Substrate diffusion voltage.
5	<code>cgate (F)</code>	Gate capacitance at zero bias.
6	<code>csubt (F)</code>	Substrate capacitance at zero bias.

Operating-Point Parameters

1	<code>pwr (W)</code>	Power.
2	<code>ids (A)</code>	Total current including velocity saturation.

Spectre Circuit Simulator Reference

Component Statements Part III

3	qb (Coul)	Substrate charge.
4	qg (Coul)	Gate charge.
5	qds (Coul)	Space charge in the channel.
6	gdsd (S)	Conductance (d ids / d vd).
7	gdsg (S)	Conductance (d ids / d vg).
8	gdss (S)	Conductance (d ids / d vs).
9	gdsb (S)	Conductance (d ids / d vb).
10	cbd (F)	Capacitance (d qb / d vd).
11	cbg (F)	Capacitance (d qb / d vg).
12	cbs (F)	Capacitance (d qb / d vs).
13	cbb (F)	Capacitance (d qb / d vb).
14	cgd (F)	Capacitance (d qg / d vd).
15	cgg (F)	Capacitance (d qg / d vg).
16	cgs (F)	Capacitance (d qg / d vs).
17	cgb (F)	Capacitance (d qg / d vb).
18	cdsd (F)	Capacitance (d qds / d vd).
19	cdsg (F)	Capacitance (d qds / d vg).
20	cdss (F)	Capacitance (d qds / d vs).
21	cdsb (F)	Capacitance (d qds / d vb).

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the

Spectre Circuit Simulator Reference Component Statements Part III

description for that parameter. For example, a reference of M-35 means the 35th model parameter.

ach	M-15	cgd	OP-14	m	I-4	tox	M-7
af	M-17	cgg	OP-15	mult	I-1	tr	M-18
area	I-2	cgs	OP-16	psat	M-5	tref	M-19
cbb	OP-13	csub	M-13	pwr	OP-1	trise	M-22
cbd	OP-10	csubt	O-6	qb	OP-3	type	M-1
cbg	OP-11	dch	M-8	qds	OP-5	vgap	M-11
cbs	OP-12	dsub	M-9	qg	OP-4	vp	M-6
cdsb	OP-21	dta	M-21	region	I-3	vsat	M-4
cdsd	OP-18	gdsb	OP-9	ron	M-2	vsatt	O-3
cdsg	OP-19	gdsd	OP-6	ront	O-1	vsub	M-10
cdss	OP-20	gdsg	OP-7	rsat	O-2	vsubt	O-4
cgate	O-5	gdss	OP-8	rsat	M-3		
cgate	M-12	ids	OP-2	tausc	M-14		
cgb	OP-17	kf	M-16	tnom	M-20		

Long Channel JFET/MOSFET Model (mos3100)

Description

This long channel JFET/MOSFET model is special developed to describe the drift region of LDMOS, EPMOS and VDMOS devices. It is described in the Philips MOST Modelbook (Dec.98) as MOS model, level 3002 (Used for DMOS). Information on how to obtain this document can be found on Source Link by searching for Philips.

Note: In noise analysis, mos3100 instances will not generate any contribution, since there are no noise sources included in the mos3100 model.

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This device is supported within altergroups.

This device is dynamically loaded from the shared object /vobs/spectre_dev/tools.sun4v/spectre/lib/cmi/3.0.doc/libphilips_sh.so

Spectre Circuit Simulator Reference

Component Statements Part III

Instance Definition

Name d g s [b] ModelName parameter=value ...

Instance Parameters

- | | | |
|---|---------------|--|
| 1 | mult=1 | Number of devices in parallel. |
| 2 | area=1 | Alias of mult. |
| 3 | region=triode | Estimated DC operating region, used as a convergence aid.
Possible values are off, triode, sat, or subth. |
| 4 | m=1 | Multiplicity factor. |

Model Definition

model modelName mos3100 parameter=value ...

Model Parameters

- | | | |
|----|-----------------------------|--|
| 1 | type=n | Transistor gender.
Possible values are n or p. |
| 2 | ron=1.0 Ω | Ohmic resistance at zero bias. |
| 3 | rsat=1.0 Ω | Space charge resistance at zero bias. |
| 4 | vsat=10.0 V | Critical drain-source voltage for hot carriers. |
| 5 | psat=1.0 | Velocity saturation coefficient. |
| 6 | vp=-1.0 V | Pinch off voltage at zero gate and substrate voltages. |
| 7 | tox=-1.0 m | Gate oxide thickness. |
| 8 | dch=1.0e21 m ⁻³ | Doping level channel. |
| 9 | dsub=1.0e21 m ⁻³ | Doping level substrate. |
| 10 | vsub=0.6 V | Substrate diffusion voltage. |
| 11 | vgap=1.2 V | Bandgap voltage channel. |

Spectre Circuit Simulator Reference

Component Statements Part III

12	<code>cgate=0.0 F</code>	Gate capacitance at zero bias.
13	<code>csub=0.0 F</code>	Substrate capacitance at zero bias.
14	<code>tausc=0.0 s</code>	Space charge transit time of the channel.
15	<code>ach=0.0</code>	Temperature coefficient resistivity of the channel.
16	<code>tr (C)</code>	Reference temperature. Default set by option <code>tnom</code> .
17	<code>tréf (C)</code>	Alias of <code>tr</code> . Default set by option <code>tnom</code> .
18	<code>tnom (C)</code>	Alias of <code>tr</code> . Default set by option <code>tnom</code> .
19	<code>dta=0.0 K</code>	Temperature offset of the device.
20	<code>trise=0.0 K</code>	Alias of <code>dta</code> .

Output Parameters

1	<code>r_{ont} (Ω)</code>	Ohmic resistance at zero bias.
2	<code>r_{sat} (Ω)</code>	Space charge resistance at zero bias.
3	<code>v_{satt} (V)</code>	Critical drain-source voltage for hot carriers.
4	<code>v_{subt} (V)</code>	Substrate diffusion voltage.
5	<code>c_{gate} (F)</code>	Gate capacitance at zero bias.
6	<code>c_{subt} (F)</code>	Substrate capacitance at zero bias.

Operating-Point Parameters

1	<code>pwr (W)</code>	Power.
2	<code>ids (A)</code>	Total current including velocity saturation.
3	<code>qb (Coul)</code>	Substrate charge.
4	<code>qg (Coul)</code>	Gate charge.

Spectre Circuit Simulator Reference

Component Statements Part III

5	qds (Coul)	Space charge in the channel.
6	$gdsd$ (S)	Conductance ($d\ i_{ds} / d\ v_d$).
7	$gdsg$ (S)	Conductance ($d\ i_{ds} / d\ v_g$).
8	$gdss$ (S)	Conductance ($d\ i_{ds} / d\ v_s$).
9	$gdsb$ (S)	Conductance ($d\ i_{ds} / d\ v_b$).
10	cbd (F)	Capacitance ($d\ q_b / d\ v_d$).
11	cbg (F)	Capacitance ($d\ q_b / d\ v_g$).
12	cbs (F)	Capacitance ($d\ q_b / d\ v_s$).
13	cbb (F)	Capacitance ($d\ q_b / d\ v_b$).
14	cgd (F)	Capacitance ($d\ q_g / d\ v_d$).
15	cgg (F)	Capacitance ($d\ q_g / d\ v_g$).
16	cgs (F)	Capacitance ($d\ q_g / d\ v_s$).
17	cgb (F)	Capacitance ($d\ q_g / d\ v_b$).
18	$cdsd$ (F)	Capacitance ($d\ q_{ds} / d\ v_d$).
19	$cdsg$ (F)	Capacitance ($d\ q_{ds} / d\ v_g$).
20	$cdss$ (F)	Capacitance ($d\ q_{ds} / d\ v_s$).
21	$cdsb$ (F)	Capacitance ($d\ q_{ds} / d\ v_b$).

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the

Spectre Circuit Simulator Reference Component Statements Part III

description for that parameter. For example, a reference of M-35 means the 35th model parameter.

ach	M-15	cgd	OP-14	m	I-4	tnom	M-18
area	I-2	cgg	OP-15	mult	I-1	tox	M-7
cbb	OP-13	cgs	OP-16	psat	M-5	tr	M-16
cbd	OP-10	csb	M-13	pwr	OP-1	tref	M-17
cbg	OP-11	csbt	O-6	qb	OP-3	trise	M-20
cbs	OP-12	dch	M-8	qds	OP-5	type	M-1
cdsb	OP-21	dsub	M-9	qg	OP-4	vgap	M-11
cdsd	OP-18	dta	M-19	region	I-3	vp	M-6
cdsg	OP-19	gdsb	OP-9	ron	M-2	vsat	M-4
cdss	OP-20	gdsd	OP-6	ront	O-1	vsatt	O-3
cgate	O-5	gdsg	OP-7	rsat	O-2	vsub	M-10
cgate	M-12	gdss	OP-8	rsat	M-3	vsubt	O-4
cgb	OP-17	ids	OP-2	tausc	M-14		

Silicon On Isolator JFET Model (mos40)

Description

This long channel JFET/MOSFET model is special developed to describe the drift region of LDMOS, EPMOS and VDMOS devices. It is described in the Philips MOST Modelbook (Dec.98) as MOS model, level 3002 (Used for DMOS). Information on how to obtain this document can be found on Source Link by searching for Philips.

Note: In noise analysis, mos40 instances will not generate any contribution, since there are no noise sources included in the mos40 model.

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This device is supported within altergroups.

This device is dynamically loaded from the shared object /vobs/spectre_dev/tools.sun4v/spectre/lib/cmi/3.0.doc/libphilips_sh.so

Spectre Circuit Simulator Reference

Component Statements Part III

Instance Definition

Name d g s [b] ModelName parameter=value ...

Instance Parameters

- | | | |
|---|---------------|--|
| 1 | mult=1 | Number of devices in parallel. |
| 2 | area=1 | Alias of mult. |
| 3 | region=triode | Estimated DC operating region, used as a convergence aid.
Possible values are off, triode, sat, or subth. |
| 4 | m=1 | Multiplicity factor. |

Model Definition

model modelName mos40 parameter=value ...

Model Parameters

- | | | |
|----|----------------------------|--|
| 1 | type=n | Transistor gender.
Possible values are n or p. |
| 2 | ron=1.0 Ω | Ohmic resistance at zero bias. |
| 3 | rsat=1.0 Ω | Space charge resistance at zero bias. |
| 4 | vsat=10.0 V | Critical drain-source voltage for hot carriers. |
| 5 | psat=1.0 | Velocity saturation coefficient. |
| 6 | vp=-1.0 V | Pinch off voltage at zero gate and substrate voltages. |
| 7 | tox=-1.0 m | Gate oxide thickness. |
| 8 | dch=1.0e21 m ⁻³ | Doping level channel. |
| 9 | tbox=-1.0 m ⁻³ | Box thicknes. |
| 10 | cbox=0.0 m ⁻³ | Wafer capacitance. |
| 11 | cgate=0.0 F | Gate capacitance at zero bias. |

Spectre Circuit Simulator Reference

Component Statements Part III

12	<code>tausc=0.0 s</code>	Space charge transit time of the channel.
13	<code>ach=0.0</code>	Temperature coefficient resistivity of the channel.
14	<code>tr (C)</code>	Reference temperature. Default set by option <code>tnom</code> .
15	<code>tref (C)</code>	Alias of <code>tr</code> . Default set by option <code>tnom</code> .
16	<code>tnom (C)</code>	Alias of <code>tr</code> . Default set by option <code>tnom</code> .
17	<code>dta=0.0 K</code>	Temperature offset of the device.
18	<code>trise=0.0 K</code>	Alias of <code>dta</code> .

Output Parameters

1	<code>ront (Ω)</code>	Ohmic resistance at zero bias.
2	<code>rsat (Ω)</code>	Space charge resistance at zero bias.
3	<code>vsatt (V)</code>	Critical drain-source voltage for hot carriers.
4	<code>vbox (V)</code>	Box voltage.
5	<code>cgate (F)</code>	Gate capacitance at zero bias.

Operating-Point Parameters

1	<code>pwr (W)</code>	Power.
2	<code>ids (A)</code>	Total current including velocity saturation.
3	<code>qb (Coul)</code>	Substrate charge.
4	<code>qg (Coul)</code>	Gate charge.
5	<code>qds (Coul)</code>	Space charge in the channel.
6	<code>gdSD (S)</code>	Conductance ($d\text{ ids} / d\text{ vd}$).
7	<code>gdsg (S)</code>	Conductance ($d\text{ ids} / d\text{ vg}$).

Spectre Circuit Simulator Reference

Component Statements Part III

8	gdss (S)	Conductance (d ids / d vs).
9	gdsb (S)	Conductance (d ids / d vb).
10	cbd (F)	Capacitance (d qb / d vd).
11	cbg (F)	Capacitance (d qb / d vg).
12	cbs (F)	Capacitance (d qb / d vs).
13	cbb (F)	Capacitance (d qb / d vb).
14	cgd (F)	Capacitance (d qg / d vd).
15	cgg (F)	Capacitance (d qg / d vg).
16	cgs (F)	Capacitance (d qg / d vs).
17	cgb (F)	Capacitance (d qg / d vb).
18	cdsd (F)	Capacitance (d qds / d vd).
19	cdsg (F)	Capacitance (d qds / d vg).
20	cdss (F)	Capacitance (d qds / d vs).
21	cdsb (F)	Capacitance (d qds / d vb).

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

ach	M-13	cgate	M-11	m	I-4	tausc	M-12
area	I-2	cgb	OP-17	mult	I-1	tbox	M-9
cbb	OP-13	cgd	OP-14	psat	M-5	tnom	M-16
cbd	OP-10	cgg	OP-15	pwr	OP-1	tox	M-7

Spectre Circuit Simulator Reference Component Statements Part III

cbg	OP-11	cgs	OP-16	qb	OP-3	tr	M-14
cbox	M-10	dch	M-8	qds	OP-5	tref	M-15
cbs	OP-12	dta	M-17	qg	OP-4	trise	M-18
cdsb	OP-21	gdsb	OP-9	region	I-3	type	M-1
cdsd	OP-18	gdsc	OP-6	ron	M-2	vbox	O-4
cdsg	OP-19	gdsg	OP-7	ront	O-1	vp	M-6
cdss	OP-20	gdss	OP-8	rsat	O-2	vsat	M-4
cgate	O-5	ids	OP-2	rsat	M-3	vsatt	O-3

Compact MOS-Transistor Model (mos705)

Description

The mos705 model is a compact MOS-transistor model, intended for the simulation of circuit behavior with emphasis on analog applications. It is described in the Philips MOST Modelbook (Dec.93) as MOS model, level 705.

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In extension to the modelbook description a minimum conductance g_{min} is inserted between the drain and source node, to aid convergence. The value of g_{min} is set by an options statement, default = 1e-12 S.

This device is supported within altergroups.

This device is dynamically loaded from the shared object /vobs/spectre_dev/tools.sun4v/spectre/lib/cmi/3.0.doc/libphilips_sh.so

Sample Instance Statement

```
mn1 (1 2 0 0) mna7 ln=120e-6 wn=12e-6
```

Sample Model Statement

```
model mna7 mos705 type=n vtn=0.853 betan=77e-6 tox=15e-9 vfb=-850e-3 tref=25  
subthn=3 phi=0.645 lap=100e-9 gkn=-350e-9 th1n=0.15 th2n=0.046 th3n=0.1 fnoise=1e-  
10
```

Spectre Circuit Simulator Reference

Component Statements Part III

Instance Definition

Name d g s [b] ModelName parameter=value ...

Instance Parameters

- | | | |
|---|----------------|---|
| 1 | wn=1.0 scale m | Drawn channel width in the lay-out of the actual transistor. Scale set by option scale. |
| 2 | ln=1.0 scale m | Drawn channel length in the lay-out of the actual transistor. Scale set by option scale. |
| 3 | w=1.0 scale m | Alias for wn. |
| 4 | l=1.0 scale m | Alias for ln. |
| 5 | mult=1 | Number of devices in parallel. |
| 6 | area=1 | Alias of mult. |
| 7 | region=triode | Estimated DC operating region, used as a convergence aid. Possible values are off, triode, sat, or subth. |
| 8 | m=1 | Multiplicity factor. |

Model Definition

model modelName mos705 parameter=value ...

Model Parameters

- | | | |
|---|--------------------|---|
| 1 | type=n | Transistor gender. Possible values are n or p. |
| 2 | vtn=0 V | Threshold voltage of the reference transistor at the reference temperature. |
| 3 | kon=0 \sqrt{V} | K_0 of the reference transistor. |
| 4 | kn=100m \sqrt{V} | K of the reference transistor. |
| 5 | vsbxn=0 V | V_{sbx} of the reference transistor. |

Spectre Circuit Simulator Reference

Component Statements Part III

6	$d_{elvx}=0$ V	D_{vsbx} of the reference transistor.
7	$th_{1n}=0$ 1/V	The_1 of the reference transistor.
8	$th_{2n}=0$ 1/ \sqrt{V}	The_2 of the reference transistor.
9	$th_{3n}=0$ 1/V	The_3 of the reference transistor at the reference temperature.
10	$gamman=0$	G_{am} of the reference transistor.
11	$shiftn=0$ $V^{(1-n)}$	Sh of the reference transistor.
12	$nn=0$	N of the reference transistor.
13	$pn=0$ 1/V	P of the reference transistor.
14	$ava=0$	A of the reference transistor.
15	$avb=1$ V	B of the reference transistor.
16	$avc=0$	C of the reference transistor.
17	$w_{ref}=100u$ m	Effective width of the reference transistor.
18	$wtol=0$ m	Difference between drawn and effective gate width.
19	$dvtn=0$ V m	Narrow-width factor of the threshold voltage at v_{sbref} .
20	$dkon=0$ \sqrt{V} m	Narrow-width factor of k_0 .
21	$dkn=0$ \sqrt{V} m	Narrow-width factor of k .
22	$dvsbxn=0$ V m	Narrow-width factor of v_{sbx} .
23	$ddelvx=0$ Vm	Narrow-width factor of d_{vsbx} .
24	$betan=20u$ A/ V^2	Gain factor of a infinite-square transistor at the reference temperature.
25	$dth_{1n}=0$ m/V	Narrow-width factor of th_{e1} .
26	$dth_{2n}=0$ m/ \sqrt{V}	Narrow-width factor of th_{e2} .

Spectre Circuit Simulator Reference

Component Statements Part III

27	$dth3n=0$ m/V	Narrow-width factor of $the3$.
28	$dgamn=0$ m	Narrow-width factor of gam .
29	$dava=0$ m	Narrow-width factor of a .
30	$davb=0$ V m	Narrow-width factor of b .
31	$davc=0$ m	Narrow-width factor of c .
32	$lref=100u$ m	Effective length of the reference transistor.
33	$ltol=0$ m	Difference between drawn and actual gate polysilicon length.
34	$gvtn=0$ V m	Short-channel factor of the threshold voltage at $vsbref$.
35	$gkon=0$ \sqrt{V} m	Short-channel factor of ko .
36	$gkn=0$ \sqrt{V} m	Short-channel factor of k .
37	$gvsbxn=0$ V m	Short-channel factor of $vsbx$.
38	$gdelvx=0$ V m	Short-channel factor of $dvsbx$.
39	$gth1n=0$ m/V	Short-channel factor of $the1$.
40	$gth2n=0$ m/ \sqrt{V}	Short-channel factor of $the2$.
41	$gth3n=0$ m/V	Short-channel factor of $the3$.
42	$ggamn=0$ m	Short-channel factor of gam .
43	$gshift=0$ $V^{(1-n)}$ m^2	Short-channel factor of sh .
44	$gnn=0$ m	Short-channel factor of n .
45	$gpn=0$ m/V	Short-channel factor of p .
46	$gava=0$ m	Short-channel factor of a .
47	$gavb=0$ V m	Short-channel factor of b .

Spectre Circuit Simulator Reference

Component Statements Part III

48	$g_{avc}=0$	m	Short-channel factor of c .
49	$l_{ap}=0$	m	Half of the effective channel-length reduction due to lateral diffusion.
50	$v_{sbref}=0$	V	Source to bulk reference voltage for parameter determination.
51	$\phi_i=600$	m V	Diffusion potential at the reference temperature.
52	$t_{cvt}=-1$	m V/K	Temperature coefficient of v_{t0} .
53	$t_{betan}=1.5$		Power temperature coefficient of β_{et} .
54	$t_{th3n}=0$	1/(V K)	Temperature coefficient of t_{he3} .
55	$t_{gth3n}=0$	m/(V K)	Temperature coefficient of the length dependence of t_{he3} .
56	$m=1.0$		Subthreshold-slope factor at reference back bias and at the reference temperature.
57	$subthn=0$		Weak-inversion factor.
58	$v_{tr}=0$	V	Depletion-MOS-transistor-transition voltage.
59	$ratio=0$		Depletion-MOS-transistor-gain ratio.
60	$v_{fb}=0$	V	Flat-band voltage.
61	$t_{ox}=100$	n m	Gate-oxide thickness.
62	$c_{ol}=0$	F/m	Gate/drain or gate/source overlap capacitance per unit length.
63	$f_{noise}=0$	$m^2 V^2$	Flicker-noise factor.
64	$t_{noise}=0$		Thermal-noise factor.

Temperature parameters

65	t_r	(C)	Reference temperature. Default set by option t_{nom} .
66	t_{ref}	(C)	Alias of t_r . Default set by option t_{nom} .

Spectre Circuit Simulator Reference

Component Statements Part III

67	<code>tnom (C)</code>	Alias of <code>tr</code> . Default set by option <code>tnom</code> .
68	<code>dta=0 K</code>	Deviation between the temperature of the transistor and the temperature of the circuit.
69	<code>trise=0 K</code>	Alias of <code>dta</code> .

Output Parameters

1	<code>weff (V)</code>	Effective channel width of the actual transistor.
2	<code>leff (V)</code>	Effective channel length of the actual transistor.
3	<code>twophif (V)</code>	Diffusion potential.
4	<code>bet (A/V²)</code>	Gain factor of the transistor.
5	<code>k (\sqrt{V})</code>	Body-effect factor.
6	<code>ko (\sqrt{V})</code>	Initial body-effect factor for dual <code>k</code> approach.
7	<code>vsbx (V)</code>	Transition voltage for dual <code>k</code> approach.
8	<code>dvsbx (V)</code>	Transition-voltage range for dual <code>k</code> approach.
9	<code>vto (V)</code>	Threshold voltage.
10	<code>von (V)</code>	Onset voltage of the superthreshold region.
11	<code>the1 (1/V)</code>	Gate-bias-controlled transverse-field mobility reduction factor.
12	<code>the2 (1/\sqrt{V})</code>	Back-bias-controlled transverse-field mobility reduction factor.
13	<code>the3 (1/V)</code>	Lateral-field mobility reduction factor (velocity saturation).
14	<code>gam</code>	Static-drain-feedback factor.
15	<code>sh (V⁽¹⁻ⁿ⁾)</code>	Threshold-voltage-shift factor.
16	<code>n</code>	Threshold-voltage-shift exponent.
17	<code>p (1/V)</code>	Back-bias-shift factor.

Spectre Circuit Simulator Reference

Component Statements Part III

18	m_e (\sqrt{V})	Auxiliary parameter for subthreshold-slope factor.
19	a	Weak-avalanche multiplier.
20	b (V)	Weak-avalanche exponent factor.
21	c	Saturation-voltage reduction factor.
22	c_{ox} (F)	Gate capacitance.
23	c_{gso} (F)	Gate/source-overlap capacitance.
24	c_{gdo} (F)	Gate/drain-overlap capacitance.
25	v_{tre} (V)	Depletion MOS transistor transition voltage.
26	ratio	Depletion MOS transistor gain ratio.
27	v_{fbe} (V)	Flat band voltage.
28	v_{temp} (V)	kT/q at actual device temperature.
29	g_{noise} (V^2)	Coefficient of the flicker noise for the actual transistor.
30	u_{noise} (J)	Coefficient of the thermal noise for the actual transistor.

Operating-Point Parameters

1	i_{de} (A)	Drain current.
2	i_{ge} (A)	Gate current.
3	i_{se} (A)	Source current.
4	i_{be} (A)	Bulk current.
5	v_{ds} (V)	Drain-source voltage.
6	v_{gs} (V)	Gate-source voltage.
7	v_{sb} (V)	Source-bulk voltage.
8	i_{ds} (A)	Drain-source current.

Spectre Circuit Simulator Reference

Component Statements Part III

9	i_{db} (A)	Drain-bulk current.
10	i_{sb} (A)	Source-bulk current.
11	pwr (W)	Power.
12	v_{ts} (V)	V_{to} including back-bias effects.
13	v_{gt} (V)	Effective gate drive including back-bias and drain effects.
14	v_{dss} (V)	Saturation voltage at actual bias.
15	g_m (S)	Transconductance ($d i_{ds} / d v_{gs}$).
16	g_{mb} (S)	Bulk transconductance ($d i_{ds} / d v_{bs}$).
17	g_{ds} (S)	Output conductance ($d i_{ds} / d v_{ds}$).
18	c_{dd} (F)	Capacitance ($d q_d / d v_d$).
19	c_{dg} (F)	Capacitance ($- d q_d / d v_g$).
20	c_{ds} (F)	Capacitance ($- d q_d / d v_s$).
21	c_{db} (F)	Capacitance ($- d q_d / d v_b$).
22	c_{gd} (F)	Capacitance ($- d q_g / d v_d$).
23	c_{gg} (F)	Capacitance ($d q_g / d v_g$).
24	c_{gs} (F)	Capacitance ($- d q_g / d v_s$).
25	c_{gb} (F)	Capacitance ($- d q_g / d v_b$).
26	c_{sd} (F)	Capacitance ($- d q_s / d v_d$).
27	c_{sg} (F)	Capacitance ($- d q_s / d v_g$).
28	c_{ss} (F)	Capacitance ($d q_s / d v_s$).
29	c_{sb} (F)	Capacitance ($- d q_s / d v_b$).
30	c_{bd} (F)	Capacitance ($- d q_b / d v_d$).

Spectre Circuit Simulator Reference

Component Statements Part III

31	cbg (F)	Capacitance (- d qb / d vg).
32	cbs (F)	Capacitance (- d qb / d vs).
33	cbb (F)	Capacitance (d qb / d vb).
34	u	Transistor gain (gm/gds).
35	rout (Ω)	Small signal output resistance (1/gds).
36	vearly (V)	Equivalent Early voltage ($ I_d /gds$).
37	keff (\sqrt{V})	Describes body effect at actual bias.
38	beff (S/V)	Effective beta at actual bias in the simple MOS model.
39	fug (Hz)	Unity gain frequency at actual bias ($gm/(2*\pi*cox)$).
40	sqrtsw (V/ \sqrt{Hz})	Input-referred RMS white noise voltage (\sqrt{sth}/gm).
41	sqrtfff (V/ \sqrt{Hz})	Input-referred RMS 1/f noise voltage at 1kHz ($\sqrt{gnoise/1000}$).
42	fknee (Hz)	Cross-over frequency above which white noise is dominant.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

a	O-19	dta	M-68	k	O-5	the3	O-13
area	I-6	dth1n	M-25	keff	OP-37	tnoise	M-64
ava	M-14	dth2n	M-26	kn	M-4	tnom	M-67
avb	M-15	dth3n	M-27	ko	O-6	tox	M-61
avc	M-16	dvsbx	O-8	kon	M-3	tr	M-65

Spectre Circuit Simulator Reference Component Statements Part III

b	O-20	dvsbxn	M-22	l	I-4	tref	M-66
beff	OP-38	dvtm	M-19	lap	M-49	trise	M-69
bet	O-4	fknee	OP-42	leff	O-2	tth3n	M-54
betan	M-24	fnoise	M-63	ln	I-2	twophif	O-3
c	O-21	fug	OP-39	lref	M-32	type	M-1
cbb	OP-33	gam	O-14	ltol	M-33	u	OP-34
cbd	OP-30	gamman	M-10	m	M-56	unoise	O-30
cbg	OP-31	gava	M-46	m	I-8	vds	OP-5
cbs	OP-32	gavb	M-47	me	O-18	vdss	OP-14
cdb	OP-21	gavc	M-48	mult	I-5	vearly	OP-36
cdd	OP-18	gdelvx	M-38	n	O-16	vfb	M-60
cdg	OP-19	gds	OP-17	nn	M-12	vfbe	O-27
cds	OP-20	ggamn	M-42	p	O-17	vgs	OP-6
cgb	OP-25	gkn	M-36	phi	M-51	vgt	OP-13
cgd	OP-22	gkon	M-35	pn	M-13	von	O-10
cgdo	O-24	gm	OP-15	pwr	OP-11	vsb	OP-7
cgg	OP-23	gmb	OP-16	ratio	M-59	vsbref	M-50
cgs	OP-24	gmn	M-44	ratio	O-26	vsbx	O-7
cgso	O-23	gnoise	O-29	region	I-7	vsbxn	M-5
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cox	O-22	gshift	M-43	sh	O-15	vtn	M-2
csb	OP-29	gth1n	M-39	shiftm	M-11	vto	O-9
csd	OP-26	gth2n	M-40	sqrtsff	OP-41	vtr	M-58
csq	OP-27	gth3n	M-41	sqrtsfw	OP-40	vtre	O-25
css	OP-28	gvsbxn	M-37	subthn	M-57	vts	OP-12
dava	M-29	gvtm	M-34	tbetan	M-53	w	I-3
davb	M-30	ibe	OP-4	tcvt	M-52	weff	O-1
davc	M-31	idb	OP-9	tgth3n	M-55	wn	I-1
ddelvx	M-23	ide	OP-1	th1n	M-7	wref	M-17
delvx	M-6	ids	OP-8	th2n	M-8	wtol	M-18
dgamn	M-28	ige	OP-2	th3n	M-9		
dkn	M-21	isb	OP-10	thel	O-11		

dkon M-20

ise OP-3

the2 O-12

Compact MOS-Transistor Model (mos902)

Description

The mos902 model is a compact MOS-transistor model, intended for the simulation of circuit behavior with emphasis on analog applications. It is described in the Philips MOST Modelbook (Feb.98) as MOS model, level 902. Information on how to obtain this document can be found on Source Link by searching for Philips.

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In extension to the modelbook description a minimum conductance `gmin` is inserted between the drain and source node, to aid convergence. The value of `gmin` is set by an options statement, default = 1e-12 S.

This device is supported within altergroups.

This device is dynamically loaded from the shared object `/vobs/spectre_dev/tools.sun4v/spectre/lib/cmi/3.0.doc/libphilips_sh.so`

Sample Instance Statement

```
mp1 (0 1 2 2) mos9pch w=10u l=2u area=1.5
```

Sample Model Statement

```
model mos9pch mos902 ler=0.93e-6 wer=20e-6 tref=27 vtor=1.11 kr=0.54 phibr=0.66  
vsbxxr=100 thelr=0.19 slk=-0.215e-6 swk=98e-9 swthe3=7.8e-9
```

Instance Definition

```
Name d g s [b] ModelName parameter=value ...
```

Instance Parameters

- | | | | |
|---|---------------------|----------------------|---|
| 1 | <code>w=1.0</code> | <code>scale m</code> | Drawn channel width in the lay-out. Scale set by option scale. |
| 2 | <code>l=1.0</code> | <code>scale m</code> | Drawn channel length in the lay-out. Scale set by option scale. |
| 3 | <code>mult=1</code> | | Number of devices in parallel. |

Spectre Circuit Simulator Reference

Component Statements Part III

- | | | |
|---|----------------------------|---|
| 4 | <code>area=1</code> | Alias of <code>mult</code> . |
| 5 | <code>region=triode</code> | Estimated DC operating region, used as a convergence aid. Possible values are <code>off</code> , <code>triode</code> , <code>sat</code> , or <code>subth</code> . |
| 6 | <code>m=1</code> | Multiplicity factor. |

Model Definition

```
model modelName mos902 parameter=value ...
```

Model Parameters

Device type parameters

- | | | |
|---|---------------------|---|
| 1 | <code>type=n</code> | Transistor gender. Possible values are <code>n</code> or <code>p</code> . |
|---|---------------------|---|

Geometry parameters

- | | | |
|---|----------------------------|--|
| 2 | <code>ler=2.5e-6 m</code> | Effective channel length of the reference transistor. |
| 3 | <code>wer=25e-6 m</code> | Effective channel width of the reference transistor. |
| 4 | <code>lvar=0.3e-6 m</code> | Difference between the actual and the programmed poly-silicon gate length. |
| 5 | <code>lap=0.1e-6 m</code> | Effective channel length reduction per side. |
| 6 | <code>wvar=3e-6 m</code> | Difference between the actual and the programmed field-oxide opening. |
| 7 | <code>wot=1e-6 m</code> | Effective channel width reduction per side. |
| 8 | <code>wdog=0 m</code> | Characteristic drawn gate width, below which dogboning appears. |

Threshold-voltage parameters

- | | | |
|---|-------------------------|--------------------------------------|
| 9 | <code>vtor=0.8 V</code> | Threshold voltage at zero back-bias. |
|---|-------------------------|--------------------------------------|

Spectre Circuit Simulator Reference

Component Statements Part III

- 10 $stvto=0.01 \text{ V/K}$ Coefficient of the temperature dependence of v_{t0} .
- 11 $slvto=0.5e-6 \text{ V m}$ Coefficient of the length dependence of v_{t0} .
- 12 $sl2vto=0 \text{ V m}^2$ Second coefficient of the length dependence of v_{t0} .
- 13 $swvto=5e-6 \text{ V m}$ Coefficient of the width dependence of v_{t0} .
- 14 $kor=0.5 \sqrt{V}$ Low-backbias body factor.
- 15 $slko=1e-6 \sqrt{V} \text{ m}$ Coefficient of the length dependence of k_0 .
- 16 $swko=10e-6 \sqrt{V} \text{ m}$ Coefficient of the width dependence of k_0 .
- 17 $kr=0.1 \sqrt{V}$ High-backbias body factor.
- 18 $slk=0.5e-6 \sqrt{V} \text{ m}$ Coefficient of the length dependence of k .
- 19 $swk=5e-6 \sqrt{V} \text{ m}$ Coefficient of the width dependence of k .
- 20 $phibr=0.65 \text{ V}$ Surface potential at strong inversion.
- 21 $vsbxr=0.9 \text{ V}$ Transition voltage for the dual-k-factor model.
- 22 $slvsbx=0.5e-6 \text{ V m}$ Coefficient of the length dependence of $vsbx$.
- 23 $swvsbx=5e-6 \text{ V m}$ Coefficient of the width dependence of $vsbx$.

Channel-current parameters

- 24 $betsq=0.1e-3 \text{ A/V}^2$ Gain factor for an infinite square transistor.
- 25 $etabet=0.5$ Exponent of the temperature dependence of the gain factor.
- 26 $thelr=0.05 \text{ 1/V}$ Coefficient of the mobility reduction due to the gate-induced field.

Spectre Circuit Simulator Reference

Component Statements Part III

- 27 $stthe1r=3e-3 \text{ 1/(V K)}$
Coefficient of the temperature dependence of t_{he1} .
- 28 $slthe1r=50e-9 \text{ m/V}$ Coefficient of the length dependence of t_{he1} .
- 29 $stlthe1=5e-9 \text{ m/(V K)}$
Coefficient of the temperature dependence of $slthe1$.
- 30 $swthe1=1e-6 \text{ m/V}$ Coefficient of the width dependence of t_{he1} .
- 31 $fthe1=0$ Coefficient describing the width dependence of t_{he1} for $w < w_{dog}$.
- 32 $the2r=17e-3 \text{ 1/}\sqrt{V}$
Coefficient of the mobility reduction due to the back-bias.
- 33 $stthe2r=0.1e-3 \text{ 1/}(\sqrt{V} \text{ K)}$
Coefficient of the temperature dependence of t_{he2} .
- 34 $slthe2r=5e-9 \text{ m/}\sqrt{V}$
Coefficient of the length dependence of t_{he2} .
- 35 $stlthe2=0.5e-9 \text{ m/}(\sqrt{V} \text{ K)}$
Coefficient of the temperature dependence of $slthe2$.
- 36 $swthe2=0.1e-6 \text{ m/}\sqrt{V}$
Coefficient of the width dependence of t_{he2} .
- 37 $the3r=37e-3 \text{ 1/V}$ Coefficient of the mobility reduction due to the lateral field.
- 38 $stthe3r=0.1e-3 \text{ 1/(V K)}$
Coefficient of the temperature dependence of t_{he3} .
- 39 $slthe3r=5e-9 \text{ m/V}$ Coefficient of the length dependence of t_{he3} .
- 40 $stlthe3=0.5e-9 \text{ m/(V K)}$
Coefficient of the temperature dependence of $slthe3$.
- 41 $swthe3=0.1e-6 \text{ m/V}$ Coefficient of the width dependence of t_{he3} .

Spectre Circuit Simulator Reference

Component Statements Part III

Drain-feedback parameters

- 42 $\text{gam1r}=40\text{e-}3 \text{ V}^{(1-\text{etads})}$ Coefficient for the drain induced threshold shift for large gate drive.
- 43 $\text{slgam1}=0.1\text{e-}6 \text{ V}^{(1-\text{etads})} \text{ m}$ Coefficient of the length dependence of gam1 .
- 44 $\text{swgam1}=1\text{e-}6 \text{ V}^{(1-\text{etads})} \text{ m}$ Coefficient of the width dependence of gam1 .
- 45 $\text{etadsr}=0.6$ Exponent of the v_{ds} dependence of gam1 .
- 46 $\text{alpr}=4\text{e-}3$ Factor of the channel-length modulation.
- 47 $\text{etaalp}=0.5$ Exponent of the length dependence of alp .
- 48 $\text{slalp}=0.14\text{e-}3 \text{ m}^{\text{etaalp}}$ Coefficient of the length dependence of alp .
- 49 $\text{swalp}=0.1\text{e-}6 \text{ m}$ Coefficient of the width dependence of alp .
- 50 $\text{vpr}=0.25 \text{ V}$ Characteristic voltage of the channel-length modulation.

Sub-threshold parameters

- 51 $\text{gamoor}=1.1\text{e-}3$ Coefficient for the drain induced threshold shift at zero gate drive.
- 52 $\text{slgamoo}=10\text{e-}15 \text{ m}^2$ Coefficient of the length dependence of gamoo .
- 53 $\text{etagamr}=2$ Exponent of the back-bias dependence of gamo .
- 54 $\text{mor}=0.3$ Factor for the subthreshold slope.
- 55 $\text{stmo}=0.01 \text{ 1/K}$ Coefficient of the temperature dependence of mo .
- 56 $\text{slmo}=1.4\text{e-}3 \sqrt{\text{m}}$ Coefficient of the length dependence of mo .
- 57 $\text{etamr}=2$ Exponent of the back-bias dependence of m .

Spectre Circuit Simulator Reference

Component Statements Part III

- 58 `zet1r=0.7` Weak-inversion correction factor.
- 59 `etazet=0.5` Exponent of the length dependence of `zet1`.
- 60 `slzet1=0.14e-6 m^etazet`
Coefficient of the length dependence of `zet1`.
- 61 `vsbtr=99 V` Limiting voltage of the `vsb` dependence of `m` and `gamma0`.
- 62 `slvsbt=10e-6 V m` Coefficient of the length dependence of `vsbt`.

Weak-avalanche parameters

- 63 `a1r=22` Factor of the weak-avalanche current.
- 64 `sta1=0.1 1/K` Coefficient of the temperature dependence of `a1`.
- 65 `sla1=10e-6 m` Coefficient of the length dependence of `a1`.
- 66 `swa1=0.1e-3 m` Coefficient of the width dependence of `a1`.
- 67 `a2r=33 V` Exponent of the weak-avalanche current.
- 68 `sla2=10e-6 V m` Coefficient of the length dependence of `a2`.
- 69 `swa2=0.1e-3 V m` Coefficient of the width dependence of `a2`.
- 70 `a3r=0.6` Factor of the drain-source voltage above which weak-avalanche occurs.
- 71 `sla3=1e-6 m` Coefficient of the length dependence of `a3`.
- 72 `swa3=10e-6 m` Coefficient of the width dependence of `a3`.

Charge parameters

- 73 `tox=20e-9 m` Thickness of the oxide layer.
- 74 `col=50e-12 F/m` Gate overlap capacitance per unit channel width.

Spectre Circuit Simulator Reference

Component Statements Part III

Noise parameters

- 75 $n_{tr}=21e-21$ J Coefficient of the thermal noise.
- 76 $n_{fr}=16e-12$ V² Coefficient of the flicker noise.

Temperature parameters

- 77 t_r (C) Reference temperature. Default set by option t_{nom} .
- 78 t_{ref} (C) Alias of t_r . Default set by option t_{nom} .
- 79 t_{nom} (C) Alias of t_r . Default set by option t_{nom} .
- 80 $dta=0$ K Temperature offset of the device.
- 81 $trise=0$ K Alias of dta .

Output Parameters

- 1 l_e (m) Effective channel length.
- 2 w_e (m) Effective channel width.
- 3 v_{to} (V) Threshold voltage at zero back-bias.
- 4 k_o (\sqrt{V}) Low-backbias body factor.
- 5 k (\sqrt{V}) High-backbias body factor.
- 6 ϕ_{ib} (V) Surface potential at strong inversion.
- 7 v_{sbx} (V) Transition voltage for the dual-k-factor model.
- 8 β_{et} (A/V²) Gain factor (* mult).
- 9 $the1$ (1/V) Coefficient of the mobility reduction due to the gate-induced field.
- 10 $the2$ (1/ \sqrt{V}) Coefficient of the mobility reduction due to the back-bias.
- 11 $the3$ (1/V) Coefficient of the mobility reduction due to the lateral field.

Spectre Circuit Simulator Reference

Component Statements Part III

12	gam1 ($V^{(1-\text{etads})}$)	Coefficient for the drain induced threshold shift for large gate drive.
13	etads	Exponent of the vds dependence of gam1 .
14	alp	Factor of the channel-length modulation.
15	vp (V)	Characteristic voltage of the channel-length modulation.
16	gamoo	Coefficient for the drain induced threshold shift at zero gate drive.
17	etagam	Exponent of the back-bias dependence of gamoo .
18	mo	Factor for the subthreshold slope.
19	etam	Exponent of the back-bias dependence of m .
20	phit (V)	Thermal voltage.
21	zet1	Weak-inversion correction factor.
22	vsbt (V)	Limiting voltage of the vsb dependence of m and gamoo .
23	a1	Factor of the weak-avalanche current.
24	a2 (V)	Exponent of the weak-avalanche current.
25	a3	Factor of the drain-source voltage above which weak-avalanche occurs.
26	cox (F)	Gate-to-channel capacitance (* mult).
27	cgdo (F)	Gate-drain overlap capacitance (* mult).
28	cgso (F)	Gate-source overlap capacitance (* mult).
29	nt (J)	Coefficient of the thermal noise.
30	nf (V^2)	Coefficient of the flicker noise (/ mult).

Operating-Point Parameters

1	ide (A)	Resistive drain current.
---	------------------	--------------------------

Spectre Circuit Simulator Reference

Component Statements Part III

2	i_{ge} (A)	Resistive gate current.
3	i_{se} (A)	Resistive source current.
4	i_{be} (A)	Resistive bulk current.
5	v_{ds} (V)	Drain-source voltage.
6	v_{gs} (V)	Gate-source voltage.
7	v_{sb} (V)	Source-bulk voltage.
8	i_{ds} (A)	Resistive drain-source current.
9	i_{db} (A)	Resistive drain-bulk current.
10	i_{sb} (A)	Resistive source-bulk current.
11	i_{avl} (A)	Substrate current.
12	pwr (W)	Power.
13	v_{t1} (V)	V_{t0} including backbias effects.
14	v_{gt2} (V)	Effective gate drive including backbias and drain effects.
15	v_{dss1} (V)	Saturation voltage at actual bias.
16	v_{sat} (V)	Saturation limit.
17	g_m (S)	Transconductance ($d i_{ds} / d v_{gs}$).
18	g_{mb} (S)	Bulk transconductance ($d i_{ds} / d v_{bs}$).
19	g_{ds} (S)	Output conductance ($d i_{ds} / d v_{ds}$).
20	c_{dd} (F)	Capacitance ($d q_d / d v_d$).
21	c_{dg} (F)	Capacitance ($- d q_d / d v_g$).
22	c_{ds} (F)	Capacitance ($- d q_d / d v_s$).
23	c_{db} (F)	Capacitance ($- d q_d / d v_b$).

Spectre Circuit Simulator Reference

Component Statements Part III

24	c_{gd} (F)	Capacitance ($-d qg / d vd$).
25	c_{gg} (F)	Capacitance ($d qg / d vg$).
26	c_{gs} (F)	Capacitance ($-d qg / d vs$).
27	c_{gb} (F)	Capacitance ($-d qg / d vb$).
28	c_{sd} (F)	Capacitance ($-d qs / d vd$).
29	c_{sg} (F)	Capacitance ($-d qs / d vg$).
30	c_{ss} (F)	Capacitance ($d qs / d vs$).
31	c_{sb} (F)	Capacitance ($-d qs / d vb$).
32	c_{bd} (F)	Capacitance ($-d qb / d vd$).
33	c_{bg} (F)	Capacitance ($-d qb / d vg$).
34	c_{bs} (F)	Capacitance ($-d qb / d vs$).
35	c_{bb} (F)	Capacitance ($d qb / d vb$).
36	u	Transistor gain (gm/gds).
37	r_{out} (Ω)	Small signal output resistance ($1/gds$).
38	v_{early} (V)	Equivalent Early voltage ($ id /gds$).
39	k_{eff} (\sqrt{V})	Describes body effect at actual bias.
40	b_{eff} (S/V)	Effective beta at actual bias in the simple MOS model ($2* ids /vgt^2$).
41	f_{ug} (Hz)	Unity gain frequency at actual bias ($gm/(2*\pi*cin)$).
42	$sqrt_{sfw}$ (V/ \sqrt{Hz})	Input-referred RMS white noise voltage ($sqrt(sth)/gm$).
43	$sqrt_{sff}$ (V/ \sqrt{Hz})	Input-referred RMS 1/f noise voltage at 1kHz ($sqrt(nf/1000)$).

Spectre Circuit Simulator Reference Component Statements Part III

44 `fknee` (Hz) Cross-over frequency above which white noise is dominant.

Parameter Index

In the following index, `I` refers to instance parameters, `M` refers to the model parameters section, `O` refers to the output parameters section, and `OP` refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of `M-35` means the 35th model parameter.

<code>a1</code>	<code>O-23</code>	<code>etazet</code>	<code>M-59</code>	<code>region</code>	<code>I-5</code>	<code>swvto</code>	<code>M-13</code>
<code>alr</code>	<code>M-63</code>	<code>fknee</code>	<code>OP-44</code>	<code>rout</code>	<code>OP-37</code>	<code>the1</code>	<code>O-9</code>
<code>a2</code>	<code>O-24</code>	<code>fthe1</code>	<code>M-31</code>	<code>sl2vto</code>	<code>M-12</code>	<code>the1r</code>	<code>M-26</code>
<code>a2r</code>	<code>M-67</code>	<code>fug</code>	<code>OP-41</code>	<code>sla1</code>	<code>M-65</code>	<code>the2</code>	<code>O-10</code>
<code>a3</code>	<code>O-25</code>	<code>gam1</code>	<code>O-12</code>	<code>sla2</code>	<code>M-68</code>	<code>the2r</code>	<code>M-32</code>
<code>a3r</code>	<code>M-70</code>	<code>gam1r</code>	<code>M-42</code>	<code>sla3</code>	<code>M-71</code>	<code>the3</code>	<code>O-11</code>
<code>alp</code>	<code>O-14</code>	<code>gamoo</code>	<code>O-16</code>	<code>slalp</code>	<code>M-48</code>	<code>the3r</code>	<code>M-37</code>
<code>alpr</code>	<code>M-46</code>	<code>gamoor</code>	<code>M-51</code>	<code>slgam1</code>	<code>M-43</code>	<code>tnom</code>	<code>M-79</code>
<code>area</code>	<code>I-4</code>	<code>gds</code>	<code>OP-19</code>	<code>slgamoo</code>	<code>M-52</code>	<code>tox</code>	<code>M-73</code>
<code>beff</code>	<code>OP-40</code>	<code>gm</code>	<code>OP-17</code>	<code>slk</code>	<code>M-18</code>	<code>tr</code>	<code>M-77</code>
<code>bet</code>	<code>O-8</code>	<code>gmb</code>	<code>OP-18</code>	<code>slko</code>	<code>M-15</code>	<code>tref</code>	<code>M-78</code>
<code>betsq</code>	<code>M-24</code>	<code>iavl</code>	<code>OP-11</code>	<code>slmo</code>	<code>M-56</code>	<code>trise</code>	<code>M-81</code>
<code>cbb</code>	<code>OP-35</code>	<code>ibe</code>	<code>OP-4</code>	<code>slthe1r</code>	<code>M-28</code>	<code>type</code>	<code>M-1</code>
<code>cbd</code>	<code>OP-32</code>	<code>idb</code>	<code>OP-9</code>	<code>slthe2r</code>	<code>M-34</code>	<code>u</code>	<code>OP-36</code>
<code>cbg</code>	<code>OP-33</code>	<code>ide</code>	<code>OP-1</code>	<code>slthe3r</code>	<code>M-39</code>	<code>vds</code>	<code>OP-5</code>
<code>cbs</code>	<code>OP-34</code>	<code>ids</code>	<code>OP-8</code>	<code>slvsbt</code>	<code>M-62</code>	<code>vdss1</code>	<code>OP-15</code>
<code>cdb</code>	<code>OP-23</code>	<code>ige</code>	<code>OP-2</code>	<code>slvsbx</code>	<code>M-22</code>	<code>vearly</code>	<code>OP-38</code>
<code>cdd</code>	<code>OP-20</code>	<code>isb</code>	<code>OP-10</code>	<code>slvto</code>	<code>M-11</code>	<code>vgs</code>	<code>OP-6</code>
<code>cdg</code>	<code>OP-21</code>	<code>ise</code>	<code>OP-3</code>	<code>slzet1</code>	<code>M-60</code>	<code>vgt2</code>	<code>OP-14</code>
<code>cds</code>	<code>OP-22</code>	<code>k</code>	<code>O-5</code>	<code>sqrtsff</code>	<code>OP-43</code>	<code>vp</code>	<code>O-15</code>
<code>cgb</code>	<code>OP-27</code>	<code>keff</code>	<code>OP-39</code>	<code>sqrtsfw</code>	<code>OP-42</code>	<code>vpr</code>	<code>M-50</code>
<code>cgd</code>	<code>OP-24</code>	<code>ko</code>	<code>O-4</code>	<code>stal</code>	<code>M-64</code>	<code>vsat</code>	<code>OP-16</code>
<code>cgdo</code>	<code>O-27</code>	<code>kor</code>	<code>M-14</code>	<code>stlthe1</code>	<code>M-29</code>	<code>vsb</code>	<code>OP-7</code>

Spectre Circuit Simulator Reference Component Statements Part III

cgg	OP-25	kr	M-17	stlthe2	M-35	vsbt	O-22
cgs	OP-26	l	I-2	stlthe3	M-40	vsbtr	M-61
cgso	O-28	lap	M-5	stmo	M-55	vsbx	O-7
col	M-74	le	O-1	stthe1r	M-27	vsbxr	M-21
cox	O-26	ler	M-2	stthe2r	M-33	vt1	OP-13
csb	OP-31	lvar	M-4	stthe3r	M-38	vto	O-3
csd	OP-28	m	I-6	stvto	M-10	vtor	M-9
csg	OP-29	mo	O-18	swa1	M-66	w	I-1
css	OP-30	mor	M-54	swa2	M-69	wdog	M-8
dta	M-80	mult	I-3	swa3	M-72	we	O-2
etaalp	M-47	nf	O-30	swalp	M-49	wer	M-3
etabet	M-25	nfr	M-76	swgam1	M-44	wot	M-7
etads	O-13	nt	O-29	swk	M-19	wvar	M-6
etadsr	M-45	ntr	M-75	swko	M-16	zet1	O-21
etagam	O-17	phib	O-6	swthe1	M-30	zet1r	M-58
etagamr	M-53	phibr	M-20	swthe2	M-36		
etam	O-19	phit	O-20	swthe3	M-41		
etamr	M-57	pwr	OP-12	swvsbx	M-23		

Compact MOS-Transistor Model (mos903)

Description

The mos903 model is a compact MOS-transistor model, intended for the simulation of circuit behavior with emphasis on analog applications. It is described in the Philips MOST Modelbook (Jun.98) as MOS model, level 903. Information on how to obtain this document can be found on Source Link by searching for Philips.

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In extension to the modelbook description a minimum conductance g_{min} is inserted between the drain and source node, to aid convergence. The value of g_{min} is set by an options statement, default = 1e-12 S.

This device is supported within altergroups.

Spectre Circuit Simulator Reference

Component Statements Part III

This device is dynamically loaded from the shared object /vobs/spectre_dev/tools.sun4v/spectre/lib/cmi/3.0.doc/libphilips_sh.so

Sample Instance Statement

```
m_1 (1 2 0 0) mos9nch w=0.35e-6 l=0.35e-6
```

Sample Model Statement

```
model mos9nch mos903 uhyylr=3.5e-7 wer=1e-5 lvar=0 lap=2.2e-8 wvar=0 wot=3e-8  
vtor=0.76 thelr=0.67 stthelr=-1.76e-3 etaalp=0 slalp=0 alpr=0.01
```

Instance Definition

```
Name d g s [b] ModelName parameter=value ...
```

Instance Parameters

- | | | |
|---|---------------|--|
| 1 | w=1.0 scale m | Drawn channel width in the lay-out. Scale set by option scale. |
| 2 | l=1.0 scale m | Drawn channel length in the lay-out. Scale set by option scale. |
| 3 | mult=1 | Number of devices in parallel. |
| 4 | area=1 | Alias of mult. |
| 5 | region=triode | Estimated DC operating region, used as a convergence aid.
Possible values are off, triode, sat, or subth. |
| 6 | m=1 | Multiplicity factor. |

Model Definition

```
model modelName mos903 parameter=value ...
```

Model Parameters

Device type parameters

- | | | |
|---|--------|---|
| 1 | type=n | Transistor gender.
Possible values are n or p. |
|---|--------|---|

Spectre Circuit Simulator Reference

Component Statements Part III

Geometry parameters

2	$l_{er}=2.5e-6$ m	Effective channel length of the reference transistor.
3	$w_{er}=25e-6$ m	Effective channel width of the reference transistor.
4	$l_{var}=0.3e-6$ m	Difference between the actual and the programmed poly-silicon gate length.
5	$l_{ap}=0.1e-6$ m	Effective channel length reduction per side.
6	$w_{var}=3e-6$ m	Difference between the actual and the programmed field-oxide opening.
7	$w_{ot}=1e-6$ m	Effective channel width reduction per side.

Threshold-voltage parameters

8	$v_{tor}=0.8$ V	Threshold voltage at zero back-bias.
9	$stv_{to}=0.01$ V/K	Coefficient of the temperature dependence of v_{to} .
10	$slv_{to}=0.5e-6$ V m	Coefficient of the length dependence of v_{to} .
11	$sl2v_{to}=0$ V m ²	Second coefficient of the length dependence of v_{to} .
12	$swv_{to}=5e-6$ V m	Coefficient of the width dependence of v_{to} .
13	$k_{or}=0.5$ \sqrt{V}	Low-backbias body factor.
14	$slk_{o}=1e-6$ \sqrt{V} m	Coefficient of the length dependence of k_{o} .
15	$swk_{o}=10e-6$ \sqrt{V} m	Coefficient of the width dependence of k_{o} .
16	$k_{r}=0.1$ \sqrt{V}	High-backbias body factor.
17	$slk=0.5e-6$ \sqrt{V} m	Coefficient of the length dependence of k .

Spectre Circuit Simulator Reference

Component Statements Part III

- 18 $swk=5e-6 \sqrt{V} \text{ m}$ Coefficient of the width dependence of k .
- 19 $phibr=0.65 \text{ V}$ Surface potential at strong inversion.
- 20 $vsbxr=0.9 \text{ V}$ Transition voltage for the dual- k -factor model.
- 21 $slvsbx=0.5e-6 \text{ V m}$ Coefficient of the length dependence of $vsbx$.
- 22 $swvsbx=5e-6 \text{ V m}$ Coefficient of the width dependence of $vsbx$.

Channel-current parameters

- 23 $betsq=0.1e-3 \text{ A/V}^2$ Gain factor for an infinite square transistor.
- 24 $etabet=0.5$ Exponent of the temperature dependence of the gain factor.
- 25 $thelr=0.05 \text{ 1/V}$ Coefficient of the mobility reduction due to the gate-induced field.
- 26 $stthelr=3e-3 \text{ 1/(V K)}$ Coefficient of the temperature dependence of $thel$.
- 27 $slthelr=50e-9 \text{ m/V}$ Coefficient of the length dependence of $thel$.
- 28 $stlthel=5e-9 \text{ m/(V K)}$ Coefficient of the temperature dependence of $slthel$.
- 29 $swthel=1e-6 \text{ m/V}$ Coefficient of the width dependence of $thel$.
- 30 $wdog=0 \text{ m}$ Characteristic drawn gate width, below which dogboning appears.
- 31 $fthel=0$ Coefficient describing the width dependence of $thel$ for $w < wdog$.
- 32 $the2r=17e-3 \text{ 1/}\sqrt{V}$ Coefficient of the mobility reduction due to the back-bias.
- 33 $stthe2r=0.1e-3 \text{ 1/(\}\sqrt{V} \text{ K)}$ Coefficient of the temperature dependence of $the2$.

Spectre Circuit Simulator Reference

Component Statements Part III

- 34 $slthe2r=5e-9 \text{ m}/\sqrt{V}$ Coefficient of the length dependence of $the2$.
- 35 $stlthe2=0.5e-9 \text{ m}/(\sqrt{V} \text{ K})$ Coefficient of the temperature dependence of $slthe2$.
- 36 $swthe2=0.1e-6 \text{ m}/\sqrt{V}$ Coefficient of the width dependence of $the2$.
- 37 $the3r=37e-3 \text{ 1}/V$ Coefficient of the mobility reduction due to the lateral field.
- 38 $stthe3r=0.1e-3 \text{ 1}/(V \text{ K})$ Coefficient of the temperature dependence of $the3$.
- 39 $slthe3r=5e-9 \text{ m}/V$ Coefficient of the length dependence of $the3$.
- 40 $stlthe3=0.5e-9 \text{ m}/(V \text{ K})$ Coefficient of the temperature dependence of $slthe3$.
- 41 $swthe3=0.1e-6 \text{ m}/V$ Coefficient of the width dependence of $the3$.

Drain-feedback parameters

- 42 $gam1r=40e-3 \text{ V}^{(1-etads)}$ Coefficient for the drain induced threshold shift for large gate drive.
- 43 $slgam1=0.1e-6 \text{ V}^{(1-etads)} \text{ m}$ Coefficient of the length dependence of $gam1$.
- 44 $swgam1=1e-6 \text{ V}^{(1-etads)} \text{ m}$ Coefficient of the width dependence of $gam1$.
- 45 $etadsr=0.6$ Exponent of the vds dependence of $gam1$.
- 46 $alpr=4e-3$ Factor of the channel-length modulation.
- 47 $etaalp=0.5$ Exponent of the length dependence of alp .
- 48 $slalp=0.14e-3 \text{ m}^{etaalp}$ Coefficient of the length dependence of alp .

Spectre Circuit Simulator Reference

Component Statements Part III

- 49 `swalp=0.1e-6 m` Coefficient of the width dependence of `alp`.
- 50 `vpr=0.25 V` Characteristic voltage of the channel-length modulation.

Sub-threshold parameters

- 51 `gamoor=1.1e-3` Coefficient for the drain induced threshold shift at zero gate drive.
- 52 `slgamoo=10e-15 m2` Coefficient of the length dependence of `gamoo`.
- 53 `etagamr=2` Exponent of the back-bias dependence of `gamoo`.
- 54 `mor=0.3` Factor for the subthreshold slope.
- 55 `stmo=0.01 1/K` Coefficient of the temperature dependence of `mo`.
- 56 `slmo=1.4e-3 \sqrt{m}` Coefficient of the length dependence of `mo`.
- 57 `etamr=2` Exponent of the back-bias dependence of `m`.
- 58 `zet1r=0.7` Weak-inversion correction factor.
- 59 `etazet=0.5` Exponent of the length dependence of `zet1`.
- 60 `slzet1=0.14e-6 metazet` Coefficient of the length dependence of `zet1`.
- 61 `vsbtr=99 V` Limiting voltage of the `vsb` dependence of `m` and `gamoo`.
- 62 `slvsbt=10e-6 V m` Coefficient of the length dependence of `vsbt`.

Weak-avalanche parameters

- 63 `a1r=22` Factor of the weak-avalanche current.
- 64 `sta1=0.1 1/K` Coefficient of the temperature dependence of `a1`.
- 65 `sla1=10e-6 m` Coefficient of the length dependence of `a1`.

Spectre Circuit Simulator Reference

Component Statements Part III

66	$swa1=0.1e-3$ m	Coefficient of the width dependence of $a1$.
67	$a2r=33$ V	Exponent of the weak-avalanche current.
68	$sla2=10e-6$ V m	Coefficient of the length dependence of $a2$.
69	$swa2=0.1e-3$ V m	Coefficient of the width dependence of $a2$.
70	$a3r=0.6$	Factor of the drain-source voltage above which weak-avalanche occurs.
71	$sla3=1e-6$ m	Coefficient of the length dependence of $a3$.
72	$swa3=10e-6$ m	Coefficient of the width dependence of $a3$.

Charge parameters

73	$tox=20e-9$ m	Thickness of the oxide layer.
74	$col=50e-12$ F/m	Gate overlap capacitance per unit channel width.

Noise parameters

75	$ntr=21e-21$ J	Coefficient of the thermal noise.
76	$nfmod=0.0$	Switch that selects either old or new flicker noise model.
77	$nfr=16e-12$ V ²	Flicker noise coefficient of the reference transistor (for $nfmod=1$).
78	$nfar=7.15e+22$ 1/(V m ⁴)	First coefficient of the flicker noise coefficient of the reference transistor (for $nfmod=1$).
79	$nfbr=2.16e+7$ 1/(V m ²)	Second coefficient of the flicker noise coefficient of the reference transistor (for $nfmod=1$).
80	$nfcr=0.0$ 1/V	Third coefficient of the flicker noise coefficient of the reference transistor (for $nfmod=1$).

Spectre Circuit Simulator Reference

Component Statements Part III

Temperature parameters

81	t_r (C)	Reference temperature. Default set by option t_{nom} .
82	t_{ref} (C)	Alias of t_r . Default set by option t_{nom} .
83	t_{nom} (C)	Alias of t_r . Default set by option t_{nom} .
84	$dt_a=0$ K	Temperature offset of the device.
85	$t_{rise}=0$ K	Alias of dt_a .

Other parameters

86	$th3mod=1$	Flag for theta3 clipping.
----	------------	---------------------------

Output Parameters

1	l_e (m)	Effective channel length.
2	w_e (m)	Effective channel width.
3	v_{to} (V)	Threshold voltage at zero back-bias.
4	k_o (\sqrt{V})	Low-backbias body factor.
5	k (\sqrt{V})	High-backbias body factor.
6	ϕ_{ib} (V)	Surface potential at strong inversion.
7	v_{sbx} (V)	Transition voltage for the dual-k-factor model.
8	β_{et} (A/V^2)	Gain factor (* mult).
9	$the1$ ($1/V$)	Coefficient of the mobility reduction due to the gate-induced field.
10	$the2$ ($1/\sqrt{V}$)	Coefficient of the mobility reduction due to the back-bias.
11	$the3$ ($1/V$)	Coefficient of the mobility reduction due to the lateral field.
12	γ_{m1} ($V^{(1-\epsilon_{tads})}$)	Coefficient for the drain induced threshold shift for large gate drive.

Spectre Circuit Simulator Reference

Component Statements Part III

13	etads	Exponent of the vds dependence of g_{am1} .
14	alp	Factor of the channel-length modulation.
15	vp (V)	Characteristic voltage of the channel-length modulation.
16	gamoo	Coefficient for the drain induced threshold shift at zero gate drive.
17	etagam	Exponent of the back-bias dependence of g_{amo} .
18	mo	Factor for the subthreshold slope.
19	etam	Exponent of the back-bias dependence of m .
20	phit (V)	Thermal voltage.
21	zet1	Weak-inversion correction factor.
22	vsbt (V)	Limiting voltage of the vsb dependence of m and g_{amo} .
23	a1	Factor of the weak-avalanche current.
24	a2 (V)	Exponent of the weak-avalanche current.
25	a3	Factor of the drain-source voltage above which weak-avalanche occurs.
26	cox (F)	Gate-to-channel capacitance (* mult).
27	cgdo (F)	Gate-drain overlap capacitance (* mult).
28	cgso (F)	Gate-source overlap capacitance (* mult).
29	nt (J)	Coefficient of the thermal noise.
30	nf (V ²)	Coefficient of the flicker noise (/ mult) (nfmod = 0).
31	nfa (1/(V m ⁴))	First coefficient of the flickernoise of the actual transistor (nfmod = 1).
32	nfba (1/(V m ²))	Second coefficient of the flickernoise of the actual transistor (nfmod = 1).

Spectre Circuit Simulator Reference

Component Statements Part III

33	nfc (1/V)	Second coefficient of the flicker noise of the actual transistor ($nfmod = 1$).
34	tox (m)	Thickness of gate oxide layer.

Operating-Point Parameters

1	ide (A)	Resistive drain current.
2	ige (A)	Resistive gate current.
3	ise (A)	Resistive source current.
4	ibe (A)	Resistive bulk current.
5	vds (V)	Drain-source voltage.
6	vgs (V)	Gate-source voltage.
7	vsb (V)	Source-bulk voltage.
8	ids (A)	Resistive drain-source current.
9	idb (A)	Resistive drain-bulk current.
10	isb (A)	Resistive source-bulk current.
11	$iavl$ (A)	Substrate current.
12	pwr (W)	Power.
13	$vt1$ (V)	V_{t0} including backbias effects.
14	$vgt2$ (V)	Effective gate drive including backbias and drain effects.
15	$vdss1$ (V)	Saturation voltage at actual bias.
16	$vsat$ (V)	Saturation limit.
17	gm (S)	Transconductance ($d\ ids / d\ vgs$).
18	gmb (S)	Bulk transconductance ($d\ ids / d\ vbs$).

Spectre Circuit Simulator Reference

Component Statements Part III

19	g_{ds} (S)	Output conductance ($d i_{ds} / d v_{ds}$).
20	c_{dd} (F)	Capacitance ($d q_d / d v_d$).
21	c_{dg} (F)	Capacitance ($- d q_d / d v_g$).
22	c_{ds} (F)	Capacitance ($- d q_d / d v_s$).
23	c_{db} (F)	Capacitance ($- d q_d / d v_b$).
24	c_{gd} (F)	Capacitance ($- d q_g / d v_d$).
25	c_{gg} (F)	Capacitance ($d q_g / d v_g$).
26	c_{gs} (F)	Capacitance ($- d q_g / d v_s$).
27	c_{gb} (F)	Capacitance ($- d q_g / d v_b$).
28	c_{sd} (F)	Capacitance ($- d q_s / d v_d$).
29	c_{sg} (F)	Capacitance ($- d q_s / d v_g$).
30	c_{ss} (F)	Capacitance ($d q_s / d v_s$).
31	c_{sb} (F)	Capacitance ($- d q_s / d v_b$).
32	c_{bd} (F)	Capacitance ($- d q_b / d v_d$).
33	c_{bg} (F)	Capacitance ($- d q_b / d v_g$).
34	c_{bs} (F)	Capacitance ($- d q_b / d v_s$).
35	c_{bb} (F)	Capacitance ($d q_b / d v_b$).
36	u	Transistor gain (g_m/g_{ds}).
37	r_{out} (Ω)	Small signal output resistance ($1/g_{ds}$).
38	v_{early} (V)	Equivalent Early voltage ($ i_d /g_{ds}$).
39	k_{eff} (\sqrt{V})	Describes body effect at actual bias.

Spectre Circuit Simulator Reference

Component Statements Part III

40	beff (S/V)	Effective beta at actual bias in the simple MOS model ($2* ids /vgt^2$).
41	fug (Hz)	Unity gain frequency at actual bias ($gm/(2*pi*cin)$).
42	sqrtsfw (V/ \sqrt{Hz})	Input-referred RMS white noise voltage (\sqrt{sth}/gm).
43	sqrtsff (V/ \sqrt{Hz})	Input-referred RMS 1/f noise voltage at 1kHz ($\sqrt{nf/1000}$).
44	fknee (Hz)	Cross-over frequency above which white noise is dominant.

Parameter Index

In the following index, I refers to instance parameters, M refers to the model parameters section, O refers to the output parameters section, and OP refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of M-35 means the 35th model parameter.

a1	O-23	fthel	M-31	phibr	M-19	swvsbx	M-22
alr	M-63	fug	OP-41	phit	O-20	swvto	M-12
a2	O-24	gam1	O-12	pwr	OP-12	th3mod	M-86
a2r	M-67	gam1r	M-42	region	I-5	thel	O-9
a3	O-25	gamoo	O-16	rout	OP-37	thelr	M-25
a3r	M-70	gamoor	M-51	sl2vto	M-11	the2	O-10
alp	O-14	gds	OP-19	sla1	M-65	the2r	M-32
alpr	M-46	gm	OP-17	sla2	M-68	the3	O-11
area	I-4	gmb	OP-18	sla3	M-71	the3r	M-37
beff	OP-40	iavl	OP-11	slalp	M-48	tnom	M-83
bet	O-8	ibe	OP-4	slgam1	M-43	tox	M-73
betsq	M-23	idb	OP-9	slgamoo	M-52	tox	O-34
cbb	OP-35	ide	OP-1	slk	M-17	tr	M-81
cbd	OP-32	ids	OP-8	slko	M-14	tref	M-82
cbg	OP-33	ige	OP-2	slmo	M-56	trise	M-85

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cbs	OP-34	isb	OP-10	slthe1r	M-27	type	M-1
cdb	OP-23	ise	OP-3	slthe2r	M-34	u	OP-36
cdd	OP-20	k	O-5	slthe3r	M-39	vds	OP-5
cdg	OP-21	keff	OP-39	slvsbt	M-62	vdss1	OP-15
cds	OP-22	ko	O-4	slvsbx	M-21	vearly	OP-38
cgb	OP-27	kor	M-13	slvto	M-10	vgs	OP-6
cgd	OP-24	kr	M-16	slzet1	M-60	vgt2	OP-14
cgdo	O-27	l	I-2	sqrtsff	OP-43	vp	O-15
cgg	OP-25	lap	M-5	sqrtsfw	OP-42	vpr	M-50
cgs	OP-26	le	O-1	stal	M-64	vsat	OP-16
cgso	O-28	ler	M-2	stlthe1	M-28	vsb	OP-7
col	M-74	lvar	M-4	stlthe2	M-35	vsbt	O-22
cox	O-26	m	I-6	stlthe3	M-40	vsbtr	M-61
csb	OP-31	mo	O-18	stmo	M-55	vsbx	O-7
csd	OP-28	mor	M-54	stthe1r	M-26	vsbxr	M-20
csg	OP-29	mult	I-3	stthe2r	M-33	vt1	OP-13
css	OP-30	nf	O-30	stthe3r	M-38	vto	O-3
dta	M-84	nfa	O-31	stvto	M-9	vtor	M-8
etaalp	M-47	nfarc	M-78	swa1	M-66	w	I-1
etabet	M-24	nfb	O-32	swa2	M-69	wdog	M-30
etads	O-13	nfbr	M-79	swa3	M-72	we	O-2
etadsr	M-45	nfc	O-33	swalp	M-49	wer	M-3
etagam	O-17	nfcrc	M-80	swgam1	M-44	wot	M-7
etagamr	M-53	nfmod	M-76	swk	M-18	wvar	M-6
etam	O-19	nfr	M-77	swko	M-15	zet1	O-21
etamr	M-57	nt	O-29	swthe1	M-29	zet1r	M-58
etazet	M-59	ntr	M-75	swthe2	M-36		
fknee	OP-44	phib	O-6	swthe3	M-41		

Microstrip Line (msli432'1'ne)

Description

This is a microstrip line based on the equations of Hammerstad and Jensen. The model contains a thickness correction to the width and frequency dependent permittivity and characteristic impedance. The dispersion equations are those of Kirschning and Jansen.

This device is supported within altergroups.

Sample Instance Statement

```
t11 (in 0 out 0) msline l=0.15 w=0.01 h=0.01
```

Instance Definition

```
Name t1 b1 t2 b2 msline parameter=value ...
```

Instance Parameters

- | | | |
|---|---------------------------|--|
| 1 | <code>l=0 m</code> | Length. |
| 2 | <code>w (m)</code> | Width. |
| 3 | <code>h (m)</code> | Substrate height. |
| 4 | <code>t=0 m</code> | Conductor thickness. |
| 5 | <code>eps=1</code> | Substrate permittivity relative to a vacuum. |
| 6 | <code>m=1</code> | Multiplicity factor. |
| 7 | <code>fmax=10e9 Hz</code> | Maximum signal frequency. |

Multi-Conductor Transmission Line (mtline)

Description

VA1 IA1 -> |-----| <- IB1 VB1

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Component Statements Part III

in1 -o-----+o=====o+-----o- out1

| |

VA2 IA2 -> | | <- IB2 VB2

in2 -o-----+o=====o+-----o- out2

. | . | .
 . | . | .
 . | . | .

VAN IAN -> | | <- IBN VBN

inN -o-----+o=====o+-----o- outN

| |
 | |

inref -o-----+o=====o+-----o- outref

|-----|

A multi-conductor transmission line (mtline) is characterized by constant RLGC matrices or frequency dependent RLGC data. An mtline can have as many conductors as there are as described in the input, however, there must be at least two conductors with one conductor used as reference to define terminal voltages. The reference conductor can be ground. The order of the conductors is the same as the order of the data in the input.

All of the conductors are assumed to have the same length, the input to mtline are conductor length, per-unit-length resistance (R), inductance (L), conductance (G), and capacitance (C) matrices. Because these matrices are generally symmetric, either full matrix description or lower half matrix description can be used. For example, to describe the resistance matrix of a four conductor mtline:

[50 10 1]

R = [10 50 10] Ohm/meter

[1 10 50]

The following two model descriptions are equivalent:

Spectre Circuit Simulator Reference

Component Statements Part III

model line mtline

+ r=[50 10 1

+ 10 50 10

+ 1 10 50]

+ ...

model line mtline

+ r=[50

+ 10 50

+ 1 10 50]

+ ...

Frequency dependent RLGC matrices are described in a data file through parameter `file`. The frequency axis can be scaled with the `scale` parameter. The frequencies in the data file are then multiplied by `scale` before the simulator uses them. The default scale factor is unity. An example data file is listed below:

```
; Comments: rl.dat
```

```
FORMAT FREQ: R1:1 R2:1 R2:2
```

```
          L1:1 L2:1 L2:2
```

```
0.001e+9:  4.444 0.000383 4.444
```

```
          4.565 0.3545  4.565
```

```
0.010e+9:  4.447 0.003834 4.447
```

```
          4.565 0.3545  4.565
```

```
0.100e+9   4.476 0.03834  4.476
```

```
          4.565 0.3545  4.565
```

```
1.000e+9   4.762 0.3834  4.762
```

```
          3.103 0.2357  3.103
```

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Component Statements Part III

10.00e+9 13.96 1.082 13.96

2.718 0.2058 2.718

100.0e+9 56.88 3.294 56.88

2.531 0.1866 2.531

Constant matrix is the first choice of input if both the constant matrix and tabular data are provided. If only one frequency point is provided in the `file`, the RLGC matrices are treated as constant matrices.

This device is not supported within altergroup.

Sample Instance Statement

```
x1 (a1 b1 a2 b2 0 0) mtline len=0.01
```

```
+ r=[ 0.3
```

```
+ 0.0 0.3 ]
```

```
+ c=[ 0.35p
```

```
+ -0.03p 0.35p ]
```

Sample Model Statement

```
model mtmodel mtline
```

```
+ r=[ 0.3
```

```
+ 0.0 0.3 ]
```

```
+ c=[ 0.35p
```

```
+ -0.03p 0.35p ]
```

```
model mtmodel mtline
```

```
+ r=[ 0.3 0.0
```

```
+ 0.0 0.3 ]
```

```
+ c=[ 0.35p -0.03p
```

Spectre Circuit Simulator Reference

Component Statements Part III

```
+ -0.03p 0.35p ]
model mtmodel mtline
+ c=[ 0.35p
+ -0.03p 0.35p ]
+ file="rl.data" scale=1
```

Instance Definition

```
Name in1 out1 in2 [out2] ... ModelName parameter=value ...
Name in1 out1 in2 [out2] ... mtline parameter=value ...
```

The last two terminals will be used as refin and refout respectively.

Instance Parameters

1 len=0.01 m Physical length of line.
2 m=1 Multiplicity factor.

RLGC data parameters

3 r=[...] Ω/m Resistance matrix per unit length.
4 l=[...] H/m Inductance matrix per unit length.
5 g=[...] S/m Conductor matrix per unit length.
6 c=[...] F/m Capacitance matrix per unit length.
7 rskin=[...] $\Omega/m \sqrt{Hz}$ Skin effect resistance matrix per unit length.
8 gdloss=[...] S/m Hz Dielectric loss conductance matrix per unit length.
9 file RLGC data file that contains the frequency dependent RLGC data.
10 freqscale=1 Frequency scale factor for frequency dependent RLGC data.

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Component Statements Part III

Rational fitting parameters

11 `fmax=2.5e10 Hz` Maximum signal frequency used to determine the relevant range of rational fitting.

Model Definition

`model modelName mtline parameter=value ...`

Model Parameters

RLGC data parameters

- 1 `r=[...] Ω/m` Resistance matrix per unit length.
- 2 `l=[...] H/m` Inductance matrix per unit length.
- 3 `g=[...] S/m` Conductor matrix per unit length.
- 4 `c=[...] F/m` Capacitance matrix per unit length.
- 5 `rskin=[...] $\Omega/m \sqrt{Hz}$` Skin effect resistance matrix per unit length.
- 6 `gdloss=[...] S/m Hz` Dielectric loss conductance matrix per unit length.
- 7 `file` RLGC data file that contains the frequency dependent RLGC data.
- 8 `freqscale=1` Frequency scale factor for frequency dependent RLGC data.

Rational fitting parameters

9 `fmax=2.5e10 Hz` Maximum signal frequency used to determine the relevant range of rational fitting.

Important Note About Rational Fitting

Spectre uses rational fitting algorithm to build a stable model that approximates the desired transmission line characteristics. Maximum signal frequency f_{max} is used to determine the

Spectre Circuit Simulator Reference

Component Statements Part III

relevant range of rational fitting. The accuracy of the mtline model is solely dependent on how well the rational approximation is over frequency range [f_{min} , f_{max}].

When constant RLGC matrices are provide, 1Hz is used as f_{min} and f_{max} is defaulted to 25GHz. Three times the inverse of rise time in the input signal can be used as a good estimation of f_{max} . When RLGC data file is provided, the lowest frequency point in the data file is used as f_{min} , and the largest frequency point in the data file is used as f_{max} . User should provide sufficient data points to cover both low-frequency and high-frequency for an accurate, stable model.

Modeling frequency dependent effects:

One can model the frequency dependent RLGC matrices by providing the data file using parameter `file`. One should always try to provide accurate and sufficient data to describe the frequency dependent RLGC matrices.

In addition, the following simplified equation can be used to model skin effect with the constant RLGC matrices

$$R(f) = r + \text{sqrt}(f) * (1 + j) * r_{\text{skin}},$$

and the following equation can be used to model dielectric loss with the constant RLGC matrices

$$G(f) = g + f * g_{\text{dloss}},$$

where f stands for frequency. User should be aware of that the equation of $G(f)$ results in a non-physical model, and it could lead to an unstable rational model.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

<code>c</code>	M-4	<code>fmax</code>	M-9	<code>gdloss</code>	I-8	<code>m</code>	I-2
<code>c</code>	I-6	<code>freqscale</code>	I-10	<code>gdloss</code>	M-6	<code>r</code>	M-1
<code>file</code>	M-7	<code>freqscale</code>	M-8	<code>l</code>	I-4	<code>r</code>	I-3
<code>file</code>	I-9	<code>g</code>	I-5	<code>l</code>	M-2	<code>rskin</code>	M-5

fmax I-11

g M-3

len I-1

rskin I-7

Mutual Inductor (mutual_inductor)

Description

The mutual inductor couples two previously specified inductors. There is no limit to the number of inductors that you can couple or to the number of couplings to a particular inductor, but you must specify separate mutual inductor statements for each coupling. Using the `dot` convention, place a `dot` on the first terminal of each inductor.

This device is not supported within `altergroup`.

The mutual inductor modifies the constitutive equations of two isolated inductors to

$$v1 = L11*di1/dt + M*di2/dt$$

$$v2 = M*di1/dt + L22*di2/dt$$

where the mutual inductance, M , is computed from the coupling coefficient, k , using $k = |M|/\sqrt{L11*L22}$.

Sample Instance Statement with Two Inductors

```
l1 (1 0) inductor
```

```
l2 (2 0) inductor
```

```
ml1 mutual_inductor coupling=1 ind1=l1 ind2=l2
```

Instance Definition

```
Name mutual_inductor parameter=value ...
```

Instance Parameters

- | | | |
|---|-------------------------|-------------------------|
| 1 | <code>coupling=0</code> | Coupling coefficient. |
| 2 | <code>ind1</code> | Inductor to be coupled. |
| 3 | <code>ind2</code> | Inductor to be coupled. |

Node Capacitance (nodcap)

Description

The nodcap model is generally used to model voltage dependent capacitances and currents of the source and drain diodes of MOS transistors and the capacitances of related interconnection areas and sidewalls (IN and PS regions). It is described in the Philips MOST Modelbook (Dec.93) as NODCAP model.

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In extension to the modelbook description a minimum conductance g_{min} is inserted between the nodcap nodes, to aid convergence. The value of g_{min} is set by an options statement, default = 1e-12 S.

The i_{max} parameter is used to aid convergence and to prevent numerical overflow. The junction characteristics of the node capacitor are accurately modeled for currents up to i_{max} . For currents above i_{max} , the junction is modeled as a linear resistor and a warning is printed.

This device is supported within altergroups.

This device is dynamically loaded from the shared object /vobs/spectre_dev/tools.sun4v/spectre/lib/cmi/3.0.doc/libphilips_sh.so

Sample Instance Statement

```
n14 (outc vcc) pcapmod ad=191.13e-6 pdg=70e-6 pdcs=75
```

Sample Model Statement

```
model pcapmod nodcap type=p js=10e-6 pt=2 cjr=0.6e-3 vdr=750e-3 tref=27 pmr=0.4  
vr=0 kpers=4.9e-6 cox=1.77e-3
```

Instance Definition

```
Name nn [ns] ModelName parameter=value ...
```

Instance Parameters

1 m=1.0 Multiplicity factor.

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Component Statements Part III

2	<code>region=rev</code>	Estimated DC operating region, used as a convergence aid. Spectre outputs number (0-2) in a rawfile. Possible values are <code>fwd</code> , <code>rev</code> or <code>brk</code> .
3	<code>cf=0.0 F</code>	Fixed capacitance.
4	<code>ad=0.0 scale² m²</code>	Diffusion area (source or drain). Scale set by option <code>scale</code> .
5	<code>pdcs=0.0 scale m</code>	Length of the side-wall of the diffusion area AD which is not under the gate. Scale set by option <code>scale</code> .
6	<code>pdg=0.0 scale m</code>	Length of the side-wall of the diffusion area AD which is under the gate. Scale set by option <code>scale</code> .
7	<code>ain=0.0 scale² m²</code>	Area of metal interconnection over thick oxide. Scale set by option <code>scale</code> .
8	<code>ag=0.0 scale² m²</code>	Area metal or poly-Si over thin oxide. Scale set by option <code>scale</code> .
9	<code>aps=0.0 scale² m²</code>	Area metal of poly-Si over thick oxide. Scale set by option <code>scale</code> .
10	<code>pin=0.0 scale m</code>	Length of the side-wall of AIN. Scale set by option <code>scale</code> .
11	<code>pps=0.0 scale m</code>	Length of the side-wall of APS. Scale set by option <code>scale</code> .

Model Definition

```
model modelName nodcap parameter=value ...
```

Model Parameters

1	<code>type=n</code>	Type of the nodcap device. Possible values are <code>n</code> or <code>p</code> .
2	<code>kpjs=0.0 m</code>	Conversion factor of the sidewall length and the gate edge length to its equivalent area for the saturation current.
3	<code>n=1.0</code>	Emission coefficient.

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Component Statements Part III

4	$bv=\infty$ V	Reverse break-down voltage.
5	$imax=1.0$ A	Explosion current.
6	$pt=2.0$	Temperature coefficient of the saturation current.
7	$vr=0.0$ V	Voltage at which CJR is specified.
8	$fc=0.5$	Forward-bias non-ideal junction capacitance coefficient.
9	$pmr=0.5$	Grading coefficient.
10	$kpers=0.0$ m	Conversion factor of the sidewall length to its equivalent area for the junction capacitance.
11	$kperg=0.0$ m	Conversion factor of the gate edge length to its equivalent area for the junction capacitance.
12	$cin=0.0$ F/m ²	Specific capacitance of the (interconnection) metal to the substrate.
13	$cox=0.0$ F/m ²	Specific capacitance of poly-silicon or aluminum to the substrate over thin oxide.
14	$cps=0.0$ F/m ²	Specific capacitance of poly-silicon to the substrate over thick oxide.
15	$cins=0.0$ F/m	IN-side-wall capacitance.
16	$cpss=0.0$ F/m	PS-side-wall capacitance.
17	$dta=0.0$ K	Temperature offset of the device with respect to TEMP.
18	$trise$ (K)	Alias of dta .
19	$tref=27.0$ C	Temperature at which the parameters are specified. Default set by option $tnom$.
20	$tnom$ (C)	Alias of $tref$.
21	tr (C)	Alias of $tref$.
22	$js=100u$ A/m ²	Saturation current density.

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Component Statements Part III

23 `cjr=0.0 F/m2` Specific junction capacitance at $V_d = V_R$.

24 `vdr=0.8 V` Diffusion voltage.

Output Parameters

1 `jst (A/m2)` Saturation current density (temperature updated).

2 `cjrt (F/m2)` Specific junction capacitance at $V_d = V_R$ (temperature updated).

3 `vdrt (V)` Diffusion voltage (temperature updated).

1 `i0 (A)` Saturation current (temperature updated).

Operating-Point Parameters

1 `region=rev` Estimated DC operating region, used as a convergence aid. Spectre outputs number (0-2) in a rawfile. Possible values are `fwd`, `rev` or `brk`.

2 `vsub (V)` Voltage across the node-capacitance, which is measured from NS (substrate or N-well) to NN (source or drain). `Vsub` is usually negative.

3 `isub (A)` Resistive leakage current.

4 `cap (F)` Junction capacitance.

5 `q (Coul)` Junction charge.

6 `gm (S)` Total differential conductance.

7 `c0 (F)` Constant part of the junction capacitance.

8 `pwr (W)` Power.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the

Spectre Circuit Simulator Reference Component Statements Part III

description for that parameter. For example, a reference of M-35 means the 35th model parameter.

ad	I-4	cox	M-13	kpers	M-10	region	OP-1
ag	I-8	cps	M-14	kpjs	M-2	region	I-2
ain	I-7	cpss	M-16	m	I-1	tnom	M-20
aps	I-9	dta	M-17	n	M-3	tr	M-21
bv	M-4	fc	M-8	pdcs	I-5	tref	M-19
c0	OP-7	gm	OP-6	pdg	I-6	trise	M-18
cap	OP-4	i0	O-1	pin	I-10	type	M-1
cf	I-3	imax	M-5	pmr	M-9	vdr	M-24
cin	M-12	isub	OP-3	pps	I-11	vdrt	O-3
cins	M-15	js	M-22	pt	M-6	vr	M-7
cjr	M-23	jst	O-1	pwr	OP-8	vsub	OP-2
cjrt	O-2	kperg	M-11	q	OP-5		

Set Node Quantities (node)

Description

Quantities are used to hold information about particular types of signals, such as their units, absolute tolerances, and maximum allowed change per Newton iteration. Use the `quantity` statement to create new quantities or to redefine properties of an existing quantity. Use this statement to set the quantities for a particular node.

For example, to indicate that the node `net1` is used for thermal signals, the following node statement could be used.

```
i17 (net1) node value=Temp flow=Pwr
```

`Temp` and `Pwr` are predefined quantities.

This device is not supported within `altergroup`.

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Component Statements Part III

Sample Instance Statement

```
node1 (1 2 3) node value="T" flow="W" strength=override //Must define T and W with
quantity statement.
```

Instance Definition

```
Name 1 [2] ... node parameter=value ...
```

Instance Parameters

- | | | |
|---|-------------------|---|
| 1 | value | Value quantity. |
| 2 | flow | Flow quantity. |
| 3 | strength=override | Quantity strength.
Possible values are indifferent, suggest, insist, or
override. |

Linear N Port (nport)

Description

An N-port takes its characteristics from an S, Y or Z-parameter data file. An N-port can have as many ports as there are in the N-port described in the data file. Each pair of terminals in the `nport` instance statement represents one port. Because there is no limit to the number of ports, there is no limit to the number of terminals. However, the terminals must be given in pairs and there must be at least one pair. The order of the pairs is the same as the order of the ports in the data file. Any missing ports should be skipped.

The S, Y or Z-parameter data file specifies the characteristics of the N-port. You can scale the frequency axis with the `scale` parameter. The frequencies in the data file are then multiplied by `scale` before the simulator uses them. The default scale factor is unity. In addition to Spectres native format, the S, Y, or Z-parameters can be in Touchstone or CITIfile format, and the data can be given as: real-imag, mag-deg, mag-rad, db-deg, or db-rad.

An internal thermal noise model is used in noise analysis. However, if the user provides frequency-dependent noise data then this data is used to build a noise model. Currently Touchstone format accepts the two-port noise parameters (NFmin, Gamma (Gopt, Bopt) and Rn), while Spectres native format accepts both two-port noise parameters and noise correlation matrix.

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Component Statements Part III

If `interp=linear` or `spline` is specified, then impulse response is calculated, and convolution method is used for simulation in time domain. It is assumed that the S, Y or Z-parameter data is complete and smooth enough to be safely interpolated or extrapolated in the frequency range from `fdelta` to `fmax`, and to DC. Be aware that N-port can be used to model many different kind of systems, the default setting of the impulse response and convolution algorithm is made for typical N-port applications. See the important notes below on when and how to set some of the controlling parameters.

Linear interpolation or cubic spline is used on the data in polar form. A simple algorithm removes 2π jumps in the phase data. Frequency points where the data is measured must therefore be close enough to avoid an excessive number of jumps. Unfortunately, noisy phase data can cause unnecessary warning messages. Number of frequency domain samples used for FFT is `fmax/fdelta` with upper limit of `maxn`. It might be necessary to increase `maxn` in the case of extremely large average group delay. When `usewindow` is set to `yes`, the data in frequency domain is multiplied by Kaiser-Bessel window function with smoothing parameter equal to one.

If `interp=rational` is specified, the data is interpolated and extrapolated using a rational function fit to the data. The degree of rational interpolation is automatically selected based on the values of `abserr` and `relerr`, unless `ratorder` is given, in which case `relerr` and `abserr` are ignored in selecting the order of the rational function interpolation. It is usually better to allow the simulator to automatically select the rational interpolation order.

If the S, Y or Z-parameter data contains noise, `abserr` and `relerr` should be set so that the fitting procedure can ignore the noise, for example, by setting `abserr` above the noise floor and/or relaxing `relerr` as necessary.

Because the fitting procedure can take a long time for complicated data, the reduced order model (ROM) file option is available to store and re-use the rational interpolation function in subsequent simulations.

It is not practical to rely on extrapolated data.

This device is not supported within `altergroup`.

Sample Instance Statement

```
x1 (a1 0 b1 0 b3 0) ndata file="sparam2.data"
```

Sample Model Statement

```
model ndata nport file="sparam.data" scale=1
```

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Component Statements Part III

Instance Definition

Name t1 b1 [t2] [b2] ... ModelName parameter=value ...

Name t1 b1 [t2] [b2] ... nport parameter=value ...

Terminals must be given in pairs.

Instance Parameters

- | | | |
|---|---------------|--|
| 1 | m=1 | Multiplicity factor. |
| 2 | file | S-parameter data file name. |
| 3 | datafmt | The format of the S-parameter data file. If this parameter is not given, Spectre will try to detect the format by itself.
Possible values are <code>spectre</code> , <code>touchstone</code> or <code>citi</code> . |
| 4 | scale=1 | Frequency scale factor. |
| 5 | interp=spline | Method to interpolate s-parameter data.
Possible values are <code>spline</code> , <code>rational</code> or <code>linear</code> . |

Spline/Linear interpolation parameters

- | | | |
|----|-----------------|---|
| 6 | fmax (Hz) | Maximum frequency of interest. Default is 3 times the highest frequency found in s-parameter file. |
| 7 | fdelta (Hz) | Frequency sampling interval. Default value is fmax/1024, or 1/tstop, whichever is smaller. |
| 8 | maxn=4096 | Maximal order of impulse response. Cannot exceed 16384. |
| 9 | imptrunc=1.0e-4 | Impulse response truncation threshold relative to the main maximum. The tail of the impulse response below imptrunc will be removed. Set imptrunc=0 to keep the tail. |
| 10 | usewindow=no | Use smooth data windowing function.
Possible values are <code>no</code> or <code>yes</code> . |

Spectre Circuit Simulator Reference

Component Statements Part III

Rational interpolation parameters

- | | | |
|----|--------------------------|--|
| 11 | <code>relerr=0.01</code> | Maximum relative allowed tolerance for rational interpolation errors. Deviations of the nport model from supplied s-parameter data of relative magnitude less than <code>relerr</code> are generally ignored. |
| 12 | <code>abserr=1e-4</code> | Maximum absolute allowed tolerance for rational interpolation errors. Deviations of the nport model from supplied s-parameter data of absolute magnitude less than <code>abserr</code> are generally ignored. |
| 13 | <code>romdatfile</code> | File used for storing time-domain reduced order model (ROM). |
| 14 | <code>ratorder</code> | Order of rational function to use in fitting the s-parameter data. If this argument is given, <code>relerr</code> and <code>abserr</code> are ignored in selecting the order of the rational function interpolation. If <code>ratorder</code> is not specified then the program will attempt to select an order of rational interpolation that satisfies the criteria implied by <code>abserr</code> and <code>relerr</code> . |

Noise parameters

- | | | |
|----|-------------------------------|---|
| 15 | <code>trise (C)</code> | Temperature rise from ambient. |
| 16 | <code>thermalnoise=yes</code> | Thermal noise.
Possible values are <code>no</code> or <code>yes</code> . |
| 17 | <code>noisemodel</code> | To use the internal thermal noise model, or the externally supplied noise data in the S-parameter data file. The default behavior is to use external data whenever it is available.
Possible values are <code>internal</code> or <code>external</code> . |

Model Definition

`model modelName nport parameter=value ...`

Model Parameters

- | | | |
|---|-------------------|-----------------------------|
| 1 | <code>file</code> | S-parameter data file name. |
|---|-------------------|-----------------------------|

Spectre Circuit Simulator Reference

Component Statements Part III

2	<code>datafmt</code>	The format of the S-parameter data file. If this parameter is not given, Spectre will try to detect the format by itself. Possible values are <code>spectre</code> , <code>touchstone</code> or <code>citi</code> .
3	<code>scale=1</code>	Frequency scale factor.
4	<code>fmax (Hz)</code>	Maximum frequency of interest. Default is 3 times the highest frequency provided in the s-parameter file.
5	<code>fdelta (Hz)</code>	Frequency sampling interval. Default value is <code>fmax/1024</code> , or <code>1/tstop</code> , whichever is smaller.
6	<code>maxn=4096</code>	Maximal order of impulse response.
7	<code>imptrunc=1.0e-4</code>	Relative truncation threshold for the impulse response.
8	<code>usewindow=no</code>	Use smooth data windowing function. Possible values are <code>no</code> or <code>yes</code> .
9	<code>trise=0 C</code>	Default temperature rise from ambient.
10	<code>thermalnoise=yes</code>	Thermal noise. Possible values are <code>no</code> or <code>yes</code> .
11	<code>noisemodel</code>	To use the internal thermal noise model, or the externally supplied noise data in the S-parameter data file. The default behavior is to use external data whenever it is available. Possible values are <code>internal</code> or <code>external</code> .

Important note about spline and linear interpolation parameters

`interp`

To calculate impulse response, either linear or spline interpolation is used to sample data points in the frequency domain, based on the S-parameter data file. When the input data points are sufficient, linear and spline interpolation produce comparable model accuracy. However, when the input data points are scarce, linear interpolation is preferred to bound jumps between data points. Interpolation/extrapolation can be avoided all together if s-parameter data is provided from zero frequency to `fmax` with uniform step `fdelta`, and number of samples is a power of 2.

`fmax`

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The default of `fmax` is three times the highest frequency in the S-parameter data file. Impulse response is calculated by sampling frequency points between DC and f_{\max} . The purpose of this extrapolation is to preserve high-frequency model accuracy. However, if user has enough bandwidth in the S-parameter data file, `fmax` needs to be set to the original highest frequency in the S-parameter data file. In general, `fmax` must be larger than expected signal bandwidth.

`fdelta`

Frequency sampling interval, with default of $f_{\max}/1024$, or $1/t_{\text{stop}}$, whichever is smaller. Note that `fdelta` will be further reduced, so that `fmax/fdelta` is a power of 2. This is required by FFT algorithm.

`maxn`

Maximal order of impulse response, or the maximum number of sampling points in the frequency domain. User should only increase this value when modeling a system with large average group delay, such as transmission line. The given value of `maxn` will be reduced to the nearest power of 2. Setting this value unnecessarily large slows down time-domain simulation. Absolute upper limit for `maxn` is 16384.

`imptrunc`

Relative truncation threshold for impulse response. The tail of the impulse response with absolute values below `imptrunc` is removed to speed up simulation.

`usewindow`

Kaiser-Bessel window with smoothing parameter 1 is used to better regulate the stability of S-parameter data. This parameter trades off model accuracy with model stability, particular useful for S-parameter data with insufficient bandwidth. Window must be used for all-pass type systems, where s-parameters are nonzero for the frequencies beyond `fmax`.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

<code>abserr</code>	I-12	<code>fmax</code>	I-6	<code>maxn</code>	I-8	<code>scale</code>	I-4
<code>datafmt</code>	I-3	<code>fmax</code>	M-4	<code>noisemodel</code>	M-11	<code>thermalnoise</code>	I-16

Spectre Circuit Simulator Reference Component Statements Part III

datafmt M-2	imptrunc I-9	noisemodel I-17	thermalnoise M-10
fdelta M-5	imptrunc M-7	ratorder I-14	trise M-9
fdelta I-7	interp I-5	relerr I-11	trise I-15
file I-2	m I-1	romdatfile I-13	usewindow I-10
file M-1	maxn M-6	scale M-3	usewindow M-8

Parameter Value Tester (paramtest)

Description

The parameter value tester tests the values of its parameters and prints a message if they satisfy the testers criteria. The tester therefore allows you to check the ranges of subcircuit parameters. If you specify more than one test, the message is printed if any test passes. The message is also printed if no tests are specified.

This device is not supported within altergroup.

Sample Instance Statement

```
tooShort paramtest errorif=(l < 0.2um) message="W of device is less than 0.2um"
```

Instance Definition

```
Name paramtest parameter=value ...
```

Instance Parameters

- | | |
|-------------------|---|
| 1 printif | Message is printed if this value is nonzero. |
| 2 warnif | Message is printed as a warning if this value is nonzero. |
| 3 errorif | Message is printed as an error and program quits if this value is nonzero. |
| 4 message | Text of message. |
| 5 severity=status | Message severity (use if printing message without test). Possible values are debug, status, warning, error, or fatal. |

Polynomial Current Controlled Current Source (pcccs)

Description

A vector of coefficients specifies the polynomial function that defines the relationship between the output current and the controlling currents. You must specify at least one coefficient.

This device is not supported within altergroup.

For a polynomial in M variables a_1, a_2, \dots, a_m , the polynomial function $F(a_0, a_1, \dots, a_m)$ is given by

$$\begin{aligned} F = & c_0 + c_1 * a_1 + c_2 * a_2 + \dots \\ & + c_{(m+1)} * a_1^2 + c_{(m+2)} * a_1 * a_2 + \dots \\ & + c_{(2m+1)} * a_2^2 + c_{(2m+2)} * a_2 * a_3 + \dots \end{aligned}$$

where the c_s are coefficients of the polynomial terms.

Sample Instance Statement

```
vpc (net1 0) pcccs probes=[vb vc ve vlp vpn] coeffs=[0 8.8e6 -8.8e6 9e6 8e6 -9e6]
```

Instance Definition

```
Name sink src pcccs parameter=value ...
```

Instance Parameters

- | | | |
|---|---------------------------|--|
| 1 | <code>file</code> | File that contains nonzero polynomial coefficients. |
| 2 | <code>coeffs=[...]</code> | Polynomial coefficients. At least one must be given. |
| 3 | <code>probes=[...]</code> | Devices through which the controlling currents flow. |
| 4 | <code>ports=[...]</code> | Indice of the probe ports through which the controlling currents flow. |
| 5 | <code>gain=1</code> | Gain Parameter. |
| 6 | <code>m=1</code> | Multiplicity factor. |

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Component Statements Part III

7	<code>min</code> (A)	Minimum output current.
8	<code>max</code> (A)	Maximum output current.
9	<code>abs=off</code>	Absolute output current. Possible values are <code>off</code> or <code>on</code> .
10	<code>delta=0</code>	Smoothing parameter. This may lead to circuit convergency. The smaller the delta is, the sharper the corner is.

Temperature effects parameters

11	<code>tc1=0</code> 1/C	Linear temperature coefficient.
12	<code>tc2=0</code> C ⁻²	Quadratic temperature coefficient.

Operating-Point Parameters

1	<code>i</code> (A)	Output current.
2	<code>v</code> (V)	Output voltage.
3	<code>pwr</code> (W)	Power dissipation.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

<code>abs</code>	I-9	<code>gain</code>	I-5	<code>min</code>	I-7	<code>tc1</code>	I-11
<code>coeffs</code>	I-2	<code>i</code>	OP-1	<code>ports</code>	I-4	<code>tc2</code>	I-12
<code>delta</code>	I-10	<code>m</code>	I-6	<code>probes</code>	I-3	<code>v</code>	OP-2
<code>file</code>	I-1	<code>max</code>	I-8	<code>pwr</code>	OP-3		

Polynomial Current Controlled Voltage Source (pccvs)

Description

The polynomial function defining the relationship between the output voltage and the controlling currents is specified by a vector of coefficients. At least one coefficient must always be specified. Current through the voltage source is calculated and is defined to be positive if it flows from the positive terminal, through the source, to the negative terminal.

This device is not supported within altergroup.

For a polynomial in M variables a_1, a_2, \dots, a_m , the polynomial function $F(a_0, a_1, \dots, a_m)$ is given by

$$\begin{aligned} F = & c_0 + c_1 * a_1 + c_2 * a_2 + \dots \\ & + c_{(m+1)} * a_1^2 + c_{(m+2)} * a_1 * a_2 + \dots \\ & + c_{(2m+1)} * a_2^2 + c_{(2m+2)} * a_2 * a_3 + \dots \end{aligned}$$

where the c_s are coefficients of the polynomial terms.

Sample Instance Statement

```
ixy (net1 0) pccvs coeffs=[0 1 0 1] probes=[vin1 vin2] gain=2
```

Instance Definition

```
Name p n pccvs parameter=value ...
```

Instance Parameters

- | | | |
|---|---------------------------|---|
| 1 | <code>file</code> | File that contains nonzero polynomial coefficients. |
| 2 | <code>coeffs=[...]</code> | Polynomial coefficients. At least one must be given. |
| 3 | <code>probes=[...]</code> | Devices through which the controlling currents flow. |
| 4 | <code>ports=[...]</code> | Indices of the probe ports through which the controlling currents flow. |
| 5 | <code>gain=1</code> | Gain Parameter. |

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Component Statements Part III

6	<code>m=1</code>	Multiplicity factor.
7	<code>min (V)</code>	Minimum output voltage.
8	<code>max (V)</code>	Maximum output voltage.
9	<code>abs=off</code>	Absolute output voltage. Possible values are <code>off</code> or <code>on</code> .
10	<code>delta=0</code>	Smoothing parameter. This may lead to circuit convergency. The smaller the delta is, the sharper the corner is.

Temperature effects parameters

11	<code>tc1=0 1/C</code>	Linear temperature coefficient.
12	<code>tc2=0 C⁻²</code>	Quadratic temperature coefficient.

Operating-Point Parameters

1	<code>i (A)</code>	Output current.
2	<code>v (V)</code>	Output voltage.
3	<code>pwr (W)</code>	Power dissipation.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

<code>abs</code>	I-9	<code>gain</code>	I-5	<code>min</code>	I-7	<code>tc1</code>	I-11
<code>coeffs</code>	I-2	<code>i</code>	OP-1	<code>ports</code>	I-4	<code>tc2</code>	I-12
<code>delta</code>	I-10	<code>m</code>	I-6	<code>probes</code>	I-3	<code>v</code>	OP-2
<code>file</code>	I-1	<code>max</code>	I-8	<code>pwr</code>	OP-3		

Physical Resistor (phy_res)

Description

A physical resistor consists of a two terminal resistor (tied between t_1 and t_2) and two diodes (tied between t_1-t_0 and t_2-t_0). The diodes are junction diodes. Under normal operation, the two diodes are reverse biased, but the parameter `subtype` can reverse the direction of the diodes. If you do not specify t_0 , ground is assumed. The instance parameters always override model parameters. If you do not specify the instance resistance value, it is calculated from the model parameters.

This device is supported within altergroups.

If $R(\text{inst})$ is not given and $R(\text{model})$ is given,

$$R(\text{inst}) = R(\text{model}).$$

Otherwise,

$$R(\text{inst}) = R_{\text{sh}} * (L - 2 * \text{etchl}) / (W - 2 * \text{etch}).$$

If the polynomial coefficients vector (`coeffs=[c1 c2 ...]`) is specified, the resistor is nonlinear. When `nonlinform` is set to `g`, the conductance is

$$\begin{aligned} G(V) &= dI / dV \\ &= (1 + c_1 * V + c_2 * V^2 + \dots) / R(\text{inst}) \end{aligned}$$

where

$$V = V(t_1) - V(t_2)$$

Here V is the controlling voltage across the resistor. It is also the controlling voltage when the model parameter `polyarg` is set to `diff`. In this form, the physical resistor is symmetric with respect to $V(t_1)$ and $V(t_2)$. The branch current as a function of the applied voltage is given by

$$I(V) = (V / R(\text{inst})) * (1 + 1/2 * c_1 * V + 1/3 * c_2 * V^2 + \dots)$$

where c_k is the k th entry in the coefficient vector.

If the model parameter `polyarg` is set to `sum`, then the controlling voltage is defined as

$$V_{\text{sum}} = ((V(t_1) - V(t_0)) + (V(t_2) - V(t_0))) / 2$$

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Component Statements Part III

Here, V_{sum} is the controlling voltage between the resistor and the substrate, t_0 . In this case, the device becomes asymmetric with respect to $V(t_1)$ and $V(t_2)$. The branch current as a function of the applied voltage for this case is given by

$$I(V_{sum}) = (V / R(inst)) * (1 + c1 * V_{sum} + c2 * V_{sum}^2 + \dots)$$

The large-signal conductance is given by

$$G(V_{sum}) = I/V = (1 + c1 * V_{sum} + c2 * V_{sum}^2 + \dots) / R(inst)$$

Note, since the device is asymmetrical, the small-signal model is more complicated than a simple conductance.

When `nonlinform` is set to `r`, the resistance is

$$\begin{aligned} R(V) &= dV / dI \\ &= R(inst) * (1 + c1 * V + c2 * V^2 + \dots). \end{aligned}$$

The resistance as a function of temperature is given by:

$$R(T) = R(tnom) * [1 + tc1 * (T - tnom) + tc2 * (T - tnom)^2].$$

where

$$T = trise(inst) + temp$$

if `trise(inst)` is given, and

$$T = trise(model) + temp$$

otherwise.

If you do not specify the junction leakage current (i_s) and j_s is specified, the leakage current is calculated from j_s and the device dimensions.

$$i_s = j_s * 0.5 * (L - 2 * etch) * (W - 2 * etch)$$

If you specify the instance capacitance or the linear model capacitance, linear capacitors are used between t_1-t_0 and t_2-t_0 . Otherwise, nonlinear junction capacitors are used and the zero-bias capacitance values are calculated from the model parameters.

If `C(inst)` is not given and `C(model)` is given,

$$C(inst) = C(model).$$

Spectre Circuit Simulator Reference

Component Statements Part III

Otherwise,

$$C(\text{inst}) = 0.5 * C_j * (L - 2 * \text{etchlc}) * (W - 2 * \text{etchc}) + C_{jsw} * (W + L - 2 * \text{etchc} - 2 * \text{etchlc}).$$

If the capacitance is nonlinear, the temperature model for the junction capacitance is used. Otherwise, the following equation is used.

$$C(T) = C(\text{tnom}) * [1 + tc1c * (T - \text{tnom}) + tc2c * (T - \text{tnom})^2].$$

Sample Instance Statement

```
res1 (net9 vcc) resphy l=1e-3 w=2e-6
```

Sample Model Statement

```
model resphy phy_res rsh=85 tc1=1.53e-3 tc2=4.67e-7 etch=0 cj=1.33e-3 cjsw=3.15e-10 tc1c=9.26e-4
```

Instance Definition

```
Name 1 2 [0] ModelName parameter=value ...
```

Instance Parameters

1	r (Ω)	Resistance.
2	c (F)	Linear capacitance.
3	l (m)	Line length.
4	w (m)	Line width.
5	region=normal	Estimated operating region. Spectre outputs number (0-1) in a rawfile. Possible values are <code>normal</code> or <code>breakdown</code> .
6	tc1=0 1/C	Linear temperature coefficient of resistor.
7	tc2=0 C ⁻²	Quadratic temperature coefficient of resistor.
8	tc1c=0 1/C	Linear temperature coefficient of linear capacitor.
9	tc2c=0 C ⁻²	Quadratic temperature coefficient of linear capacitor.

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10 `trise (C)` Temperature rise from ambient.

11 `m=1` Multiplicity factor.

The `w` and `l` parameters are scaled by the option parameters `scale` and `scalem`. The values of `w` and `l` printed by Spectre are those given in the input file. These values may not have the correct units if the scaling factors are not unity. The actual effective resistor dimensions are stored in the output parameters. You can obtain these dimensions with the `info` statement. You can delete the diodes from the device by either setting `is=0` or `subtype=poly`. You can also set both `mj` and `mjsw` to zero to make the capacitance linear but still calculated from the device geometry. If `subtype=poly`, the linear capacitors will always be used irrespective of the values of `mj` and `mjsw`.

Model Definition

```
model modelName phy_res parameter=value ...
```

Model Parameters

Substrate type parameters

1 `subtype=p` Substrate type.
Possible values are `n`, `p` or `poly`.

Resistance parameters

2 `r=∞ Ω` Default resistance.

3 `rsh=∞ Ω/sqr` Sheet resistance.

4 `minr=0.1 Ω` Minimum resistance.

5 `coeffs=[...]` Vector of polynomial conductance or resistance coefficients.

6 `nonlinform=g` The form of the nonlinear resistance.
Possible values are `g` or `r`.

7 `polyarg=diff` Polynomial model argument type.
Possible values are `sum` or `diff`.

Spectre Circuit Simulator Reference

Component Statements Part III

Temperature effects parameters

8	<code>tc1=0</code>	$1/C$	Linear temperature coefficient of resistor.
9	<code>tc2=0</code>	C^{-2}	Quadratic temperature coefficient of resistor.
10	<code>tc1c=0</code>	C^{-2}	Linear temperature coefficient of linear capacitor.
11	<code>tc2c=0</code>	C^{-2}	Quadratic temperature coefficient of linear capacitor.
12	<code>tnom</code>	(C)	Parameters measurement temperature. Default set by <code>options</code> .
13	<code>trise=0</code>	C	Temperature rise from ambient.

Junction diode model parameters

14	<code>is</code>	(A)	Saturation current.
15	<code>js=0</code>	A/m^2	Saturation current density.
16	<code>n=1</code>		Emission coefficient.
17	<code>eg=1.11</code>	V	Band gap.
18	<code>x_{ti}=3</code>		Saturation current temperature exponent.
19	<code>imelt='imaxA'</code>		Explosion current, diode is linearized beyond this current to aid convergence.
20	<code>jmelt='jmeltA/m'²</code>		Explosion current density, diode is linearized beyond this current to aid convergence.
21	<code>imax=1</code>	A	Maximum current, currents above this limit generate a warning.
22	<code>jmax=1e8</code>	A/m^2	Maximum current density, currents above this limit generate a warning.
23	<code>dskip=yes</code>		Use simple piece-wise linear model for diode currents below $0.1 \cdot i_{abstol}$. Possible values are <code>no</code> or <code>yes</code> .

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24 $b_{vj}=\infty$ V Junction reverse breakdown voltage.

Junction capacitance model parameters

25 $c=0$ F Default linear capacitance.

26 $c_j=0$ F/m² Zero-bias junction bottom capacitance density.

27 $c_{jsw}=0$ F/m Zero-bias junction sidewall capacitance density.

28 $m_j=1/2$ Junction bottom grading coefficient.

29 $m_{jsw}=1/3$ Junction sidewall grading coefficient.

30 $p_b=0.8$ V Junction bottom built-in potential.

31 $p_{bsw}=0.8$ V Junction sidewall built-in potential.

32 $f_c=0.5$ Junction bottom capacitor forward-bias threshold.

33 $f_{csw}=0.5$ Junction sidewall capacitor forward-bias threshold.

34 $t_t=0$ s Transit time.

Device size parameters

35 $l=\infty$ m Default line length.

36 $w=1e-6$ m Default line width.

37 $etch=0$ m Narrowing due to etching.

38 $etchl=0$ m Length reduction due to etching.

39 $etchc=etch$ m Narrowing due to etching for capacitances.

40 $etchlc=etchl$ m Length reduction due to etching for capacitances.

41 $scaler=1$ Resistance scaling factor.

42 $scalec=1$ Capacitance scaling factor.

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Component Statements Part III

Noise model parameters

43	<code>kf=0</code>	Flicker (1/f) noise coefficient.
44	<code>af=1</code>	Flicker (1/f) noise exponent.
45	<code>wdexp=1</code>	Flicker (1/f) noise W exponent.
46	<code>ldexp=1</code>	Flicker (1/f) noise L exponent.
47	<code>weexp=0</code>	Flicker (1/f) noise W effective exponent.
48	<code>leexp=0</code>	Flicker (1/f) noise L effective exponent.
49	<code>fexp=1</code>	Flicker (1/f) noise frequency exponent.

Output Parameters

1	<code>leff (m)</code>	Effective line length.
2	<code>weff (m)</code>	Effective line width.
3	<code>iseff (A)</code>	Effective saturation current.
4	<code>reff (Ω)</code>	Effective resistance.
5	<code>ceff (F)</code>	Effective zero-bias capacitance.

Operating-Point Parameters

1	<code>subtype=p</code>	Substrate type. Possible values are <code>n</code> , <code>p</code> or <code>poly</code> .
2	<code>region=normal</code>	Estimated operating region. Spectre outputs number (0-1) in a rawfile. Possible values are <code>normal</code> or <code>breakdown</code> .
3	<code>i (A)</code>	Current through the resistor.
4	<code>capd1 (F)</code>	Capacitance at the positive node.
5	<code>capd2 (F)</code>	Capacitance at the negative node.

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6	id1 (A)	Current between nodes t1 and t0.
7	id2 (A)	Current between nodes t2 and t0.
8	res (Ω)	Resistance between nodes t1 and t2.
9	resd1 (Ω)	Resistance between nodes t1 and t0.
10	resd2 (Ω)	Resistance between nodes t2 and t0.
11	pwr (W)	Power at op point.

Parameter Index

In the following index, I refers to instance parameters, M refers to the model parameters section, O refers to the output parameters section, and OP refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of M-35 means the 35th model parameter.

af	M-44	i	OP-3	mjsw	M-29	subtype	OP-1
bvj	M-24	id1	OP-6	n	M-16	tc1	I-6
c	I-2	id2	OP-7	nonlinform	M-6	tc1	M-8
c	M-25	imax	M-21	pb	M-30	tc1c	I-8
capd1	OP-4	imelt	M-19	pbsw	M-31	tc1c	M-10
capd2	OP-5	is	M-14	polyarg	M-7	tc2	I-7
ceff	O-5	iseff	O-3	pwr	OP-11	tc2	M-9
cj	M-26	jmax	M-22	r	I-1	tc2c	I-9
cjsw	M-27	jmelt	M-20	r	M-2	tc2c	M-11
coeffs	M-5	js	M-15	reff	O-4	tnom	M-12
dskip	M-23	kf	M-43	region	I-5	trise	I-10
eg	M-17	l	M-35	region	OP-2	trise	M-13
etch	M-37	l	I-3	res	OP-8	tt	M-34
etchc	M-39	ldexp	M-46	resd1	OP-9	w	I-4
etchl	M-38	leexp	M-48	resd2	OP-10	w	M-36
etchlc	M-40	leff	O-1	rsh	M-3	wdexp	M-45

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Component Statements Part III

fc	M-32	m	I-11	scalec	M-42	weexp	M-47
fcsw	M-33	minr	M-4	scaler	M-41	weff	O-2
fexp	M-49	mj	M-28	subtype	M-1	xti	M-18

Independent Resistive Source (port)

Description

A port is a resistive source that is tied between `pos` and `neg`. It is equivalent to a voltage source in series with a resistor, and the reference resistance of the port is the value of the resistor. The DC value given for the port voltage specifies the DC voltage across the port when it is terminated in its reference resistance (in other words, the DC voltage of the internal voltage source is double the user specified DC value, `dc`). The same is true for the values for the transient, AC and PAC signals of the port. However, the amplitude of the sine wave in the transient and PAC analyses can alternatively be specified as the power in dBm delivered by the port when terminated with the reference resistance.

While generally useful as a stimulus in high frequency circuits, the port has three unique capabilities. First, it acts to define the ports of the circuit to the S-parameter analysis. Second, it has an intrinsic noise source, and so allows the noise analysis to directly compute the noise figure of the circuit. And finally, it is the only source for which the amplitude can be specified in terms of power.

This device is not supported within altergroup.

The value of the DC voltage as a function of the temperature is given by:

$$V(T) = V * [1 + tc1 * (T - tnom) + tc2 * (T - tnom)^2].$$

Sample Instance Statement

```
p20 (2 0) port num=2 r=50 type=pulse period=1e-9 rise=1e-10 fall=1e-10 vall=1
width=0.5n mag=1
```

Instance Definition

```
Name p n port parameter=value ...
```

Instance Parameters

1 `dc=0 v` DC value.

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Component Statements Part III

General waveform parameters

- 2 `type=dc` Waveform type.
Possible values are `dc`, `pulse`, `pwl`, `sine`, or `exp`.
- 3 `fundname` Name of the fundamental frequency. Must be specified if the source is active during a `pdisto` analysis or it is the active clock during an `envlp` analysis.
- 4 `delay=0 s` Waveform delay time.

Pulse waveform parameters

- 5 `val0=0 V` Zero value used in pulse and exponential waveforms.
- 6 `val1=1 V` One value used in pulse and exponential waveforms.
- 7 `period= ∞ s` Period of waveform.
- 8 `rise (s)` Rise time for pulse waveform (time for transition from `val0` to `val1`).
- 9 `fall (s)` Fall time for pulse waveform (time for transition from `val1` to `val0`).
- 10 `width= ∞ s` Pulse width (duration of `val1`).

PWL waveform parameters

- 11 `file` Name of file containing waveform.
- 12 `wave=[...]` Vector of time/value pairs that defines waveform.
- 13 `offset=0 V` DC offset for the PWL waveform.
- 14 `scale=1` Scale factor for the PWL waveform.
- 15 `pwldbms (dBm)` Power of PWL waveform in dBm (alternative to `scale`).
- 16 `stretch=1` Scale factor for time given for the PWL waveform.

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Component Statements Part III

- 17 `allbrkpts` All the points in the PWL waveform are breakpoints if set to yes. Default is yes if the number of points is less than 20. Possible values are `no` or `yes`.
- 18 `pwlperiod (s)` Period of the periodic PWL waveform.
- 19 `twidth=pwlperiod/1000 s` Transition width used when making PWL waveforms periodic.

Sinusoidal waveform parameters

- 20 `sinedc=dc V` DC level for sinusoidal waveforms.
- 21 `ampl=1 V` Peak amplitude of sinusoidal waveform.
- 22 `dbm (dBm)` Amplitude of sinusoidal waveform in dBm (alternative to `ampl`).
- 23 `freq=0 Hz` Frequency of sinusoidal waveform.
- 24 `sinephase=0 °` Phase of sinusoid when `t=delay`.
- 25 `ampl2=1 V` Peak amplitude of second sinusoidal waveform.
- 26 `dbm2 (dBm)` Amplitude of second sinusoidal waveform in dBm (alternative to `ampl2`).
- 27 `freq2=0 Hz` Frequency of second sinusoidal waveform.
- 28 `sinephase2=0 °` Phase of second sinusoid when `t=delay`.
- 29 `fundname2` Name of the fundamental frequency associated with `freq2`. Must be specified if `freq2` is used in a `pdisto` analysis.
- 30 `fmodindex=0` FM index of modulation for sinusoidal waveform.
- 31 `fmodfreq=0 Hz` FM modulation frequency for sinusoidal waveform.
- 32 `ammodindex=0` AM index of modulation for sinusoidal waveform.
- 33 `ammodfreq=0 Hz` AM modulation frequency for sinusoidal waveform.
- 34 `ammodphase=0 °` AM phase of modulation for sinusoidal waveform.

Spectre Circuit Simulator Reference

Component Statements Part III

35 `damp=0` 1/s Damping factor for sinusoidal waveform.

Exponential waveform parameters

36 `td1=0` s Rise start time for exponential wave.

37 `tau1` (s) Rise time constant for exponential wave.

38 `td2` (s) Fall start time for exponential wave.

39 `tau2` (s) Fall time constant for exponential wave.

Noise Parameters

40 `noisefile` Name of file containing excess spot noise data in the form of frequency-noise pairs.

41 `noisevec=[...]` V^2/Hz Excess spot noise as a function of frequency in the form of frequency-noise pairs.

42 `noisetemp` (C) Noise temperature of port. If not specified, the noise temperature is taken to be 290 K.

Port parameters

43 `r=50` Ω Reference resistance.

44 `num` Port number.

45 `m=1` Multiplicity factor.

Small signal parameters

46 `mag=0` V Small signal voltage.

47 `phase=0` $^\circ$ Small signal phase.

48 `xfmag=1` V/V Transfer function analysis magnitude.

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Component Statements Part III

- 49 `pacmag=0 V` Periodic AC analysis magnitude.
- 50 `pacdbm (dBm)` Periodic AC analysis magnitude in dBm (alternative to `pacmag`).
- 51 `pacphase=0 °` Periodic AC analysis phase.

Temperature effects parameters

- 52 `tc1=0 1/C` First order temperature coefficient.
- 53 `tc2=0 C-2` Second order temperature coefficient.
- 54 `tnom (C)` Parameters measurement temperature. Default set by options.

If you do not specify the DC value, it is assumed to be the `time=0` value of the waveform.

Sinusoidal waveform in the time interval $0 < t < \text{delay}$ has constant value equal to that at $t = \text{delay}$.

In DC analyses, the only active parameters are `dc`, `m`, and the temperature coefficient parameters. In AC analyses, the only active parameters are `m`, `mag` and `phase`. In transient analyses, all parameters are active except the small signal parameters and the noise parameters. The `type` parameter selects which type of waveform is generated. You may specify parameters for more than one waveform type, and use the `alter` statement to change the waveform type between analyses.

A vector of time-value pairs describes the piecewise linear waveform. As an alternative, you can read the waveform from a file. In this case, you give time-value pairs one pair per line with a space or tab between the time and the value.

If you set `allbrkpts` to `yes`, you force the simulator to place time points at each point specified in a PWL waveform during a transient analysis. This can be very expensive for waveforms with many points. If you set `allbrkpts` to `no`, Spectre inspects the waveform, looking for abrupt changes, and forces time points only at those changes.

The PWL waveform is periodic if you specify `pwlperiod`. If the value of the waveform specified is not exactly the same at both its beginning its end, then you must provide a nonzero value `twidth`. Before repeating, the waveform changes linearly in an interval of `twidth` from its value at $(\text{period} - \text{twidth})$ to its value at the beginning of the waveform. Thus `twidth` must always be less than `period`.

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Component Statements Part III

You can give the excess noise of the source as a file or specify it with a vector of frequency-noise pairs. For a file, give the frequency-noise pairs one pair per line with a space or tab between the frequency and noise values.

When computing the noise figure of a circuit driven at its input by a port, the noise temperature (noisetemp) of the port should be set to 16.85C (290K) in order to match the standard IEEE definition of noise figure. In addition, all other sources of noise in the port (noisefile and noisevec) should be disabled. If a noiseless port is desired, set the noise temperature to absolute zero or below, and do not specify a noise file or noise vector.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

allbrkpts	I-17	fmmodindex	I-30	pacphase	I-51	tc1	I-52
ammodfreq	I-33	freq	I-23	period	I-7	tc2	I-53
ammodindex	I-32	freq2	I-27	phase	I-47	td1	I-36
ammodphase	I-34	fundname	I-3	pwldbms	I-15	td2	I-38
ampl	I-21	fundname2	I-29	pwlperiod	I-18	tnom	I-54
ampl2	I-25	m	I-45	r	I-43	twidth	I-19
damp	I-35	mag	I-46	rise	I-8	type	I-2
dbm	I-22	noisefile	I-40	scale	I-14	val0	I-5
dbm2	I-26	noisetemp	I-42	sinedc	I-20	val1	I-6
dc	I-1	noisevec	I-41	sinephase	I-24	wave	I-12
delay	I-4	num	I-44	sinephase2	I-28	width	I-10
fall	I-9	offset	I-13	stretch	I-16	xfmag	I-48
file	I-11	pacdbm	I-50	tau1	I-37		
fmmodfreq	I-31	pacmag	I-49	tau2	I-39		

Poly-Si TFT (psitft)

Description

Poly-Si TFT model RPI level=16

This device is supported within altergroups.

Sample Instance Statement

```
m4 (0 2 1 1) nch w=2u l=0.8u
```

Sample Model Statement

```
model nch psitft type=p
```

Instance Definition

```
Name d g s [b] ModelName parameter=value ...
```

Instance Parameters

1	w (m)	Channel width.
2	l (m)	Channel length.
3	nrd (m/m)	Drain squares.
4	nrs (m/m)	Source squares.
5	m=1	Multiplicity factor (number of MOSFETs in parallel).
6	region=triode	Estimated operating region. Possible values are <code>off</code> , <code>triode</code> , <code>sat</code> , or <code>subth</code> .

Model Definition

```
model modelName psitft parameter=value ...
```

Model Parameters

Device type parameters

1 type=n Transistor type.
Possible values are n or p.

Drain current model parameters

2 vto=0 V Threshold voltage at zero body bias.

3 lambda=0.048 1/V Channel length modulation parameter.

4 tox=1e-7 m Gate oxide thickness.

5 eta=7 Subthreshold ideality factor.

6 asat=1 Proportionality constant of Vsat.

7 delta=4 Transition width parameter.

8 mus=1 cm²/V s Subthreshold mobility.

9 mu0=100 cm²/V s High field mobility.

10 mu1=0.004 cm²/V s Low field mobility parameter.

11 mmu=1.7 Low field mobility exponent.

12 vfb=-0.1 V Flat band voltage.

13 dd=1400e-10 m Vds field constant.

14 dg=2000e-10 m Vds field constant.

15 blk=0.001 Leakage barrier lowering constant.

16 clk=6 A/m Leakage scaling constant.

17 lkink=19e-6 m Kink effect constant.

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Component Statements Part III

18	mkink=1.3	Kink effect exponent.
19	vkink=9.1 V	Kink effect voltage.
20	rs=0 Ω	Source resistance.
21	rd=0 Ω	Drain resistance.
22	rsx=0 Ω	Resistance in series with Cgs.
23	rdx=0 Ω	Resistance in series with Cgd.
24	at=3e-8 m/V	DIBL parameter 1.
25	bt=1.9e-6 m/V	DIBL parameter 2.
26	eb=0.68 eV	Barrier height of diode.
27	i00=150 A/m	Reverse diode saturation current.
28	etac0	Capacitance subthreshold ideality factor at zero drain bias.
29	etac00=0 1/V	Capacitance subthreshold coefficient of drain bias.
30	mc=3	Capacitance knee shape parameter.
31	dvto=0 V/C	Temperature coefficient of VTO.
32	dmu1=0 cm ² /V s C	Temperature coefficient of MU1.
33	dasat=0 1/C	Temperature coefficient of ASAT.
34	lasat=0 m	Coefficient of length dependence of ASAT.
35	von=0 V	On-Voltage.
36	cgso=0 F/m	Gate-source overlap capacitance.
37	cgdo=0 F/m	Gate-drain overlap capacitance.
38	vsigma=0.2 V	Above threshold DIBL parameter.
39	vsigmat=1.7 V	Above threshold DIBL parameter.

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Component Statements Part III

40	<code>me=2.5</code>	Long channel saturation transition parameter.
41	<code>meta=1</code>	ETA floating-body parameter.
42	<code>ls=35e-9 m</code>	Channel-length modulation coefficient 1.
43	<code>vp=0.2 V</code>	Channel-length modulation coefficient 2.
44	<code>isubmod=0</code>	Channel-length modulation model version.
45	<code>vmax=4e4 m/s</code>	Carrier saturation velocity.
46	<code>theta=0 1/V</code>	Mobility modulation coefficient.
47	<code>mss=1.5</code>	Vdse transition parameter parameter.
48	<code>kss=0</code>	Fractions of the channel resistance coefficient.
49	<code>rsh=0 Ω/sqr</code>	Source/drain diffusion sheet resistance.
50	<code>capmod=0</code>	Intrinsic charge model.
51	<code>zeroc=0</code>	Zero gate-source (gate-drain) capacitance flag (Cgs=Cgd=0 if zeroc=1 and capmod=1).
52	<code>intdsnod=0</code>	Intrinsic source and drain nodes usage flag.

Temperature effects parameters

53	<code>tnom (C)</code>	Parameter measurement temperature.
54	<code>trise=0 C</code>	Temperature rise from ambient.

Operating region warning control parameters

55	<code>alarm=none</code>	Forbidden operating region. Possible values are none, off, triode, sat, subth, or rev.
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Spectre Circuit Simulator Reference

Component Statements Part III

Default device parameters

56	<code>w=3e-6 m</code>	Default channel width.
57	<code>l=3e-6 m</code>	Default channel length.
58	<code>nrd=1 m/m</code>	Default drain squares.
59	<code>nrs=1 m/m</code>	Default source squares.

Auto Model Selector parameters

60	<code>wmax=1 m</code>	Maximum channel width for which the model is valid.
61	<code>wmin=0 m</code>	Minimum channel width for which the model is valid.
62	<code>lmax=1 m</code>	Maximum channel length for which the model is valid.
63	<code>lmin=0 m</code>	Minimum channel length for which the model is valid.

Operating-Point Parameters

1	<code>type=n</code>	Transistor type. Possible values are <code>n</code> or <code>p</code> .
2	<code>region=triode</code>	Estimated operating region. Possible values are <code>off</code> , <code>triode</code> , <code>sat</code> , or <code>subth</code> .
3	<code>reversed</code>	Reverse mode indicator. Possible values are <code>no</code> or <code>yes</code> .
4	<code>ids (A)</code>	Resistive drain-to-source current.
5	<code>vgs (V)</code>	Gate-source voltage.
6	<code>vds (V)</code>	Drain-source voltage.
7	<code>vth (V)</code>	Threshold voltage.
8	<code>vdsat (V)</code>	Drain-source saturation voltage.
9	<code>gm (S)</code>	Common-source transconductance.

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Component Statements Part III

10	gds (S)	Common-source output conductance.
11	cgd (F)	Gate-drain capacitance.
12	cgs (F)	Gate-source capacitance.
13	ron (Ω)	On-resistance.
14	id (A)	Resistive drain current.
15	pwr (W)	Power at op point.
16	igs (A)	Gate-to-source current.
17	igd (A)	Gate-to-drain current.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

alarm	M-55	gm	OP-9	mmu	M-11	trise	M-54
asat	M-6	i00	M-27	mss	M-47	type	OP-1
at	M-24	id	OP-14	mu0	M-9	type	M-1
blk	M-15	ids	OP-4	mu1	M-10	vds	OP-6
bt	M-25	igd	OP-17	mus	M-8	vdsat	OP-8
capmod	M-50	igs	OP-16	nrd	I-3	vfb	M-12
cgd	OP-11	intdsnod	M-52	nrd	M-58	vgs	OP-5
cgdo	M-37	isubmod	M-44	nrs	I-4	vkink	M-19
cgs	OP-12	kss	M-48	nrs	M-59	vmax	M-45
cgso	M-36	l	M-57	pwr	OP-15	von	M-35
clk	M-16	l	I-2	rd	M-21	vp	M-43
dasat	M-33	lambda	M-3	rdx	M-23	vsigma	M-38
dd	M-13	lasat	M-34	region	OP-2	vsigmat	M-39

Spectre Circuit Simulator Reference Component Statements Part III

delta	M-7	lkink	M-17	region	I-6	vth	OP-7
dg	M-14	lmax	M-62	reversed	OP-3	vto	M-2
dmul	M-32	lmin	M-63	ron	OP-13	w	M-56
dvto	M-31	ls	M-42	rs	M-20	w	I-1
eb	M-26	m	I-5	rsh	M-49	wmax	M-60
eta	M-5	mc	M-30	rsx	M-22	wmin	M-61
etac0	M-28	me	M-40	theta	M-46	zeroc	M-51
etac00	M-29	meta	M-41	tnom	M-53		
gds	OP-10	mkink	M-18	tox	M-4		

Polynomial Voltage Controlled Current Source (pvccs)

Description

A vector of coefficients specifies the polynomial function that defines the relationship between the output current and the controlling voltages. You must specify at least one coefficient. Current exits the source node and enters the sink node.

This device is not supported within altergroup.

For a polynomial in M variables a_1, a_2, \dots, a_m , the polynomial function $F(a_0, a_1, \dots, a_m)$ is given by

$$\begin{aligned} F = & c_0 + c_1 * a_1 + c_2 * a_2 + \dots \\ & + c_{(m+1)} * a_1^2 + c_{(m+2)} * a_1 * a_2 + \dots \\ & + c_{(2m+1)} * a_2^2 + c_{(2m+2)} * a_2 * a_3 + \dots \end{aligned}$$

where the c_s are coefficients of the polynomial terms.

Sample Instance Statement

```
v2 (net1 0 net2 0) pvccs coeffs=[0 -2e-3 - 10e-3] gain=2 m=1
```

Instance Definition

```
Name sink src ps1 ns1 ... pvccs parameter=value ...
```


Spectre Circuit Simulator Reference

Component Statements Part III

Instance Parameters

1	<code>file</code>	File that contains nonzero polynomial coefficients.
2	<code>coeffs=[...]</code>	Polynomial coefficients. At least one must be given.
3	<code>gain=1</code>	Gain Parameter.
4	<code>m=1</code>	Multiplicity factor.
5	<code>min (A)</code>	Minimum output current.
6	<code>max (A)</code>	Maximum output current.
7	<code>abs=off</code>	Absolute output current. Possible values are <code>off</code> or <code>on</code> .
8	<code>delta=0</code>	Smoothing parameter. This may lead to circuit convergency. The smaller the delta is, the sharper the corner is.

Temperature effects parameters

9	<code>tc1=0 1/C</code>	Linear temperature coefficient.
10	<code>tc2=0 C⁻²</code>	Quadratic temperature coefficient.

Operating-Point Parameters

1	<code>i (A)</code>	Output current.
2	<code>v (V)</code>	Output voltage.
3	<code>pwr (W)</code>	Power dissipation.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the

Spectre Circuit Simulator Reference

Component Statements Part III

description for that parameter. For example, a reference of M-35 means the 35th model parameter.

abs	I-7	gain	I-3	min	I-5	v	OP-2
coeffs	I-2	i	OP-1	pwr	OP-3		
delta	I-8	m	I-4	tc1	I-9		
file	I-1	max	I-6	tc2	I-10		

Polynomial Voltage Controlled Voltage Source (pvcvs)

Description

A vector of coefficients specifies the polynomial function that defines the relationship between the output voltage and the controlling voltages. You must specify at least one coefficient. Current through the voltage source is calculated and is defined to be positive if it flows from the positive terminal, through the source, to the negative terminal.

This device is not supported within altergroup.

For a polynomial in M variables a_1, a_2, \dots, a_m , the polynomial function $F(a_0, a_1, \dots, a_m)$ is given by

$$\begin{aligned} F = & c_0 + c_1 * a_1 + c_2 * a_2 + \dots \\ & + c_{(m+1)} * a_1^2 + c_{(m+2)} * a_1 * a_2 + \dots \\ & + c_{(2m+1)} * a_2^2 + c_{(2m+2)} * a_2 * a_3 + \dots \end{aligned}$$

where the c_s are coefficients of the polynomial terms.

Sample Instance Statement

```
v1 (p 0 c1 0) pvcvs coeffs=[0 0 0 0.1 1 1] gain=1
```

Instance Definition

```
Name p n psl nsl ... pvcvs parameter=value ...
```

Spectre Circuit Simulator Reference

Component Statements Part III

Instance Parameters

1	<code>file</code>	File that contains nonzero polynomial coefficients.
2	<code>coeffs=[...]</code>	Polynomial coefficients. At least one must be given.
3	<code>gain=1</code>	Gain Parameter.
4	<code>m=1</code>	Multiplicity factor.
5	<code>min (V)</code>	Minimum output voltage.
6	<code>max (V)</code>	Maximum output voltage.
7	<code>abs=off</code>	Absolute output voltage. Possible values are <code>off</code> or <code>on</code> .
8	<code>delta=0</code>	Smoothing parameter. This may lead to circuit convergency. The smaller the delta is, the sharper the corner is.

Temperature effects parameters

9	<code>tc1=0 1/C</code>	Linear temperature coefficient.
10	<code>tc2=0 C⁻²</code>	Quadratic temperature coefficient.

Operating-Point Parameters

1	<code>i (A)</code>	Output current.
2	<code>v (V)</code>	Output voltage.
3	<code>pwr (W)</code>	Power dissipation.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the

Spectre Circuit Simulator Reference

Component Statements Part III

description for that parameter. For example, a reference of M-35 means the 35th model parameter.

abs	I-7	gain	I-3	min	I-5	v	OP-2
coeffs	I-2	i	OP-1	pwr	OP-3		
delta	I-8	m	I-4	tc1	I-9		
file	I-1	max	I-6	tc2	I-10		

Quantity Information (quantity)

Description

Quantities are used to hold information about particular types of signals, such as their units, absolute tolerances, and maximum allowed change per Newton iteration. Two predefined quantities are voltage and current. A node indicates the type of its signals by keeping pointers to quantities, thus an electrical node points to the voltage quantity for its value, and to the current for its residue. Since many electrical nodes point to the same quantities, changing an attribute on the quantity, such as the absolute tolerance, changes it for many nodes. Use this statement to create new quantities or to redefine properties of an existing quantity. Use the `node` statement to set the quantities for a particular node.

The predefined quantities are as follows:

I: Electrical current in Amperes.

Units = A

Absolute tolerance = 1 pA

Huge value = 45.036 MA

Blowup value = 1 GA

MMF: Magnetomotive force in Ampere-Turns.

Units = A*turn

Absolute tolerance = 1 pA*turn

Huge value = 45.036 MA*turn

Spectre Circuit Simulator Reference

Component Statements Part III

Blowup value = 1 GA*turn

Pwr: Power in Watts.

Units = W

Absolute tolerance = 1 nW

Huge value = 45.036 MW

Blowup value = 1 GW

Temp: Temperatures in Celsius.

Units = C

Absolute tolerance = 100 uC

Huge value = 45.036 MC

Blowup value = 1 GC

U: Unitless signals scaled to unity.

Absolute tolerance = 1e-06

Huge value = 45.036e+06

Blowup value = 1e+09

V: Electomotive force in Volts.

Units = V

Absolute tolerance = 1 uV

Maximum change = 300 mV if limit=delta

Huge value = 1 kV

Blowup value = 1 GV

Wb: Magnetic flux in Webers.

Units = Wb

Absolute tolerance = 1 nWb

Spectre Circuit Simulator Reference

Component Statements Part III

Huge value = 45.036 MWb

Blowup value = 1 GWb

This device is not supported within altergroup.

Sample Instance Statement

```
voltageQ quantity name="V" abstol=3u maxdelta=500m huge=10K blowup=1G
```

Instance Definition

```
Name quantity parameter=value ...
```

Instance Parameters

- | | | |
|---|-------------|--------------------------|
| 1 | name | Name. |
| 2 | description | Description of quantity. |
| 3 | units | Units. |

Newton parameters

- | | | |
|---|--------------------|--|
| 4 | abstol | Absolute tolerance. |
| 5 | maxdelta= ∞ | Maximum change allowed on a Newton iteration when
limit=delta. |
| 6 | huge=45.036e+06 | Maximum change allowed on a Newton iteration otherwise. |
| 7 | blowup=1e+09 | If a signal exceeds this value, the simulation will terminate with
an error. It is assumed that the circuit is unstable and is blowing
up. |

Diffusion Resistor Model (rdiff)

Description

The rdiff model is a diffusion resistor model, which accurately models the temperature, applied bias and back-bias dependencies of NWell, N+, and P+ resistors. It is described in the paper MODEL FOR DIFFUSION RESISTORS (NWell, N+, P+) USED IN CMOS IC DESIGNS by M.J.B.Bolt, FASELEC Process Development Group, PDG-93029, Modified 3rd May 1995.

Some extensions to that description are applied:

Appropriate model and instance parameter default values are used.

No clipping of parameters is performed. Parameter values are checked for validity. If invalid parameter values occur, the job is aborted with an error message.

For exact inverse behavior of the model in case of V_h less than V_l , the setting of $V_{bl} = \text{abs}(V_b - V_l)$ is replaced by $V_{bl} = \text{min}(\text{abs}(V_b - V_h), \text{abs}(V_b - V_l))$. Additionally, the direction of I_h is inverted in this case.

Note: In noise analysis, rdiff instances will not generate any contribution, since there are no noise sources included in the rdiff model.

(c) Philips Electronics N.V. 1993, 1995

This device is supported within altergroups.

This device is dynamically loaded from the shared object /vobs/spectre_dev/tools.sun4v/spectre/lib/cmi/3.0.doc/libphilips_sh.so

Sample Instance Statement

```
r2 (1 2 0) rdsn l=9u w=2u nb=0 m=1
```

Sample Model Statement

```
model rdsn rdiff level=1 tr=27 dta=0 rshr=2.5e3 wtol=0.22u rint=3.5u swvp=13.4u  
power=2 tcrl=1.5e-3 tcr2=1e-5 vpr=40
```

Instance Definition

```
Name h l [b] ModelName parameter=value ...
```

Spectre Circuit Simulator Reference

Component Statements Part III

Instance Parameters

1	<code>l=1.0 scale m</code>	Drawn length of resistor. Must be greater than zero. Scale set by option <code>scale</code> .
2	<code>w=1.0 scale m</code>	Drawn width of resistor. Must be greater than zero. Scale set by option <code>scale</code> .
3	<code>nb=0.0</code>	Number of bends in the resistor. Must be greater or equal zero.
4	<code>m=1.0</code>	Multiplicity factor. Must be greater than zero.

Model Definition

```
model modelName rdiff parameter=value ...
```

Model Parameters

1	<code>level=1.0</code>	Level of this model. Must be 1.
2	<code>tr (C)</code>	Reference temperature. Default set by option <code>tnom</code> .
3	<code>tref (C)</code>	Alias of <code>tr</code> . Default set by option <code>tnom</code> .
4	<code>tnom (C)</code>	Alias of <code>tr</code> . Default set by option <code>tnom</code> .
5	<code>dta=0 K</code>	Temperature offset of the device.
6	<code>trise=0 K</code>	Alias of <code>dta</code> .
7	<code>rshr=1.0e+3 Ω/sqr</code>	Sheet resistance at reference temperature. Must be greater than zero.
8	<code>wtol=0.0 m</code>	Offset between the drawn and effective resistor width.
9	<code>tcr1=0.0 1/K</code>	Linear temperature coefficient of the resistor.
10	<code>tcr2=0.0 1/K²</code>	Quadratic temperature coefficient of the resistor.
11	<code>vpr=100.0 V</code>	Reference Pinch-off voltage.
12	<code>swvp=0.0 V/m</code>	Coefficient of the width dependence of <code>vpr</code> .

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Component Statements Part III

13	<code>power=1.5</code>	Voltage exponent. Must be greater than zero.
14	<code>vdr=1.0 V</code>	Diffusion voltage at reference temperature.
15	<code>r_{int}=0.0 Ω m</code>	Interface resistance at reference temperature.
16	<code>tcr_{int1}=0.0 1/K</code>	Linear temperature coefficient of the interface resistor.

Output Parameters

1	<code>vd (V)</code>	Diffusion voltage. Must be greater than zero.
2	<code>r_{sh} (Ω/sqr)</code>	Sheet resistance. Must be greater than zero.
3	<code>vp (V)</code>	Pinch-off voltage. Must be greater than zero.
4	<code>r₀ (Ω)</code>	Zero bias resistance. Must be greater than zero.

Operating-Point Parameters

1	<code>v_{hl} (V)</code>	Absolute value of the applied bias across the resistor.
2	<code>v_{bl} (V)</code>	Absolute value of the back-bias across the resistor.
3	<code>i_h (A)</code>	DC current into the resistor.
4	<code>r (Ω)</code>	Actual resistance value.
5	<code>pwr (W)</code>	Power.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

<code>dta</code>	M-5	<code>r</code>	OP-4	<code>tcr_{int1}</code>	M-16	<code>v_{hl}</code>	OP-1
------------------	------------	----------------	-------------	---------------------------------	-------------	-----------------------------	-------------

Spectre Circuit Simulator Reference Component Statements Part III

ih	OP-3	r0	O-4	tnom	M-4	vp	O-3
l	I-1	rint	M-15	tr	M-2	vpr	M-11
level	M-1	rsh	O-2	tref	M-3	w	I-2
m	I-4	rshr	M-7	trise	M-6	wtol	M-8
nb	I-3	swvp	M-12	vbl	OP-2		
power	M-13	tcr1	M-9	vd	O-1		
pwr	OP-5	tcr2	M-10	vdr	M-14		

Four Terminal Relay (relay)

Description

The four-terminal relay is a voltage controlled relay tied between terminals $t1$ and $t2$. The voltage between terminals ps and ns controls the relay resistance. The relay resistance varies nonlinearly between r_{open} and r_{closed} , the open relay resistance and closed relay resistance, respectively. These resistance values correspond to control voltages of $vt1$ and $vt2$ respectively. The four parameters, $vt1$, $vt2$, r_{open} , and r_{closed} , can be instance or model parameters.

As an alternative, you can specify the threshold voltage v_{th} and a transition width $trans$ rather than specifying $vt1$ and $vt2$. These two parameters are then calculated from v_{th} and $trans$. If all four parameters are specified, v_{th} and $trans$ override $vt1$ and $vt2$. However, $vt1$ and $vt2$ values you specify on the instance override any model parameter specifications.

The final model parameter, $hysteresis$, designates a hysteresis with the on voltage shifted from v_{th} by an amount $hysteresis$ and the off voltage shifted by the same amount in the opposite direction. The direction of shift depends on the sign of $trans$ (or the relative magnitudes of $vt1$ and $vt2$): if $trans$ is positive, the on voltage shifts by $+hysteresis$; if $trans$ is negative (implying that the relay is "normally on"), the on-voltage shifts by $-hysteresis$.

This device is not supported within altergroup.

Operating conductance is calculated from the instance parameters as follows:

When V_c lies between $vt1$ and $vt2$,

$$G = G_{min} + (G_{min} - G_{max}) * [2 * (V_c - vt1)^3 - 3 * (vt2 - vt1) * (V_c - vt1)^2] / (vt2 - vt1)^3$$

Spectre Circuit Simulator Reference

Component Statements Part III

Otherwise, if $v_{t1} < v_{t2}$, then

$G = G_{min}$ for $V_c < v_{t1}$ and

$G = G_{max}$ for $V_c > v_{t2}$.

If $v_{t1} > v_{t2}$,

$G = G_{min}$ for $V_c > v_{t1}$ and

$G = G_{max}$ for $V_c < v_{t2}$.

where $G_{min} = 1 / r_{open}$, $G_{max} = 1 / r_{closed}$, and $V_c = V(ps) - V(ns)$.

Sample Instance Statement

```
rel1 (1 2 ps ns) my_relay ropen=1G rclosed=2
```

Sample Model Statement

```
model my_relay relay vt1=2.5 vt2=5 ropen=100M rclosed=0.1
```

Instance Definition

```
Name 1 2 ps ns ModelName parameter=value ...
```

```
Name 1 2 ps ns relay parameter=value ...
```

Instance Parameters

- | | | |
|---|--|---|
| 1 | <code>vt1 (V)</code> | Relay resistance is <code>ropen</code> at this voltage. |
| 2 | <code>vt2=vt1+1.0 V</code> | Relay resistance is <code>rclosed</code> at this voltage. |
| 3 | <code>ropen=∞ Ω</code> | Resistance of a fully open relay. |
| 4 | <code>rclosed=1.0 Ω</code> | Resistance of a fully closed relay. |
| 5 | <code>m=1.0</code> | Multiplicity factor. |
| 6 | <code>region=off</code> | Estimated operating region. Spectre outputs number (0-1) in a rawfile.
Possible values are <code>off</code> or <code>on</code> . |

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Component Statements Part III

Model Definition

model modelName relay parameter=value ...

Model Parameters

- 1 vt1 (V) Relay resistance is `ropen` at this voltage.
- 2 vt2=vt1+1.0 V Relay resistance is `rclosed` at this voltage.
- 3 ropen= ∞ Ω Resistance of a fully open relay.
- 4 rclosed=1.0 Ω Resistance of a fully closed relay.
- 5 hysteresis=0.0 V Switching Hysteresis.
- 6 vth=0.0 V Threshold Voltage.
- 7 trans=0.0 V Switch Transition Region Width.

Operating-Point Parameters

- 1 region=off Estimated operating region. Spectre outputs number (0-1) in a rawfile.
Possible values are `off` or `on`.
- 2 res (Ω) Relay resistance.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

hysteresis	M-5	region	I-6	ropen	M-3	vt2	I-2
m	I-5	region	OP-1	trans	M-7	vt2	M-2
rclosed	I-4	res	OP-2	vt1	M-1	vth	M-6

rclosed M-4 ropen I-3 vt1 I-1

Two Terminal Resistor (resistor)

Description

You can give the resistance explicitly or allow it to be computed from the physical length and width of the resistor. In either case, the resistance can be a function of temperature or applied voltage.

This device is supported within altergroups.

If $R(\text{inst})$ is not given,

$$R(\text{inst}) = R(\text{model})$$

if $R(\text{model})$ is given, and

$$R(\text{inst}) = R_{\text{sh}} * (L - 2 * \text{etchl}) / (W - 2 * \text{etch})$$

otherwise.

If the polynomial coefficients vector ($\text{coeffs}=[c1 \ c2 \ \dots]$) is specified, the resistor is nonlinear. When nonlinform is set to g , the conductance is

$$\begin{aligned} G(V) &= dI / dV \\ &= (1 + c1 * V + c2 * V^2 + \dots) / R(\text{inst}). \end{aligned}$$

The branch current as a function of applied voltage is

$$I(V) = (V / R(\text{inst})) * (1 + 1/2 * c1 * V + 1/3 * c2 * V^2 + \dots)$$

When nonlinform is set to r , the resistance is

$$\begin{aligned} R(V) &= dV / dI \\ &= R(\text{inst}) * (1 + c1 * V + c2 * V^2 + \dots). \end{aligned}$$

where c_k is the k th entry in the coefficient vector.

The value of the resistor as a function of the temperature is given by:

Spectre Circuit Simulator Reference

Component Statements Part III

$$R(T) = R(tnom) * [1 + tc1 * (T - tnom) + tc2 * (T - tnom)^2]$$

where

$$T = trise(inst) + temp$$

if $trise(inst)$ is given, and

$$T = trise(model) + temp$$

otherwise.

If you specify capacitance the model is a wire RC model. You can give the capacitance explicitly or allow it to be computed from the physical length and width of the resistor. The model parameter `cratio` can be used to allocate the parasitic capacitance of the wire element between the models input capacitor and output capacitor.

If $C(inst)$ is not given and $C(model)$ is given,

$$C(inst) = C(model).$$

Otherwise,

$$C(inst) = Cj * (L - 2 * etchl) * (W - 2 * etchw) + 2 * Cjsw * (W + L - 2 * etchl - 2 * etchw).$$

The parasitic capacitance assigned to the input and output nodes is given by:

$$C(1) = C(inst) * cratio$$

$$C(2) = C(inst) * (1 - cratio).$$

The value of each capacitor as a function of the temperature is given by:

$$C(T) = C(tnom) * [1 + tc1c * (T - tnom) + tc2c * (T - tnom)^2].$$

Sample Instance Statement

without model:

```
r1 (1 2) resistor r=1.2K m=2
```

with model:

```
r1 (1 2) resmod l=8u w=1u
```

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Sample Model Statement

```
model resmod resistor rsh=150 l=2u w=2u etch=0.05u tc1=0.1 tnom=27 kf=1
```

Instance Definition

```
Name 1 2 [0] ModelName parameter=value ...
```

```
Name 1 2 [0] resistor parameter=value ...
```

Instance Parameters

1	<code>r</code> (Ω)	Resistance.
2	<code>l</code> (m)	Resistor length.
3	<code>w</code> (m)	Resistor width.
4	<code>m=1</code>	Multiplicity factor.
5	<code>scale=1</code>	Scale factor.
6	<code>resform</code>	Use the resistance form for this instance. Default is <code>yes</code> if <code>r < thresh</code> . Possible values are <code>no</code> or <code>yes</code> .
7	<code>tc1=0</code> $1/C$	Linear temperature coefficient.
8	<code>tc2=0</code> C^{-2}	Quadratic temperature coefficient.
9	<code>trise</code> (C)	Temperature rise from ambient.
10	<code>isnoisy=yes</code>	Should resistor generate noise. Possible values are <code>no</code> or <code>yes</code> .
11	<code>c</code> (F)	Capacitance.
12	<code>tc1c=0</code> $1/C$	Linear temperature coefficient of capacitor.
13	<code>tc2c=0</code> C^{-2}	Quadratic temperature coefficient of capacitor.

The instance parameter `scale`, if specified, overrides the value given by the `option` parameter `scale`. The `w` and `l` parameters are scaled by the resulting `scale`, and the `option` parameter `scalem`. The parameters `w` and `l` are scaled also by the model parameter `shrink`. The values of `w` and `l` printed out by `spectre` are those given in the input, and these

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values might not have the correct units if the scaling factors are not unity. The actual effective resistor dimensions are stored in the output parameters. You can obtain these dimensions by using the info statement.

Model Definition

```
model modelName resistor parameter=value ...
```

Model Parameters

Resistance parameters

- | | | |
|---|------------------------------|--|
| 1 | <code>r=∞ Ω</code> | Default resistance. |
| 2 | <code>rsh=∞ Ω/sqr</code> | Sheet resistance. |
| 3 | <code>thresh=1.0e-3 Ω</code> | Resistances smaller than this will use the resistance form, as opposed to the standard conductance form. |

Resistor size parameters

- | | | |
|---|------------------------|---|
| 4 | <code>l=∞ m</code> | Default resistor length. |
| 5 | <code>w=1e-6 m</code> | Default resistor width. |
| 6 | <code>etch=0 m</code> | Width narrowing due to etching per side. |
| 7 | <code>etchl=0 m</code> | Length narrowing due to etching per side. |
| 8 | <code>scaler=1</code> | Resistance scaling factor. |

Temperature effects parameters

- | | | |
|----|-----------------------------------|---|
| 9 | <code>tc1=0 1/C</code> | Linear temperature coefficient. |
| 10 | <code>tc2=0 C⁻²</code> | Quadratic temperature coefficient. |
| 11 | <code>tnom (C)</code> | Parameters measurement temperature. Default set by <code>options</code> . |

Spectre Circuit Simulator Reference

Component Statements Part III

12 `trise=0 C` Default temperature rise from ambient.

Nonlinear resistance

13 `coeffs=[...]` Vector of polynomial conductance coefficients.

14 `nonlinform=g` The form of the nonlinear resistance.
Possible values are `g` or `r`.

15 `symmetric=none` Use symmetric resistor model.
Possible values are `none` or `absolute`.

Noise model parameters

16 `kf=0` Flicker (1/f) noise coefficient.

17 `af=2` Flicker (1/f) noise exponent.

18 `wdexp=1` Flicker (1/f) noise W exponent.

19 `ldexp=1` Flicker (1/f) noise L exponent.

20 `weexp=0` Flicker (1/f) noise W effective exponent.

21 `leexp=0` Flicker (1/f) noise L effective exponent.

22 `fexp=1` Flicker (1/f) noise frequency exponent.

DC-mismatch model parameters

23 `mr=0.0` Resistor mismatch dependence.

24 `mr1=0.0` Resistor mismatch length dependence.

25 `mr1p=0.0` Resistor mismatch length power dependence.

26 `mrw=0.0` Resistor mismatch width dependence.

27 `mrwp=0.0` Resistor mismatch width power dependence.

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Component Statements Part III

- 28 `mr1w1=0.0` Resistor mismatch area 1 dependence.
- 29 `mr1w1p=0.0` Resistor mismatch area 1 power dependence.
- 30 `mr1w2=0.0` Resistor mismatch area 2 dependence.
- 31 `mr1w2p=0.0` Resistor mismatch area 2 power dependence.

Wire RC parameters

- 32 `c=0 F` Default capacitance.
- 33 `cj=0 F/m2` Bottomwall capacitance.
- 34 `cjsw=0 F/m` Sidewall fringing capacitance.
- 35 `thick=0 m` Dielectric thickness.
- 36 `di=0` Relative dielectric constant.
- 37 `cratio=0.5` Cratio.
- 38 `tc1c=0 C-2` Linear temperature coefficient of capacitor.
- 39 `tc2c=0 C-2` Quadratic temperature coefficient of capacitor.
- 40 `shrink=1` Shrink factor.
- 41 `scalec=1` Capacitance scaling factor.

The instance parameter `resform` and the model parameter `thresh` control whether a resistor is formulated in the standard conductance form, or in the resistance form. If the value of the resistor is smaller than `thresh`, Spectre uses the resistance form; otherwise it uses the conductance form. If `resform` is set on an instance, it overrides the `thresh` parameter. The resistance form is appropriate for very small resistances and the conductance form is intended for larger resistances. Using the conductance form for very small resistances or the resistance form for very large resistances can cause convergence problems.

With the resistance form, the resistance can be zero; with the conductance form, the resistance can be infinite. The resistance form is less efficient than the conductance form. You cannot change the formulation of a resistor once it has been determined. Spectre makes this choice by comparing the initial value of the resistance to `thresh`.

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Component Statements Part III

If the polynomial coefficients vector is specified and `symmetric=absolute`, the resistor model is symmetric. When `nonlinform` is set to `g`, the conductance is

$$G(V) = dI / dV$$
$$= (1 + c1 * |V| + c2 * |V|^2 + ...) / R(\text{inst})$$

where $|V| = \text{ABS}(V(1) - V(2))$ and c_k is the k th entry in the coefficient vector.

When `nonlinform` is set to `r`, the resistance is

$$R(V) = dV / dI$$
$$= R(\text{inst}) * (1 + c1 * |V| + c2 * |V|^2 + ...)$$

where $|V| = \text{ABS}(V(1) - V(2))$.

Modeling AC resistance

In certain situations, a part of a circuit that is required to calculate the DC operating point needs to be removed during a subsequent AC analysis or visa versa. An example of a situation in which this occurs is when measuring the loop gain of a feedback amplifier. In this case the feedback loop must be removed when computing the AC response of the amplifier. In Spectre, the most accurate method of doing this is to use an ideal switch component (see `spectre -h switch`), e.g.

```
Vin (pin 0) vsource mag=1
```

```
OA1 (pin nin out) opamp
```

```
Sw1 (nin out 0) switch position=1 ac_position=2
```

```
LoopGain ac start=1 stop=1MHz
```

Another possibility is that the resistance of an instance changes from one analysis to another. The following subcircuit models a resistance whose value is given by the parameter `rac` during AC analyses, and `rdc` for all other analyses.

```
subckt ac_res (a b)
```

```
parameters rdc=1 rac=2
```

```
R1 (a i) resistor r=rdc
```

```
Rac (i b) resistor r=rac-rdc
```

Spectre Circuit Simulator Reference Component Statements Part III

Sw (i b) switch position=1 ac_position=0

ends ac_res

Output Parameters

- 1 leff (m) Effective resistor length.
- 2 weff (m) Effective resistor width.
- 3 reff (Ω) Effective resistance.
- 4 ceff (F) Effective capacitance.

Operating-Point Parameters

- 1 v (V) Voltage at operating point.
- 2 i (A) Current through the resistor.
- 3 res (Ω) Resistance at op point.
- 4 pwr (W) Power dissipation.

Parameter Index

In the following index, I refers to instance parameters, M refers to the model parameters section, O refers to the output parameters section, and OP refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of M-35 means the 35th model parameter.

af	M-17	l	M-4	r	I-1	tc2	I-8
c	I-11	ldexp	M-19	r	M-1	tc2c	I-13
c	M-32	leexp	M-21	reff	O-3	tc2c	M-39
ceff	O-4	leff	O-1	res	OP-3	thick	M-35
cj	M-33	m	I-4	resform	I-6	thresh	M-3
cjsw	M-34	mr	M-23	rsh	M-2	tnom	M-11

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coeffs	M-13	mrl	M-24	scale	I-5	trise	I-9
cratio	M-37	mrlp	M-25	scalec	M-41	trise	M-12
di	M-36	mrlw1	M-28	scaler	M-8	v	OP-1
etch	M-6	mrlw1p	M-29	shrink	M-40	w	M-5
etchl	M-7	mrlw2	M-30	symmetric	M-15	w	I-3
fexp	M-22	mrlw2p	M-31	tc1	M-9	wdexp	M-18
i	OP-2	mrw	M-26	tc1	I-7	weexp	M-20
isnoisy	I-10	mrwp	M-27	tc1c	I-12	weff	O-2
kf	M-16	nonlinform	M-14	tc1c	M-38		
l	I-2	pwr	OP-4	tc2	M-10		

s-Domain Linear Current Controlled Current Source (scccs)

Description

The device output is defined through a transfer function given as a ratio of two polynomials in the complex variable s . Polynomials can be specified in terms of either coefficients or roots. The roots of the numerator are the zeros of the transfer function and the roots of the denominator are the poles.

To specify the polynomial in terms of the coefficients, you enter them as a vector in ascending order of the power of the variable s , starting from the constant term. For example, to specify a denominator of $3s^2 + 4s + 1$, use `denom=[1 4 3]`.

To specify a polynomial in terms of its roots, you give the roots as a vector of complex frequencies (frequencies should be in radians/second). You must give both the real and imaginary parts of the root, even when the root is real. For the transfer function to be stable, all poles must have negative real values. When specifying a complex root, you should also specify its complex conjugate. However, if you omit the conjugate root, Spectre will supply the missing root and print a warning that a missing root was supplied. The order of the roots is not important. For example, to specify poles of $s = -1$, $s = 4j$, $s = -4j$, $s = -2 + 2j$, and $s = -2 - 2j$; use `poles=[-1 0 0 4 0 -4 -2 2 -2 -2]`.

Either the numerator or the denominator specification can be omitted. An omitted denominator or numerator is taken to be 1.

Spectre Circuit Simulator Reference

Component Statements Part III

The parameter `gain` is interpreted either as the DC gain or, if the function has zeros or poles at the origin, as a constant factor.

This device is not supported within `altergroup`.

Sample Instance Statement

```
l1 (2 1) inductor l=15
scl (1 0) scccs probe=l1 zeros=[0 6 0 -6 2 -8 2 8] poles=[-1 0 0 64 0 -64 -2 8 -2 -8]
```

Instance Definition

```
Name sink src scccs parameter=value ...
```

Instance Parameters

1	<code>probe</code>	Device through which the controlling current flows.
2	<code>port=0</code>	Index of the probe port through which the controlling current flows.
3	<code>gain=1</code>	Transfer function gain.
4	<code>numer=[...]</code>	Vector of numerator coefficients.
5	<code>denom=[...]</code>	Vector of denominator coefficients.
6	<code>zeros=[...]</code>	Vector of complex zeros.
7	<code>poles=[...]</code>	Vector of complex poles.
8	<code>m=1</code>	Multiplicity factor.

Operating-Point Parameters

1	<code>i (A)</code>	Input current.
2	<code>v (V)</code>	Output voltage.
3	<code>pwr (W)</code>	Power dissipation.

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Component Statements Part III

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

denom	I-5	m	I-8	port	I-2	v	OP-2
gain	I-3	numer	I-4	probe	I-1	zeros	I-6
i	OP-1	poles	I-7	pwr	OP-3		

s-Domain Current Controlled Voltage Source (sccvs)

Description

The device output is defined through a transfer function given as a ratio of two polynomials in the complex variable s . Polynomials can be specified in terms of either coefficients or roots. The roots of the numerator are the zeros of the transfer function and the roots of the denominator are the poles.

To specify the polynomial in terms of the coefficients, you enter them as a vector in ascending order of the power of the variable s , starting from the constant term. For example, to specify a denominator of $3s^2 + 4s + 1$, use `denom=[1 4 3]`.

To specify a polynomial in terms of its roots, you give the roots as a vector of complex frequencies (frequencies should be in radians/second). You must give both the real and imaginary parts of the root, even when the root is real. For the transfer function to be stable, all poles must have negative real values. When specifying a complex root, you should also specify its complex conjugate. However, if you omit the conjugate root, Spectre will supply the missing root and print a warning that a missing root was supplied. The order of the roots is not important. For example, to specify poles of $s = -1$, $s = 4j$, $s = -4j$, $s = -2 + 2j$, and $s = -2 - 2j$; use `poles=[-1 0 0 4 0 -4 -2 2 -2 -2]`.

Either the numerator or the denominator specification can be omitted. An omitted denominator or numerator is taken to be 1.

The parameter `gain` is interpreted either as the DC gain or, if the function has zeros or poles at the origin, as a constant factor.

Spectre Circuit Simulator Reference

Component Statements Part III

This device is not supported within altergroup.

Sample Instance Statement

```
myv (1 0) vsource type=sine freq=10K
scc1 (2 0) sccvs probe=myv gain=0.5 numer=[2] denom=[5]
```

Instance Definition

```
Name p n sccvs parameter=value ...
```

Instance Parameters

1	probe	Device through which the controlling current flows.
2	port=0	Index of the probe port through which the controlling current flows.
3	gain=1	Transfer function gain.
4	numer=[...]	Vector of numerator coefficients.
5	denom=[...]	Vector of denominator coefficients.
6	zeros=[...]	Vector of complex zeros.
7	poles=[...]	Vector of complex poles.
8	m=1	Multiplicity factor.

Operating-Point Parameters

1	i (A)	Output current.
2	v (V)	Output voltage.
3	pwr (W)	Power dissipation.

Spectre Circuit Simulator Reference

Component Statements Part III

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

denom	I-5	m	I-8	port	I-2	v	OP-2
gain	I-3	numer	I-4	probe	I-1	zeros	I-6
i	OP-1	poles	I-7	pwr	OP-3		

s-Domain Linear Voltage Controlled Current Source (svccs)

Description

The device output is defined through a transfer function given as a ratio of two polynomials in the complex variable s . Polynomials can be specified in terms of either coefficients or roots. The roots of the numerator are the zeros of the transfer function and the roots of the denominator are the poles.

To specify the polynomial in terms of the coefficients, you enter them as a vector in ascending order of the power of the variable s , starting from the constant term. For example, to specify a denominator of $3s^2 + 4s + 1$, use `denom=[1 4 3]`.

To specify a polynomial in terms of its roots, you give the roots as a vector of complex frequencies (frequencies should be in radians/second). You must give both the real and imaginary parts of the root, even when the root is real. For the transfer function to be stable, all poles must have negative real values. When specifying a complex root, you should also specify its complex conjugate. However, if you omit the conjugate root, Spectre will supply the missing root and print a warning that a missing root was supplied. The order of the roots is not important. For example, to specify poles of $s = -1$, $s = 4j$, $s = -4j$, $s = -2 + 2j$, and $s = -2 - 2j$; use `poles=[-1 0 0 4 0 -4 -2 2 -2 -2]`.

Either the numerator or the denominator specification can be omitted. An omitted denominator or numerator is taken to be 1.

Spectre Circuit Simulator Reference

Component Statements Part III

The parameter `gain` is interpreted either as the DC gain or, if the function has zeros or poles at the origin, as a constant factor.

This device is not supported within `altergroup`.

Sample Instance Statement

```
s2 (1 0 control 0) svccs gain=0.4 numer=[2 3] denom=[4 5 1]
```

Instance Definition

```
Name sink src ps ns svccs parameter=value ...
```

Instance Parameters

- | | | |
|---|--------------------------|-------------------------------------|
| 1 | <code>gain=1</code> | Transfer function gain. |
| 2 | <code>numer=[...]</code> | Vector of numerator coefficients. |
| 3 | <code>denom=[...]</code> | Vector of denominator coefficients. |
| 4 | <code>zeros=[...]</code> | Vector of complex zeros. |
| 5 | <code>poles=[...]</code> | Vector of complex poles. |
| 6 | <code>m=1</code> | Multiplicity factor. |

Operating-Point Parameters

- | | | |
|---|----------------------|--------------------|
| 1 | <code>i (A)</code> | Output current. |
| 2 | <code>v (V)</code> | Output voltage. |
| 3 | <code>pwr (W)</code> | Power dissipation. |

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the

Spectre Circuit Simulator Reference

Component Statements Part III

description for that parameter. For example, a reference of M-35 means the 35th model parameter.

denom	I-3	m	I-6	pwr	OP-3
gain	I-1	numer	I-2	v	OP-2
i	OP-1	poles	I-5	zeros	I-4

s-Domain Voltage Controlled Voltage Source (svcv)

Description

The device output is defined through a transfer function given as a ratio of two polynomials in the complex variable s . Polynomials can be specified in terms of either coefficients or roots. The roots of the numerator are the zeros of the transfer function and the roots of the denominator are the poles.

To specify the polynomial in terms of the coefficients, you enter them as a vector in ascending order of the power of the variable s , starting from the constant term. For example, to specify a denominator of $3s^2 + 4s + 1$, use `denom=[1 4 3]`.

To specify a polynomial in terms of its roots, you give the roots as a vector of complex frequencies (frequencies should be in radians/second). You must give both the real and imaginary parts of the root, even when the root is real. For the transfer function to be stable, all poles must have negative real values. When specifying a complex root, you should also specify its complex conjugate. However, if you omit the conjugate root, Spectre will supply the missing root and print a warning that a missing root was supplied. The order of the roots is not important. For example, to specify poles of $s = -1$, $s = 4j$, $s = -4j$, $s = -2 + 2j$, and $s = -2 - 2j$; use `poles=[-1 0 0 4 0 -4 -2 2 -2 -2]`.

Either the numerator or the denominator specification can be omitted. An omitted denominator or numerator is taken to be 1.

The parameter `gain` is interpreted either as the DC gain or, if the function has zeros or poles at the origin, as a constant factor.

This device is not supported within altergroup.

Sample Instance Statement

```
e1 (1 0 control 0) svccs gain=5 poles=[-1 0 1 0] zero=[0 0 1 0]
```

Spectre Circuit Simulator Reference

Component Statements Part III

Instance Definition

Name p n ps ns svcvs parameter=value ...

Instance Parameters

- | | | |
|---|-------------|-------------------------------------|
| 1 | gain=1 | Transfer function gain. |
| 2 | numer=[...] | Vector of numerator coefficients. |
| 3 | denom=[...] | Vector of denominator coefficients. |
| 4 | zeros=[...] | Vector of complex zeros. |
| 5 | poles=[...] | Vector of complex poles. |
| 6 | m=1 | Multiplicity factor. |

Operating-Point Parameters

- | | | |
|---|---------|--------------------|
| 1 | i (A) | Output current. |
| 2 | v (V) | Output voltage. |
| 3 | pwr (W) | Power dissipation. |

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

denom	I-3	m	I-6	pwr	OP-3
gain	I-1	numer	I-2	v	OP-2
i	OP-1	poles	I-5	zeros	I-4

Ideal Switch (switch)

Description

Ideal switch is a single-pole multiple-throw switch with infinite `off` resistance and zero `on` resistance. The switch is provided to allow you to reconfigure your circuit between analyses. You can only change the switch state between analyses (using the `alter` statement), not during an analysis.

When the switch is set to position 0 it is open. In other words, no terminal is connected to any other. When the switch is set to position 1, terminal 1 is connected to terminal 0, and all others are unconnected. When the position is set to 2, terminal 2 is connected to terminal 0, etc.

An offset voltage is supported. It is placed in series with the common terminal. The negative side of the source is connected to the common terminal.

The switch can change its position based on which analysis type is being performed using the `xxx_position` parameters. This feature should be used carefully. Careless use can generate discontinuities that result in convergence problems. Once an analysis specific position has been specified using `xxx_position`, it will always dominate over a position given with the `position` parameter. To disable an analysis specific position, alter it to its default value of unspecified.

This device is not supported within `altergroup`.

Sample Instance Statement

```
sw1 (t1 t2 t3) switch dc_position=0 ac_position=1 tran_position=2
```

Instance Definition

```
Name t0 t1 ... switch parameter=value ...
```

Instance Parameters

- | | |
|------------------------------|---|
| 1 <code>position=0</code> | Switch position (0, 1, 2, ...). |
| 2 <code>dc_position</code> | Position to which switch is set at start of DC analysis. |
| 3 <code>ac_position</code> | Position to which switch is set at start of AC analysis. |
| 4 <code>tran_position</code> | Position to which switch is set at start of transient analysis. |

Spectre Circuit Simulator Reference

Component Statements Part III

- | | | |
|---|--------------------------|--|
| 5 | <code>ic_position</code> | Position to which switch is set at start of IC analysis (precedes transient analysis). |
| 6 | <code>offset=0</code> | Offset voltage in series with common terminal. |
| 7 | <code>m=1.0</code> | Multiplicity factor. |

Output Parameters

- | | | |
|---|-------------------------------|--------------------------|
| 1 | <code>present_position</code> | Current switch position. |
|---|-------------------------------|--------------------------|

Parameter Index

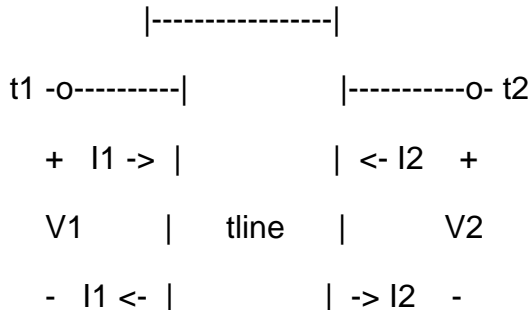
In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

<code>ac_position</code> I-3	<code>ic_position</code> I-5	<code>offset</code> I-6	<code>present_position</code> O-1
<code>dc_position</code> I-2	<code>m</code> I-7	<code>position</code> I-1	<code>tran_position</code> I-4

Transmission Line (tline)

Description

Lossy or lossless transmission line.



Spectre Circuit Simulator Reference

Component Statements Part III



The lossy transmission line model includes dielectric and conductor loss effects. The conductor loss includes skin effect assuming finite or infinite conductor thickness.

Only the odd mode is modeled, so only the voltage difference across each port is important. (The absolute voltage of each terminal is not significant.) Also, the current into one node of a port exactly equals the current leaving the other node of the port.

This device is supported within altergroups.

Sample Instance Statement

```
t1 (1 0 2 0) lmodel z0=100
```

Sample Model Statement

```
model lmodel tline f=10M z0=50 alphac=8501 fc=10M dcr=88
```

Instance Definition

```
Name t1 b1 t2 b2 ModelName parameter=value ...
```

```
Name t1 b1 t2 b2 tline parameter=value ...
```

Instance Parameters

- | | | |
|---|----------------------|--|
| 1 | <code>z0=50 Ω</code> | Characteristic impedance of lossless line. |
| 2 | <code>td (s)</code> | Time delay of a lossless line in seconds, a measure of the electrical length. |
| 3 | <code>f (Hz)</code> | Reference frequency (used in conjunction to the normalized length to specify electrical length of line). |
| 4 | <code>n1=0.25</code> | Normalized electrical length in wavelengths at <code>f</code> of a lossless line. |
| 5 | <code>vel=1</code> | Propagation velocity of the line given as a multiple of <code>c</code> , the speed of light in free space. (<code>vel <= 1</code>). |
| 6 | <code>len=0 m</code> | Physical length (used with <code>vel</code> to specify electrical length of line). |

Spectre Circuit Simulator Reference

Component Statements Part III

7 `m=1` Multiplicity factor.

Conductor loss parameters

8 `corner=0 Hz` Corner frequency for skin effect, frequency where skin depth equals the conductors wall thickness.

9 `dcr=0 Ω /m` DC series resistance per unit length.

10 `fc (Hz)` Conductor loss measurement frequency (use with `r`, `qc`, or `alphac`).

11 `r=0 Ω /m` Conductor (series) resistance per unit length at `fc`.

12 `alphac=0 dB/m` Conductor loss at `fc` (low loss approximation).

13 `qc= ∞` Conductor loss quality factor at `fc` (low loss approximation).

Dielectric loss parameters

14 `fd (Hz)` Dielectric loss measurement frequency (use with `qd`).

15 `g=0 S/m` Dielectric (shunt) conductance per unit length.

16 `alphad=0 dB/m` Dielectric loss (low loss approximation).

17 `qd= ∞` Dielectric loss quality factor at `fd` (low loss approximation).

Model Definition

```
model modelName tline parameter=value ...
```

Model Parameters

1 `z0=50 Ω` Characteristic impedance of lossless line.

2 `f (Hz)` Reference frequency (used in conjunction to the normalized length to specify electrical length of line).

3 `vel=1` Propagation velocity of the line given as a multiple of `c`, the speed of light in free space. (`vel <= 1`).

Spectre Circuit Simulator Reference

Component Statements Part III

Conductor loss parameters

4	<code>corner=0</code> Hz	Corner frequency for skin effect, frequency where skin depth equals the conductors wall thickness.
5	<code>dcr=0</code> Ω/m	DC series resistance per unit length.
6	<code>fc</code> (Hz)	Conductor loss measurement frequency (use with <code>r</code> , <code>qc</code> , or <code>alphac</code>).
7	<code>r=0</code> Ω/m	Conductor (series) resistance per unit length at <code>fc</code> .
8	<code>alphac=0</code> dB/m	Conductor loss at <code>fc</code> (low loss approximation).
9	<code>qc=∞</code>	Conductor loss quality factor at <code>fc</code> (low loss approximation).

Dielectric loss parameters

10	<code>fd</code> (Hz)	Dielectric loss measurement frequency (use with <code>gd</code>).
11	<code>g=0</code> S/m	Dielectric (shunt) conductance per unit length.
12	<code>alphad=0</code> dB/m	Dielectric loss (low loss approximation).
13	<code>gd=∞</code>	Dielectric loss quality factor at <code>fd</code> (low loss approximation).

Lossless Case

The lossless transmission line is specified with parameters z_0 and t_d . The device behavior is then:

$$V_1(t) - z_0 I_1(t) = V_2(t-t_d) + z_0 I_2(t-t_d)$$

and

$$V_2(t) - z_0 I_2(t) = V_1(t-t_d) + z_0 I_1(t-t_d).$$

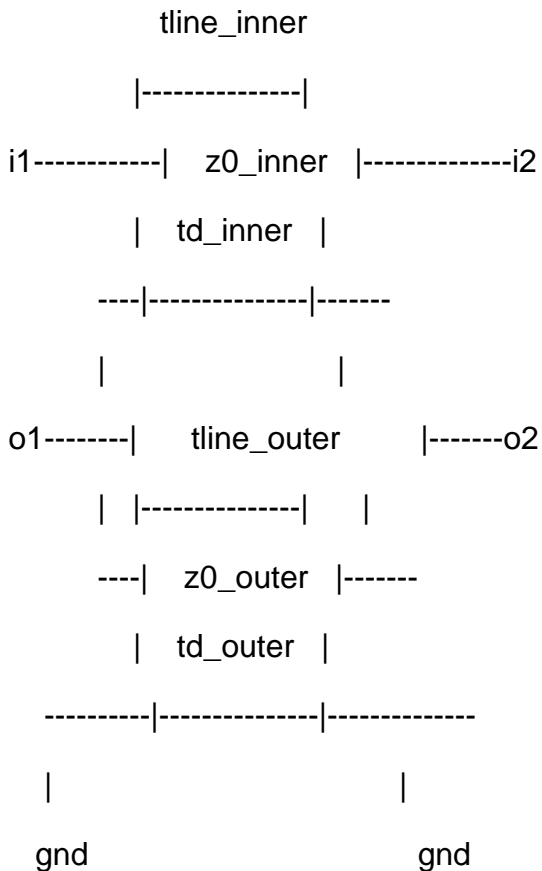
where t is time and t_d is the delay. Note, if the device is terminated by a matched impedance of z_0 (across t_2 and b_2), then it becomes an ideal delay. i.e $V_2(t) = V_1(t-t_d)$.

Spectre Circuit Simulator Reference

Component Statements Part III

To model both even and odd modes

Use two lines as shown below.



This model is suitable for a coax where `tline_inner` models the inner/outer conductor line (or the odd mode) while `tline_outer` models the outer/ground line (or the even mode). Note that this model is non-symmetric.

Lossy Case

In the frequency-domain the device is modeled by

$$V1(j\omega) - Z(j\omega) \cdot I1(j\omega) = S12(j\omega) \cdot [V2(j\omega) + Z(j\omega) \cdot I2(j\omega)]$$

and

$$V2(j\omega) - Z(j\omega) \cdot I2(j\omega) = S21(j\omega) \cdot [V1(j\omega) + Z(j\omega) \cdot I1(j\omega)]$$

Spectre Circuit Simulator Reference

Component Statements Part III

where $j = \sqrt{-1}$ and w is the angular frequency in radians/s. The loss coefficient is computed from

$$S_{21}(jw) = S_{12}(jw) = \exp(-\text{Gamma}(jw) \cdot \text{len})$$

where

$$\text{Gamma}(jw) = \sqrt{Z_c(jw) \cdot Y_d(jw)}$$

where Z_c represents the per-unit-length series impedance and Y_d represents the per-unit-length shunt admittance loss (as described below). The characteristic impedance (Z) is computed from

$$Z(jw) = \sqrt{Z_c(jw) / Y_d(jw)}$$

The time-domain behavior of the lossy transmission line is computed through a recursive convolution algorithm.

The dielectric loss (Y_d) is computed from

$$Y_d(jw) = G + j \cdot w / (z_0 \cdot c \cdot \text{vel})$$

where G is the per-unit-length shunt conductance and can be specified in three ways.

- 1) $G = g$ { when g is given }
- 2) $G = 2/z_0 \cdot \text{alphad}$ { when alphad is given }
- 3) $G = 2/z_0 \cdot f_d / (2 \cdot q_d \cdot c \cdot \text{vel})$ { when f_d and q_d are given }

where c is the speed of light.

The series impedance (Z_c) is computed from

$$Z_c(jw) = Z_i + j \cdot w \cdot z_0 / (c \cdot \text{vel})$$

where Z_i represents the internal loss. When skin effect is not present then

$$Z_i = d_{cr}$$

where d_{cr} is the DC series per-unit-length resistance.

Skin effect assuming finite thickness

In this case the internal impedance (Z_i) is computed from

Spectre Circuit Simulator Reference

Component Statements Part III

current, which results in lower loss. However, at the corner frequency, the skin depth equals the radius of the conductor. Decreasing the frequency below that point does not further reduce the loss.

The corner frequency (W_{corner}) can be specified in two ways.

1) When d_{cr} and corner are given, then

$$W_{\text{corner}} = 2 \cdot \pi \cdot \text{corner}.$$

2) When d_{cr} , r , and f_c are given, then

$$W_{\text{corner}} = 2 \cdot \pi \cdot f_c \cdot (d_{\text{cr}}/r)^2$$

In addition, there are two alternative ways to specify r .

1) $r = 2 \cdot z_0 \cdot \text{alphac}$ { when alphac is given }

2) $r = 2 \cdot z_0 \cdot f_c / (2 \cdot q_c \cdot c \cdot \text{vel})$ { when q_c is given }

where c is the speed of light are defined below.

Skin effect assuming infinite thickness

In this case there is no corner frequency (and no d_{cr}), and the internal loss (Z_i) is computed from

$$Z_i = R_i + j \cdot \omega \cdot L_i$$

where $R_i = r \cdot \sqrt{\omega / (2 \cdot \pi \cdot f_c)}$ and $L_i = r / \sqrt{\omega \cdot 2 \cdot \pi \cdot f_c}$.

Again, r can be specified directly, or using alphac or q_c as described above in the case of finite thickness.

Three ways to specify vel , td , and len

1) When vel and len are given

$$\text{td} = \text{len} / (\text{vel} \cdot c)$$

2) When td and vel are given

$$\text{len} = \text{td} \cdot \text{vel} \cdot c$$

Spectre Circuit Simulator Reference

Component Statements Part III

3) When f , $n1$ and vel are given

$$td = n1/f$$

$$len = (n1/f)*vel*c$$

The parameter len is the physical length, c is the speed of light and vel is the propagation velocity as a multiple of c (Recall that $velocity = c/\sqrt{\text{relative dielectric constant}}$). The parameter f is a reference frequency and $n1$ is the normalized electrical length in wavelengths at f .

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

α_c	I-12	f	M-2	len	I-6	r	I-11
α_c	M-8	f	I-3	m	I-7	td	I-2
α_d	M-12	f_c	M-6	$n1$	I-4	vel	M-3
α_d	I-16	f_c	I-10	q_c	M-9	vel	I-5
$corner$	M-4	f_d	M-10	q_c	I-13	$z0$	M-1
$corner$	I-8	f_d	I-14	q_d	I-17	$z0$	I-1
dcr	M-5	g	I-15	q_d	M-13		
dcr	I-9	g	M-11	r	M-7		

GaAs MESFET (tom2)

Description

TOM2 stands for Triquint Own Model version-2. It is an improved GaAs MESFET developed by David H. Smith. The charge model in TOM2 is similar to that of the Statz model and does not conserve charge. Therefore, this model should not be used to simulate circuits that requires charge conservation such as charge-pump circuits. TOM2 GaAs MESFET instances require that you use a model statement.

Spectre Circuit Simulator Reference

Component Statements Part III

This device is supported within altergroups.

There are some convergence problems with this model because of C_{gs} going to zero beyond pinchoff. The problems occur when the gate is driven from an inductive source, and there is no other capacitance at the gate. To prevent these problems, avoid setting C_{gd} to zero and add side wall capacitance to the gate-source and gate-drain junctions. A good estimate for these capacitors is $C = \pi \cdot \epsilon \cdot w/2$ where w is the gate width in microns and $\epsilon = 0.116$ fF/micron.

Sample Instance Statement

```
mt1 (2 1 0) tom2mos area=1 region=fwd
```

Sample Model Statement

```
model tom2mos tom2 vto=-0.55 alpha=3.9 beta=0.001 gamma=0.075 delta=100 ng=1 rd=550  
rs=550 rg=1 is=0.295e-14 n=1.2 cgs=1.4e-15 cgd=2e-16 cds=3e-16
```

Instance Definition

```
Name d g s ModelName parameter=value ...
```

Instance Parameters

- | | | |
|---|------------|---|
| 1 | area=1 | Junction area factor. |
| 2 | m=1 | Multiplicity factor. |
| 3 | region=fwd | Estimated operating region. Spectre outputs number (0-3) in a rawfile.
Possible values are off, triode, sat, or subth. |

Model Definition

```
model modelName tom2 parameter=value ...
```

Model Parameters

Device type parameters

- | | | |
|---|--------|---|
| 1 | type=n | Transistor type.
Possible values are n or p. |
|---|--------|---|

Spectre Circuit Simulator Reference

Component Statements Part III

Drain current parameters

2	<code>vto=-2.5 V</code>	Threshold voltage.
3	<code>alpha=2 1/V</code>	Knee-voltage parameter.
4	<code>beta=0.1 A/V²</code>	Transconductance parameter.
5	<code>gamma=0 1/V</code>	Threshold shifting parameter.
6	<code>delta=0.2 V</code>	Output feedback parameter.
7	<code>q=2</code>	Power-law parameter.

Subthreshold parameters

8	<code>ng=0</code>	Subthreshold slope gate parameter.
9	<code>nd=0 1/V</code>	Subthreshold slope drain pull parameter.

Parasitic resistance parameters

10	<code>rd=0 Ω</code>	Drain resistance (/area).
11	<code>rs=0 Ω</code>	Source resistance (/area).
12	<code>rg=0 Ω</code>	Gate resistance (/area).
13	<code>minr=0.1 Ω</code>	Minimum source/drain/gate resistance.

Junction diode model parameters

14	<code>is=1e-14 A</code>	Gate diode saturation current (*area).
15	<code>n=1</code>	Emission coefficient for the gate junction.
16	<code>imelt=`imaxA'</code>	Explosion current (*area).

Spectre Circuit Simulator Reference

Component Statements Part III

17 `dskip=yes` Use simple piece-wise linear model for diode currents below $0.1 \cdot i_{abstol}$.
Possible values are `no` or `yes`.

Junction capacitance model parameters

18 `capmod=2` Charge model selector.

19 `cgs=0 F` Gate-source zero-bias junction capacitance (*area).

20 `cgd=0 F` Gate-drain zero-bias junction capacitance (*area).

21 `cds=0 F` Drain-to-source capacitance.

22 `vbi=1 V` Gate diode built-in potential.

23 `vmax=0.95` Gate diode capacitance limiting voltage.

24 `vdelta=0.2 V` Capacitance transition voltage.

25 `tau=0 s` Conduction current delay time.

Temperature effects parameters

26 `tnom (C)` Parameters measurement temperature. Default set by options.

27 `xti=0` Temperature exponent for effect on `is`.

28 `eg=1.11 V` Energy band gap.

29 `vtotc=0 V/C` Temperature coefficient for `vto`.

30 `vbitc=0 V/C` Temperature coefficient for `vbi`.

31 `alphatce=0 1/C` Temperature coefficient for `alpha`.

32 `betatce=0 1/C` Temperature coefficient for `beta`.

33 `gammatc=0 1/C` Temperature coefficient for `gamma`.

34 `trsl=0 1/C` Temperature parameter for source resistance.

Spectre Circuit Simulator Reference

Component Statements Part III

- 35 `trd1=0 1/C` Temperature parameter for drain resistance.
- 36 `trg1=0 1/C` Temperature parameter for gate resistance.
- 37 `cgdtce=0 1/C` Drain junction capacitance temperature coefficient.
- 38 `cgstce=0 1/C` Source junction capacitance temperature coefficient.

Operating region warning control parameters

- 39 `imax=1 A` Maximum allowable current (*area).
- 40 `bvj= ∞ V` Junction reverse breakdown voltage.

Noise model parameters

- 41 `kf=0` Flicker (1/f) noise coefficient.
- 42 `af=1` Flicker (1/f) noise exponent.
- 43 `kfd=0` Flicker noise (1/f) coefficient for gate diodes.
- 44 `afg=1` Flicker noise (1/f) exponent for gate diodes.

The `imax` parameter aids convergence and prevents numerical overflow. The junction characteristics of the FET are accurately modeled for currents up to `imax`. For currents above `imax`, the junction is modeled as a linear resistor and a warning is printed. The `bv` parameter detects the junction breakdown only. The breakdown currents of the junctions are not modeled.

Operating-Point Parameters

- 1 `type=n` Transistor type.
Possible values are `n` or `p`.
- 2 `region=fwd` Estimated operating region. Spectre outputs number (0-3) in a rawfile.
Possible values are `off`, `triode`, `sat`, or `subth`.
- 3 `vgs (V)` Gate-source voltage.

Spectre Circuit Simulator Reference

Component Statements Part III

4	<code>vds</code> (V)	Drain-source voltage.
5	<code>id</code> (A)	Drain current.
6	<code>ig</code> (A)	Gate current.
7	<code>ids</code> (A)	Drain-to-source current.
8	<code>gm</code> (S)	Common-source transconductance.
9	<code>gds</code> (S)	Common-source output conductance.
10	<code>vth</code> (V)	Threshold voltage.
11	<code>cgs</code> (F)	Gate-source capacitance.
12	<code>cgd</code> (F)	Gate-drain capacitance.
13	<code>cds</code> (F)	Drain-source capacitance.
14	<code>pwr</code> (W)	Power at op point.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

<code>af</code>	M-42	<code>cgstce</code>	M-38	<code>m</code>	I-2	<code>trsl</code>	M-34
<code>afg</code>	M-44	<code>delta</code>	M-6	<code>minr</code>	M-13	<code>type</code>	OP-1
<code>alpha</code>	M-3	<code>dskip</code>	M-17	<code>n</code>	M-15	<code>type</code>	M-1
<code>alphatce</code>	M-31	<code>eg</code>	M-28	<code>nd</code>	M-9	<code>vbi</code>	M-22
<code>area</code>	I-1	<code>gamma</code>	M-5	<code>ng</code>	M-8	<code>vbitc</code>	M-30
<code>beta</code>	M-4	<code>gammatc</code>	M-33	<code>pwr</code>	OP-14	<code>vdelta</code>	M-24
<code>betatce</code>	M-32	<code>gds</code>	OP-9	<code>q</code>	M-7	<code>vds</code>	OP-4
<code>bvj</code>	M-40	<code>gm</code>	OP-8	<code>rd</code>	M-10	<code>vgs</code>	OP-3

Spectre Circuit Simulator Reference Component Statements Part III

capmod	M-18	id	OP-5	region	I-3	vmax	M-23
cds	OP-13	ids	OP-7	region	OP-2	vth	OP-10
cds	M-21	ig	OP-6	rg	M-12	vto	M-2
cgd	OP-12	imax	M-39	rs	M-11	vtotc	M-29
cgd	M-20	imelt	M-16	tau	M-25	x _{ti}	M-27
cgdtce	M-37	is	M-14	tnom	M-26		
cgs	OP-11	kf	M-41	trd1	M-35		
cgs	M-19	kfd	M-43	trg1	M-36		

GaAs MESFET (tom3)

Description

TOM3 stands for Triquint Own Model version-3. It is an improved GaAs MESFET developed by David H. Smith.

This device is supported within altergroups.

Sample Instance Statement

```
mt1 (2 1 0) tom3mos area=1 region=fwd
```

Sample Model Statement

```
model tom3mos tom3 vto=-0.55 alpha=3.9 beta=0.001 gamma=0.075 delta=100 rd=550  
rs=550 rg=1 is=1.0e-30 cds=3e-16
```

Instance Definition

```
Name d g s ModelName parameter=value ...
```

Instance Parameters

- | | | |
|---|---------------|--|
| 1 | area=1 | Junction area factor. |
| 2 | m=1 | Multiplicity factor. |
| 3 | region=triode | Estimated operating region.
Possible values are off, triode, sat, or subth. |

Spectre Circuit Simulator Reference

Component Statements Part III

Model Definition

model modelName tom3 parameter=value ...

Model Parameters

Device type parameters

1 type=n Transistor type.
Possible values are n or p.

Drain current parameters

2 vto=-2.5 V Threshold voltage.
3 alpha=2 1/V Knee-voltage parameter.
4 beta=0.1 A/V² Transconductance parameter.
5 gamma=0 1/V Threshold shifting parameter.
6 lambda=0.0 V Slope of drain characteristic.
7 q=2 Power-law parameter.
8 k=2.0 knee-function factor.

Subthreshold parameters

9 vst=1 V Subthreshold slope.
10 mst=0 1/V Subthreshold slope darin parameter.

Parasitic resistance parameters

11 rd=0 Ω Drain resistance (/area).
12 rs=0 Ω Source resistance (/area).
13 rg=0 Ω Gate resistance (/area).

Spectre Circuit Simulator Reference

Component Statements Part III

14 `minr=0.1 Ω` Minimum source/drain/gate resistance.

Junction diode model parameters

15 `is=0.0 A` Gate diode saturation current (*area).

16 `n=1` Emission coefficient for the gate junction.

17 `imelt=`imaxA'` Explosion current (*area).

18 `dskip=yes` Use simple piece-wise linear model for diode currents below $0.1 \cdot i_{abstol}$.
Possible values are `no` or `yes`.

19 `ilk=0.0 A` Gate leakage diode saturation current (*area).

20 `plk=1.0 V` Gate leakage diode potential.

Junction capacitance model parameters

21 `cds=0 F` Drain-to-source capacitance.

22 `tau=0 s` Conduction current delay time.

23 `qgqh=0.0` Charge parameter.

24 `qgsh=0.0` Charge parameter.

25 `qgdh=0.0` Charge parameter.

26 `qgio=1.0e-06` Charge parameter.

27 `qgql=0.0` Charge parameter.

28 `qgag=1.0` Charge parameter.

29 `qgad=1.0` Charge parameter.

30 `qgc1=0.0` Charge parameter.

31 `qggb=1.0` Charge parameter.

Spectre Circuit Simulator Reference

Component Statements Part III

32 `qggo=0.0` Charge parameter.

Temperature effects parameters

33 `tnom (C)` Parameters measurement temperature. Default set by options.

34 `xti=0` Temperature exponent for effect on *i_s*.

35 `eg=1.11 V` Energy band gap.

36 `vtotc=0 V/C` Temperature coefficient for *v_{to}*.

37 `alphatce=0 1/C` Temperature coefficient for *alpha*.

38 `betatce=0 1/C` Temperature coefficient for *beta*.

39 `gammatc=0 1/C` Temperature coefficient for *gamma*.

40 `trsl=0 1/C` Temperature parameter for source resistance.

41 `trdl=0 1/C` Temperature parameter for drain resistance.

42 `trgl=0 1/C` Temperature parameter for gate resistance.

43 `vsttc=0 1/C` Temperature coefficient for *V_{st}*.

44 `msttc=0 1/C` Temperature coefficient for *M_{st}*.

Operating region warning control parameters

45 `imax=1 A` Maximum allowable current (*area).

46 `bvj=∞ V` Junction reverse breakdown voltage.

Noise model parameters

47 `kf=0` Flicker (1/f) noise coefficient.

48 `af=1` Flicker (1/f) noise exponent.

Spectre Circuit Simulator Reference

Component Statements Part III

49 `kfd=0` Flicker noise (1/f) coefficient for gate diodes.

50 `afd=1` Flicker noise (1/f) exponent for gate diodes.

The `imax` parameter aids convergence and prevents numerical overflow. The junction characteristics of the FET are accurately modeled for currents up to `imax`. For currents above `imax`, the junction is modeled as a linear resistor and a warning is printed. The `bv` parameter detects the junction breakdown only. The breakdown currents of the junctions are not modeled.

Operating-Point Parameters

- | | | |
|----|----------------------------|--|
| 1 | <code>type=n</code> | Transistor type.
Possible values are <code>n</code> or <code>p</code> . |
| 2 | <code>region=triode</code> | Estimated operating region.
Possible values are <code>off</code> , <code>triode</code> , <code>sat</code> , or <code>subth</code> . |
| 3 | <code>vgs (V)</code> | Gate-source voltage. |
| 4 | <code>vds (V)</code> | Drain-source voltage. |
| 5 | <code>id (A)</code> | Drain current. |
| 6 | <code>ig (A)</code> | Gate current. |
| 7 | <code>ids (A)</code> | Drain-to-source current. |
| 8 | <code>gm (S)</code> | Common-source transconductance. |
| 9 | <code>gds (S)</code> | Common-source output conductance. |
| 10 | <code>vth (V)</code> | Threshold voltage. |
| 11 | <code>cgs (F)</code> | Gate-source capacitance. |
| 12 | <code>cgd (F)</code> | Gate-drain capacitance. |
| 13 | <code>cds (F)</code> | Drain-source capacitance. |
| 14 | <code>qg (Coul)</code> | Gate charge. |
| 15 | <code>qd (Coul)</code> | Drain charge. |

Spectre Circuit Simulator Reference

Component Statements Part III

- 16 `qs` (Coul) Source charge.
- 17 `pwr` (W) Power at op point.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

<code>af</code> M-48	<code>id</code> OP-5	<code>q</code> M-7	<code>rs</code> M-12
<code>afd</code> M-50	<code>ids</code> OP-7	<code>qd</code> OP-15	<code>tau</code> M-22
<code>alpha</code> M-3	<code>ig</code> OP-6	<code>qg</code> OP-14	<code>tnom</code> M-33
<code>alphatce</code> M-37	<code>ilk</code> M-19	<code>qgad</code> M-29	<code>trd1</code> M-41
<code>area</code> I-1	<code>imax</code> M-45	<code>qgag</code> M-28	<code>trgl</code> M-42
<code>beta</code> M-4	<code>imelt</code> M-17	<code>qgcl</code> M-30	<code>trsl</code> M-40
<code>betatce</code> M-38	<code>is</code> M-15	<code>qgdh</code> M-25	<code>type</code> OP-1
<code>bvj</code> M-46	<code>k</code> M-8	<code>qggb</code> M-31	<code>type</code> M-1
<code>cds</code> OP-13	<code>kf</code> M-47	<code>qggo</code> M-32	<code>vds</code> OP-4
<code>cds</code> M-21	<code>kfd</code> M-49	<code>qgio</code> M-26	<code>vgs</code> OP-3
<code>cgd</code> OP-12	<code>lambda</code> M-6	<code>qgqh</code> M-23	<code>vst</code> M-9
<code>cgs</code> OP-11	<code>m</code> I-2	<code>qgql</code> M-27	<code>vsttc</code> M-43
<code>dskip</code> M-18	<code>minr</code> M-14	<code>qgsh</code> M-24	<code>vth</code> OP-10
<code>eg</code> M-35	<code>mst</code> M-10	<code>qs</code> OP-16	<code>vto</code> M-2
<code>gamma</code> M-5	<code>msttc</code> M-44	<code>rd</code> M-11	<code>vtotc</code> M-36
<code>gammatc</code> M-39	<code>n</code> M-16	<code>region</code> OP-2	<code>xti</code> M-34
<code>gds</code> OP-9	<code>plk</code> M-20	<code>region</code> I-3	
<code>gm</code> OP-8	<code>pwr</code> OP-17	<code>rg</code> M-13	

Linear Two Winding Ideal Transformer (transformer)

Description

Winding 1 connects terminals t_1 and b_1 , and winding 2 connects t_2 and b_2 . The number of turns on windings 1 and 2 are given by n_1 and n_2 , respectively, and n_2 must not be zero. The absolute number of turns of each winding is not important, only the ratio of n_1 to n_2 . Current through winding 1 is computed.

This device is not supported within altergroup.

An ideal transformer is modeled, so it acts as a transformer at DC. Thus

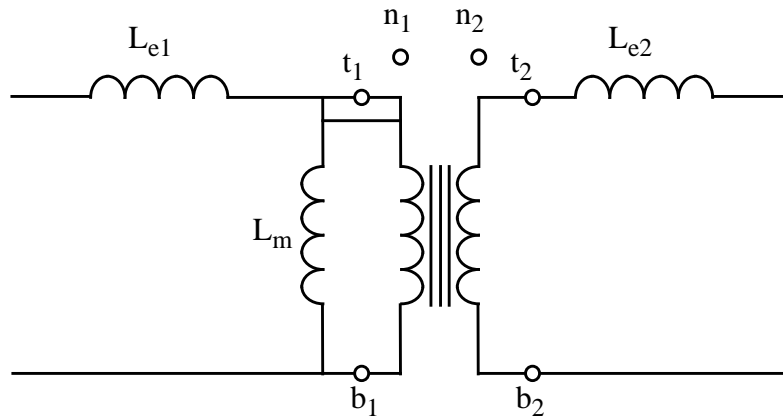
$$\frac{v_1}{v_2} = \frac{t_1}{t_2} = \frac{i_2}{i_1}$$

To model a physical transformer with L_1 and L_2 as the inductance of the windings and k as the coupling coefficient, add an inductor $L_m = k.L_1$ in parallel with winding 1 and inductors $L_{e1} = L_1.(1 - k)$ and $L_{e2} = L_2.(1 - k)$ in series with windings 1 and 2, respectively. The turns ratio can be computed with

$$\frac{n_1}{n_2} = \sqrt{\frac{L_1}{L_2}}$$

k can be calculated from the L_1 (the inductance of winding 1 with

$$k = \sqrt{1 - \frac{L_s}{L_1}}$$



Instance Definition

Name `t1 b1 t2 b2 transformer` parameter=value ...

Instance Parameters

- | | | |
|---|-------------------|-------------------------------|
| 1 | <code>n1=1</code> | Number of turns on winding 1. |
| 2 | <code>n2=1</code> | Number of turns on winding 2. |
| 3 | <code>m=1</code> | Multiplicity factor. |

VBIC Bipolar Transistor (`vbic`)

Description

The VBIC model was developed as a replacement for the SPICE G-P model. The model has four electrical terminals, two thermal terminals, and up to nine internal nodes, depending on the model parameters that the user specifies. VBIC 1.1.5 and VBIC 1.2 are implemented and controlled by model parameter version. Default is version=1.1.5. Detailed description of the

Spectre Circuit Simulator Reference

Component Statements Part III

model and equations are given in the Affirma Spectre Circuit Simulator Device Model Equations manual.

This device is supported within altergroups.

Sample Instance Statement

```
q1 (1 2 0 0 0) vbjt area=1
```

Sample Model Statement

```
model vbjt vbic type=npn is=2e-16 iben=4.5e-15 isp=1e-15 gamm=1.55e-11 ikf=0.0021  
ikr=0.0021 vef=15 ver=7 rbi=35 rbx=7 re=3 rs=15 cje=1.5e-14 tf=15e-12 selft=yes  
rth=1K
```

Instance Definition

```
Name c b e [s] [dt] [t1] ModelName parameter=value ...
```

`t1` node is the local temperature and the `dt` node is the rise above the local temperature caused by the thermal power dissipated by the device being modeled by VBIC. Consequently, the `t1` node can be connected to a thermal network that models heat flow through the substrate and/or between devices. It is not necessary to specify the substrate and the two thermal terminals. If left unspecified, the substrate and the `t1` thermal terminal are connected to ground. But if the self-heating flag is turned on and `dt` is not given, an internal node is created for self-heating. You must specify the substrate terminal if you specify `dt` and both substrate and `dt` must be given if `t1` needs to be specified.

It is strongly recommended to avoid using the `t1` node. It is not fully supported and using it will degrade convergence properties. This node is removed in the VBIC 1.2 version. All thermal effects can be modeled with just the `dt` node.

Model Definition

```
model modelName vbic parameter=value ...
```

Instance Parameters

- | | | |
|---|--------|-------------------------|
| 1 | area=1 | Transistor area factor. |
| 2 | m=1 | Multiplicity factor. |

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Component Statements Part III

3	<code>region=fwd</code>	Estimated operating region. Spectre outputs number (0-4) in a rawfile. Possible values are <code>off</code> , <code>fwd</code> , <code>rev</code> , <code>sat</code> , or <code>breakdown</code> .
4	<code>trise</code>	Temperature rise from ambient.
5	<code>dtemp</code>	Alias to <code>trise</code> .
6	<code>dtmp</code>	Alias to <code>trise</code> .

Model Parameters

Structural parameters

1	<code>type=npn</code>	Transistor type. Possible values are <code>npn</code> or <code>pnP</code> .
---	-----------------------	--

Saturation current parameters

2	<code>is=1e-16 A</code>	Transport saturation current (*area).
3	<code>ibei=1e-18 A</code>	Ideal B-E saturation current. (*area).
4	<code>iben=0 A</code>	Nonideal B-E saturation current (*area).
5	<code>ibci=1e-16 A</code>	Ideal B-C saturation current. (*area).
6	<code>ibcn=0 A</code>	Nonideal B-C saturation current (*area).
7	<code>isp=0 A</code>	Parasitic transport saturation current. (*area).
8	<code>ibeip=0 A</code>	Ideal parasitic B-E saturation current (*area).
9	<code>ibenp=0 A</code>	Nonideal parasitic B-E saturation current (*area).
10	<code>ibcip=0 A</code>	Ideal parasitic B-C saturation current (*area).
11	<code>ibcnp=0 A</code>	Nonideal parasitic B-C saturation current (*area).
12	<code>vo=0 V</code>	Epi drift saturation voltage.

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Component Statements Part III

13	gamm=0 V	Epi doping parameter.
14	hrcf=1	High current RC factor.
15	wbe=1	Portion of Ibei from Vbei.
16	wsp=1	Portion of Iccp from Vbep.

Emission coefficient parameters

17	nf=1	Forward emission coefficient.
18	nr=1	Reverse emission coefficient.
19	nei=1	Ideal B-E emission coefficient.
20	nen=2	Nonideal B-E emission coefficient.
21	nci=1	Ideal B-C emission coefficient.
22	ncn=2	Nonideal B-C emission coefficient.
23	nfp=1	Parasitic forward emission coefficient.
24	ncip=1	Ideal parasitic B-C emission coefficient.
25	ncnp=2	Nonideal parasitic B-C emission coefficient.

Current gain parameters

26	ikf= ∞ A	Forward knee current (*area).
27	ikr= ∞ A	Reverse knee current (*area).
28	ikp= ∞ A	Parasitic knee current (*area).

Early voltage parameters

29	vef= ∞ V	Forward Early voltage.
----	-----------------	------------------------

Spectre Circuit Simulator Reference

Component Statements Part III

30 $ver=\infty$ V Reverse Early voltage.

Breakdown voltage parameters

31 $avc1=0$ B-C weak avalanche parameter.

32 $avc2=0$ B-C weak avalanche parameter.

Parasitic resistance parameters

33 $rbi=0$ Ω Intrinsic base resistance (/area).

34 $rbx=0$ Ω Extrinsic base resistance (/area).

35 $re=0$ Ω Emitter resistance (/area).

36 $rs=0$ Ω Substrate resistance (/area).

37 $rbp=0$ Ω Parasitic base resistance (/area).

38 $rcx=0$ Ω Extrinsic collector resistance (/area).

39 $rci=0$ Ω Intrinsic collector resistance (/area).

Junction capacitance parameters

40 $cje=0$ F B-E zero-bias capacitance (*area).

41 $pe=0.75$ V B-E built-in potential.

42 $me=0.33$ B-E grading coefficient.

43 $aje=-0.5$ B-E capacitance smoothing factor.

44 $fc=0.9$ Forward-bias depletion capacitance limit.

45 $cbeo=0$ F Extrinsic B-E overlap capacitance (*area).

46 $cjc=0$ F B-C zero-bias capacitance (*area).

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Component Statements Part III

47	<code>cjep=0 F</code>	B-C extrinsic zero-bias capacitance (*area).
48	<code>pc=0.75 V</code>	B-C built-in potential.
49	<code>mc=0.33</code>	B-C grading coefficient.
50	<code>ajc=-0.5</code>	B-C capacitance smoothing factor.
51	<code>cbco=0 F</code>	Extrinsic B-C overlap capacitance (*area).
52	<code>qco=0 Coul</code>	Epi charge parameter.
53	<code>cjcp=0 F</code>	S-C zero-bias capacitance (*area).
54	<code>ps=0.75 V</code>	S-C built-in potential.
55	<code>ms=0.33</code>	S-C grading coefficient.
56	<code>ajs=-0.5</code>	S-C capacitance smoothing factor.

Transit time and excess phase parameters

57	<code>tf=0 s</code>	Forward transit time.
58	<code>tr=0 s</code>	Reverse transit time.
59	<code>td=0 s</code>	Forward excess-phase delay time.
60	<code>qtf=0</code>	Variation of t_f with base width modulation.
61	<code>xtf=0</code>	Coefficient of t_f with bias dependence.
62	<code>vtf=0</code>	Coefficient of t_f dependence on V_{bc} .
63	<code>itf=0</code>	Coefficient of t_f dependence on I_c .

Temperature effects parameters

64	<code>selft=0</code>	Flag denoting self-heating. Possible values are <code>no</code> or <code>yes</code> .
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Spectre Circuit Simulator Reference

Component Statements Part III

65	<code>tnom</code> (C)	Parameters measurement temperature. Default set by options.
66	<code>trise=0</code> C	Temperature rise from ambient.
67	<code>rth=0</code> Ω	Thermal resistance, must be given for self-heating.
68	<code>cth=0</code> F	Thermal capacitance.
69	<code>xis=3</code> V	Temperature exponent of <code>Is</code> .
70	<code>xii=3</code> V	Temperature exponent of <code>Ibei</code> , <code>Ibci</code> , <code>Ibeip</code> , and <code>Ibcip</code> .
71	<code>xin=3</code> V	Temperature exponent of <code>Iben</code> , <code>Ibcn</code> , <code>Ibenp</code> , and <code>Ibcnp</code> .
72	<code>tnf=0</code> V	Temperature coefficient of <code>Nf</code> .
73	<code>tavc=0</code> V	Temperature coefficient of <code>Avc2</code> .
74	<code>ea=1.12</code> V	Activation energy for <code>is</code> .
75	<code>eaie=1.12</code> V	Activation energy for <code>Ibei</code> .
76	<code>eaic=1.12</code> V	Activation energy for <code>Ibci/Ibeip</code> .
77	<code>eais=1.12</code> V	Activation energy for <code>Ibcip</code> .
78	<code>eane=1.12</code> V	Activation energy for <code>Iben</code> .
79	<code>eanc=1.12</code> V	Activation energy for <code>Ibcn/Ibenp</code> .
80	<code>eans=1.12</code> V	Activation energy for <code>Ibcnp</code> .
81	<code>xre=0</code>	Temperature exponent of <code>re</code> .
82	<code>xrb=0</code>	Temperature exponent of <code>rb</code> .
83	<code>xrc=0</code>	Temperature exponent of <code>rc</code> .
84	<code>xrs=0</code>	Temperature exponent of <code>rs</code> .
85	<code>xvo=0</code>	Temperature exponent of <code>vo</code> .
86	<code>dtmax=226.85</code> C	Maximum expected device temperature. (500 K).

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Component Statements Part III

Noise model parameters

- 87 `kfn=0` B-E flicker (1/f) noise coefficient.
- 88 `afn=1` B-E flicker (1/f) noise exponent.
- 89 `bfm=1` B-E flicker (1/f) noise dependence.

Junction diode model control parameters

- 90 `dskip=yes` Skip junction calculations if they are reverse-saturated.
Possible values are `no` or `yes`.
- 91 `imelt=10 A` Explosion current (*area).

Operating region warning control parameters

- 92 `bvbe= ∞ V` B-E breakdown voltage.
- 93 `bvbc= ∞ V` B-C breakdown voltage.
- 94 `bvce= ∞ V` C-E breakdown voltage.
- 95 `bvsub= ∞ V` Substrate junction breakdown voltage.
- 96 `vbefwd=0.2 V` B-E forward voltage.
- 97 `vbcfwd=0.2 V` B-C forward voltage.
- 98 `vsubfwd=0.2 V` Substrate junction forward voltage.
- 99 `imax=1 A` Maximum allowable base current (*area).
- 100 `imax1=imax A` Maximum allowable collector current (*area).
- 101 `alarm=none` Forbidden operating region.
Possible values are `none`, `off`, `fwd`, `rev`, or `sat`.

Spectre Circuit Simulator Reference

Component Statements Part III

DC-mismatch model parameters

102 `mvt0=0.0 V` Threshold mismatch intercept.

New model parameter for Vbic 1.2

103 `vrt=0 V` B-C reach-through limiting voltage.

104 `art=0.1 V` B-C reach-through limiting smoothing factor .

105 `ccso=0 F` Fixed collector-substrate capacitance.

106 `qbm=0` Parameter to select SGP qb formulation.

107 `nkf=0.5` High current beta rolloff parameter.

108 `xikf=0` Parameter of temperature dependence to `ikf`.

109 `xrcx=0` Parameter of temperature dependence to `rcx`.

110 `xrbx=0` Parameter of temperature dependence to `rbx`.

111 `xrbp=0` Parameter of temperature dependence to `rbp`.

112 `isrr=1` Parameter to separate `is` for forward and reversed parts.

113 `xisr=0` Temperature exponent coefficient of `isrr`.

114 `dear=0 V` Activation energy for `isrr`.

115 `eap=1.12 V` Activation energy for `isp`.

116 `vbbe=0 V` B-E breakdown voltage.

117 `nbbe=1.0` B-E breakdown emission coefficient.

118 `ibbe=1.0e-6 A` B-E breakdown current.

119 `tvbbe1=0` First temperature coefficient of `vbbe`.

120 `tvbbe2=0` Second temperature coefficient of `vbbe`.

Spectre Circuit Simulator Reference

Component Statements Part III

121	<code>tnbbe=0</code>	Temperature coefficient for <code>nbbe</code> .
122	<code>vers=1.15</code>	Version control parameter.
123	<code>vrev=0</code>	Revision control parameter.
124	<code>dtemp (C)</code>	Alias to <code>trise</code> .
125	<code>dtmp (C)</code>	Alias to <code>trise</code> .
126	<code>version</code>	Alias to <code>vers</code> .
127	<code>rev</code>	Alias to <code>vrev</code> .
128	<code>xrbi</code>	Alias to <code>xrb</code> .
129	<code>xrci</code>	Alias to <code>xrc</code> .

Imax and Imelt

The `imax` parameter aids convergence and prevents numerical overflow. The junction characteristics of the device are accurately modeled for current up to `imax`. If `imax` is exceeded during iterations, the linear model is substituted until the current drops below `imax` or until convergence is achieved. If convergence is achieved with the current exceeding `imax`, the results are inaccurate, and Spectre prints a warning.

A separate model parameter, `imelt`, is used as a limit warning for the junction current. This parameter can be set to the maximum current rating of the device. When any component of the junction current exceeds `imelt`, note that base and collector currents are composed of many exponential terms, Spectre issues a warning and the results become inaccurate. The junction current is linearized above the value of `imelt` to prevent arithmetic exception, with the exponential term replaced by a linear equation at `imelt`.

Operating-Point Parameters

1	<code>type=npn</code>	Transistor type. Possible values are <code>npn</code> or <code>pnp</code> .
2	<code>region=fwd</code>	Estimated operating region. Spectre outputs number (0-4) in a <code>rawfile</code> . Possible values are <code>off</code> , <code>fwd</code> , <code>rev</code> , <code>sat</code> , or <code>breakdown</code> .

Spectre Circuit Simulator Reference

Component Statements Part III

3	vbe (V)	Base-emitter voltage.
4	vbc (V)	Base-collector voltage.
5	vce (V)	Collector-emitter voltage.
6	vcs (V)	Collector-substrate voltage.
7	temp (C)	Device temperature.
8	ith (A)	Thermal source.
9	ic (A)	Intrinsic DC collector current. ($I_{cc} - I_{bc} + I_{gc}$).
10	ib (A)	Intrinsic DC base current. ($I_{be} + I_{bc} - I_{gc}$).
11	icc (A)	C-E current.
12	ibe (A)	Intrinsic B-E junction current.
13	ibc (A)	Intrinsic B-C junction current.
14	ibex (A)	BX-E junction current.
15	igc (A)	Breakdown current.
16	iccp (A)	Parasitic transistor C-E current.
17	ibep (A)	Parasitic transistor B-E current.
18	ibcp (A)	Parasitic transistor B-C current.
19	betadc (A/A)	Ratio of external collector current to external base current. (I_{c_ext}/I_{b_ext}).
20	gm (S)	Intrinsic small-signal transconductance. ($g_m = dI_{cc_dVbe} + dI_{cc_dVbc}$).
21	gpi (S)	Intrinsic small-signal input conductance. ($g_{pi} = dI_{be_dVbe}$).
22	go (S)	Intrinsic small-signal output conductance. ($g_o = -dI_{cc_dVbc}$).

Spectre Circuit Simulator Reference

Component Statements Part III

23	<code>gmu</code> (S)	Intrinsic small-signal Collector-Base conductance. ($gmu = dlbc_dVbc$).
24	<code>cpi</code> (F)	Intrinsic small-signal B-E capacitance. Same as <code>cje</code> .
25	<code>cmu</code> (F)	Intrinsic small signal B-C capacitance. Same as <code>cjc</code> .
26	<code>betaac</code> (A/A)	Small-signal common-emitter current gain. (gm/gpi).
27	<code>ft</code> (Hz)	Unity small-signal current-gain frequency.
28	<code>dic_dvbe</code> (S)	Intrinsic $dlc/dVbe$.
29	<code>dic_dvbc</code> (S)	Intrinsic dlc_dVbc .
30	<code>dib_dvbe</code> (S)	Intrinsic dlb_dVbe .
31	<code>dib_dvbc</code> (S)	Intrinsic dlb_dVbc .
32	<code>rbi</code> (Ω)	Intrinsic base resistance.
33	<code>rci</code> (Ω)	Intrinsic collector resistance.
34	<code>rbp</code> (Ω)	Parasitic transistor base resistance.
35	<code>cje</code> (F)	Intrinsic B-E capacitance.
36	<code>cjc</code> (F)	Intrinsic B-C capacitance.
37	<code>cbex</code> (F)	BX-E junction capacitance.
38	<code>cbcx</code> (F)	B-CX junction capacitance.
39	<code>cbep</code> (F)	Parasitic B-E junction capacitance.
40	<code>cbcp</code> (F)	Parasitic B-C junction capacitance.
41	<code>pwr</code> (W)	Power dissipation.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point

Spectre Circuit Simulator Reference Component Statements Part III

parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of M-35 means the 35th model parameter.

afn	M-88	eaic	M-76	ith	OP-8	tnbbe	M-121
ajc	M-50	eaie	M-75	kfn	M-87	tnf	M-72
aje	M-43	eaiss	M-77	m	I-2	tnom	M-65
ajs	M-56	eanc	M-79	mc	M-49	tr	M-58
alarm	M-101	eane	M-78	me	M-42	trise	M-66
area	I-1	eans	M-80	ms	M-55	trise	I-4
art	M-104	eap	M-115	mvt0	M-102	tvbbe1	M-119
avc1	M-31	fc	M-44	nbbe	M-117	tvbbe2	M-120
avc2	M-32	ft	OP-27	nci	M-21	type	OP-1
betaac	OP-26	gamm	M-13	ncip	M-24	type	M-1
betadc	OP-19	gm	OP-20	ncn	M-22	vbbe	M-116
bfm	M-89	gmu	OP-23	ncnp	M-25	vbc	OP-4
bvbc	M-93	go	OP-22	nei	M-19	vbcfwd	M-97
bvbe	M-92	gpi	OP-21	nen	M-20	vbe	OP-3
bvce	M-94	hrcf	M-14	nf	M-17	vbefwd	M-96
bvsub	M-95	ib	OP-10	nfp	M-23	vce	OP-5
cbco	M-51	ibbe	M-118	nkf	M-107	vcs	OP-6
cbcp	OP-40	ibc	OP-13	nr	M-18	vef	M-29
cbcx	OP-38	ibci	M-5	pc	M-48	ver	M-30
cbeo	M-45	ibcip	M-10	pe	M-41	vers	M-122
cbep	OP-39	ibcn	M-6	ps	M-54	version	M-126
cbex	OP-37	ibcnp	M-11	pwr	OP-41	vo	M-12
ccso	M-105	ibcp	OP-18	qbm	M-106	vrev	M-123
cjc	OP-36	ibe	OP-12	qco	M-52	vrt	M-103
cjc	M-46	ibei	M-3	qtf	M-60	vsubfwd	M-98
cjcp	M-53	ibeip	M-8	rbi	M-33	vtf	M-62
cje	OP-35	iben	M-4	rbi	OP-32	wbe	M-15
cje	M-40	ibenp	M-9	rbp	M-37	wsp	M-16

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cjep	M-47	ibep	OP-17	rbp	OP-34	xii	M-70
cmu	OP-25	ibex	OP-14	rbx	M-34	xikf	M-108
cpi	OP-24	ic	OP-9	rci	M-39	xin	M-71
cth	M-68	icc	OP-11	rci	OP-33	xis	M-69
dear	M-114	iccp	OP-16	rcx	M-38	xisr	M-113
dib_dvbc	OP-31	igc	OP-15	re	M-35	xrb	M-82
dib_dvbe	OP-30	ikf	M-26	region	I-3	xrbi	M-128
dic_dvbc	OP-29	ikp	M-28	region	OP-2	xrbp	M-111
dic_dvbe	OP-28	ikr	M-27	rev	M-127	xrbx	M-110
dskip	M-90	imax	M-99	rs	M-36	xrc	M-83
dtemp	M-124	imax1	M-100	rth	M-67	xrci	M-129
dtemp	I-5	imelt	M-91	selft	M-64	xrcx	M-109
dtmax	M-86	is	M-2	tavc	M-73	xre	M-81
dtmp	M-125	isp	M-7	td	M-59	xrs	M-84
dtmp	I-6	isrr	M-112	temp	OP-7	xtf	M-61
ea	M-74	itf	M-63	tf	M-57	xvo	M-85

Linear Voltage Controlled Current Source (vccs)

Description

Positive current exits the source node and enters the sink node.

This device can also model ideal digital gates, voltage controlled resistors and voltage controlled capacitors.

This device is supported within altergroups.

Sample Instance Statement

```
v1 (1 0 2 3) gm=-1 m=2
```

Instance Definition

```
Name sink src ps ns ... vccs parameter=value ...
```


Spectre Circuit Simulator Reference

Component Statements Part III

Instance Parameters

- | | | |
|---|------------------------|---|
| 1 | <code>m=1</code> | Multiplicity factor. |
| 2 | <code>type=vccs</code> | Type of the source.
Possible values are <code>vccs</code> , <code>vcr</code> , <code>vccap</code> , <code>and</code> , <code>nand</code> , <code>or</code> , or <code>nor</code> . |
| 3 | <code>delta=0</code> | Smoothing parameter. This may lead to circuit convergency. The smaller the delta is, the sharper the corner is. |

Linear source parameters

- | | | |
|---|----------------------|---|
| 4 | <code>gm=0 S</code> | Transconductance. |
| 5 | <code>min (A)</code> | Minimum output current. |
| 6 | <code>max (A)</code> | Maximum output current. |
| 7 | <code>abs=off</code> | Absolute output current.
Possible values are <code>off</code> or <code>on</code> . |

PWL source parameters

- | | | |
|----|------------------------|---|
| 8 | <code>file</code> | Name of file containing voltage/current pairs that define the PWL transfer function. |
| 9 | <code>pwl=[...]</code> | Vector of voltage/current pairs that defines the PWL transfer function. The format of the vector is <code>[in1 out1 in2 out2 ...]</code> . |
| 10 | <code>scale=1</code> | Scale factor for the PWL output current. |
| 11 | <code>stretch=1</code> | Scale factor for the PWL controlling voltage. |

Temperature effects parameters

- | | | |
|----|-----------------------------------|------------------------------------|
| 12 | <code>tc1=0 1/C</code> | Linear temperature coefficient. |
| 13 | <code>tc2=0 C⁻²</code> | Quadratic temperature coefficient. |

Spectre Circuit Simulator Reference

Component Statements Part III

Operating-Point Parameters

- | | | |
|---|----------------------|--------------------|
| 1 | <code>i</code> (A) | Output current. |
| 2 | <code>v</code> (V) | Output voltage. |
| 3 | <code>pwr</code> (W) | Power dissipation. |

Parameter Index

In the following index, `I` refers to instance parameters, `M` refers to the model parameters section, `O` refers to the output parameters section, and `OP` refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of `M-35` means the 35th model parameter.

<code>abs</code>	I-7	<code>i</code>	OP-1	<code>pwl</code>	I-9	<code>tc1</code>	I-12
<code>delta</code>	I-3	<code>m</code>	I-1	<code>pwr</code>	OP-3	<code>tc2</code>	I-13
<code>file</code>	I-8	<code>max</code>	I-6	<code>scale</code>	I-10	<code>type</code>	I-2
<code>gm</code>	I-4	<code>min</code>	I-5	<code>stretch</code>	I-11	<code>v</code>	OP-2

Linear Voltage Controlled Voltage Source (`vcvs`)

Description

Current through the voltage source is calculated and is defined to be positive if it flows from the positive terminal, through the source, to the negative terminal.

This device can also model ideal digital gates.

This device is supported within altergroups.

Sample Instance Statement

```
e1 (out1 0 pos neg) vcvs gain=10
```

Spectre Circuit Simulator Reference

Component Statements Part III

Instance Definition

Name p n ps ns ... vcvs parameter=value ...

Instance Parameters

- | | | |
|---|-----------|---|
| 1 | m=1 | Multiplicity factor. |
| 2 | type=vcvs | Type of the source.
Possible values are <code>vcvs</code> , <code>and</code> , <code>nand</code> , <code>or</code> , or <code>nor</code> . |
| 3 | delta=0 | Smoothing parameter. This may lead to circuit convergency. The smaller the delta is, the sharper the corner is. |

Linear source parameters

- | | | |
|---|------------|---|
| 4 | gain=0 V/V | Voltage gain. |
| 5 | min (V) | Minimum output voltage. |
| 6 | max (V) | Maximum output voltage. |
| 7 | abs=off | Absolute output voltage.
Possible values are <code>off</code> or <code>on</code> . |

PWL source parameters

- | | | |
|----|-----------|---|
| 8 | file | Name of file containing voltage/voltage pairs that define the PWL transfer function. |
| 9 | pwl=[...] | Vector of voltage/voltage pairs that defines the PWL transfer function. The format of the vector is [in1 out1 in2 out2 ...]. |
| 10 | scale=1 | Scale factor for the PWL output voltage. |
| 11 | stretch=1 | Scale factor for the PWL controlling voltage. |

Temperature effects parameters

- | | | |
|----|-----------|---------------------------------|
| 12 | tc1=0 1/C | Linear temperature coefficient. |
|----|-----------|---------------------------------|

Spectre Circuit Simulator Reference

Component Statements Part III

13 $tc2=0$ C^{-2} Quadratic temperature coefficient.

Operating-Point Parameters

1 i (A) Output current.
2 v (V) Output voltage.
3 pwr (W) Power dissipation.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

abs	I-7	i	OP-1	pwl	I-9	$tc1$	I-12
$delta$	I-3	m	I-1	pwr	OP-3	$tc2$	I-13
$file$	I-8	max	I-6	$scale$	I-10	$type$	I-2
$gain$	I-4	min	I-5	$stretch$	I-11	v	OP-2

Independent Voltage Source (vsource)

Description

Current through the source is computed and is defined to be positive if it flows from the positive node, through the source, to the negative node.

This device is supported within altergroups.

The value of the DC voltage as a function of the temperature is given by:

$$V(T) = V(tnom) * [1 + tc1 * (T - tnom) + tc2 * (T - tnom)^2].$$

Spectre Circuit Simulator Reference

Component Statements Part III

Sample Instance Statement

```
vpulse1 (1 0) vsource type=pulse val0=0 val1=5 period=100n rise=10n fall=10n
width=40n
vpwl1 (1 0) vsource type=pwl wave=[1n 0 1.1n 2 1.5n 0.5 2n 3 5n 5] pwlperiod=5n
```

Instance Definition

Name p n vsource parameter=value ...

Instance Parameters

1 dc=0 V DC value.

General waveform parameters

2 type=dc Waveform type.
Possible values are dc, pulse, pwl, sine, or exp.

3 fundname Name of the fundamental frequency. Must be specified if the source is active during a pdisto analysis or it is the active clock during an envlp analysis.

4 delay=0 s Waveform delay time.

Pulse waveform parameters

5 val0=0 V Zero value used in pulse and exponential waveforms.

6 val1=1 V One value used in pulse and exponential waveforms.

7 period= ∞ s Period of waveform.

8 rise (s) Rise time for pulse waveform (time for transition from val0 to val1).

9 fall (s) Fall time for pulse waveform (time for transition from val1 to val0).

10 width= ∞ s Pulse width (duration of val1).

Spectre Circuit Simulator Reference

Component Statements Part III

PWL waveform parameters

11	<code>file</code>	Name of file containing waveform.
12	<code>wave=[...]</code>	Vector of time/value pairs that defines waveform.
13	<code>offset=0 V</code>	DC offset for the PWL waveform.
14	<code>scale=1</code>	Scale factor for the PWL waveform.
15	<code>stretch=1</code>	Scale factor for time given for the PWL waveform.
16	<code>allbrkpts</code>	All the points in the PWL waveform are breakpoints if set to yes. Default is yes if the number of points is less than 20. Possible values are <code>no</code> or <code>yes</code> .
17	<code>pwlperiod (s)</code>	Period of the periodic PWL waveform.
18	<code>twidth=pwlperiod/1000 s</code>	Transition width used when making PWL waveforms periodic.

Sinusoidal waveform parameters

19	<code>sinedc=dc V</code>	DC level for sinusoidal waveforms.
20	<code>ampl=1 V</code>	Peak amplitude of sinusoidal waveform.
21	<code>freq=0 Hz</code>	Frequency of sinusoidal waveform.
22	<code>sinephase=0 °</code>	Phase of sinusoid when <code>t=delay</code> .
23	<code>ampl2=1 V</code>	Peak amplitude of second sinusoidal waveform.
24	<code>freq2=0 Hz</code>	Frequency of second sinusoidal waveform.
25	<code>sinephase2=0 °</code>	Phase of second sinusoid when <code>t=delay</code> .
26	<code>fundname2</code>	Name of the fundamental frequency associated with <code>freq2</code> . Must be specified if <code>freq2</code> is used in a <code>pdisto</code> analysis.
27	<code>fmodindex=0</code>	FM index of modulation for sinusoidal waveform.

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Component Statements Part III

- 28 `fmodfreq=0 Hz` FM modulation frequency for sinusoidal waveform.
- 29 `ammodindex=0` AM index of modulation for sinusoidal waveform.
- 30 `ammodfreq=0 Hz` AM modulation frequency for sinusoidal waveform.
- 31 `ammodphase=0 °` AM phase of modulation for sinusoidal waveform.
- 32 `damp=0 1/s` Damping factor for sinusoidal waveform.

Exponential waveform parameters

- 33 `td1=0 s` Rise start time for exponential wave.
- 34 `tau1 (s)` Rise time constant for exponential wave.
- 35 `td2 (s)` Fall start time for exponential wave.
- 36 `tau2 (s)` Fall time constant for exponential wave.

Noise Parameters

- 37 `noisefile` Name of file containing excess spot noise data in the form of frequency-noise pairs.
- 38 `noisevec=[...] V2/Hz` Excess spot noise as a function of frequency in the form of frequency-noise pairs.

Small signal parameters

- 39 `mag=0 V` Small signal voltage.
- 40 `phase=0 °` Small signal phase.
- 41 `xfmag=1 V/V` Transfer function analysis magnitude.
- 42 `pacmag=0 V` Periodic AC analysis magnitude.
- 43 `pacphase=0 °` Periodic AC analysis phase.

Spectre Circuit Simulator Reference

Component Statements Part III

Multiplication factor parameters

44 `m=1` Multiplicity factor.

Temperature effects parameters

45 `tc1=0 1/C` First order temperature coefficient.

46 `tc2=0 C-2` Second order temperature coefficient.

47 `tnom=27 C` Parameter measurement temperature. Default set by options.

If you do not specify the DC value, it is assumed to be the `time=0` value of the waveform.

Sinusoidal waveform in the time interval $0 < t < \text{delay}$ has constant value equal to that at $t = \text{delay}$.

In DC analyses, the only active parameters are `dc`, `m`, and the temperature coefficient parameters. In AC analyses, the only active parameters are `m`, `mag` and `phase`. In transient analyses, all parameters are active except the small signal parameters and the noise parameters. The `type` parameter selects which type of waveform is generated. You may specify parameters for more than one waveform type, and use the `alter` statement to change the waveform type between analyses.

A vector of time-value pairs describes the piecewise linear waveform. As an alternative, you can read the waveform from a file. In this case, you give time-value pairs one pair per line with a space or tab between the time and the value.

If you set `allbrkpts` to `yes`, you force the simulator to place time points at each point specified in a PWL waveform during a transient analysis. This can be very expensive for waveforms with many points. If you set `allbrkpts` to `no`, Spectre inspects the waveform, looking for abrupt changes, and forces time points only at those changes.

The PWL waveform is periodic if you specify `pwlperiod`. If the value of the waveform specified is not exactly the same at both its beginning and its end, then you must provide a nonzero value `twidth`. Before repeating, the waveform changes linearly in an interval of `twidth` from its value at $(\text{period} - \text{twidth})$ to its value at the beginning of the waveform. Thus `twidth` must always be less than `period`.

You can give the excess noise of the source as a file or specify it with a vector of frequency-noise pairs. For a file, give the frequency-noise pairs one pair per line with a space or tab between the frequency and noise values.

Spectre Circuit Simulator Reference

Component Statements Part III

Operating-Point Parameters

- 1 `v` (V) Voltage across the source.
- 2 `i` (A) Current through the source.
- 3 `pwr` (W) Power dissipation.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

<code>allbrkpts</code> I-16	<code>freq</code> I-21	<code>phase</code> I-40	<code>td1</code> I-33
<code>ammodfreq</code> I-30	<code>freq2</code> I-24	<code>pwlperiod</code> I-17	<code>td2</code> I-35
<code>ammodindex</code> I-29	<code>fundname</code> I-3	<code>pwr</code> OP-3	<code>tnom</code> I-47
<code>ammodphase</code> I-31	<code>fundname2</code> I-26	<code>rise</code> I-8	<code>twidht</code> I-18
<code>amp1</code> I-20	<code>i</code> OP-2	<code>scale</code> I-14	<code>type</code> I-2
<code>amp12</code> I-23	<code>m</code> I-44	<code>sinedc</code> I-19	<code>v</code> OP-1
<code>damp</code> I-32	<code>mag</code> I-39	<code>sinphase</code> I-22	<code>val0</code> I-5
<code>dc</code> I-1	<code>noisefile</code> I-37	<code>sinphase2</code> I-25	<code>val1</code> I-6
<code>delay</code> I-4	<code>noisevec</code> I-38	<code>stretch</code> I-15	<code>wave</code> I-12
<code>fall</code> I-9	<code>offset</code> I-13	<code>tau1</code> I-34	<code>width</code> I-10
<code>file</code> I-11	<code>pacmag</code> I-42	<code>tau2</code> I-36	<code>xfmag</code> I-41
<code>fmmodfreq</code> I-28	<code>pacphase</code> I-43	<code>tc1</code> I-45	
<code>fmmodindex</code> I-27	<code>period</code> I-7	<code>tc2</code> I-46	

Winding for Magnetic Core (winding)

Description

This winding is used in conjunction with magnetic cores to model coils and transformers with hysteresis. Each winding must be associated with a single core, though a core may have any number of windings.

Winding connects terminals `t1` and `b1`. Current through the winding is computed.

This device is not supported within `altergroup`.

Sample Instance Statement

```
c1 (1 0) core_mod area=1.2 len=8.1 id=0.45 id=0.55 gap=0.25
y1 (2 0) winding turn=5 core=c1 resis=1m
```

Instance Definition

```
Name t b winding parameter=value ...
```

Instance Parameters

- | | | |
|---|--|---|
| 1 | <code>turn=1</code> | Number of turns on winding. |
| 2 | <code>resis (Ω)</code> | Resistance of the winding. |
| 3 | <code>m=1</code> | Multiplicity factor. |
| 4 | <code>core</code> | Name of core around which winding is wrapped. |

Initial Conditions

- | | | |
|---|-----------------------|-----------------------------------|
| 5 | <code>ic=0.0 A</code> | Initial condition on the winding. |
|---|-----------------------|-----------------------------------|

z-Domain Linear Current Controlled Current Source (zcccs)

Description

The output is defined with a transfer function given as the ratio of two polynomials in the complex variable z . Each polynomial can be specified using either its coefficients or its roots. The roots of the numerator are the zeros of the transfer function and the roots of the denominator are the poles.

You may specify polynomials either in the complex variable z or $1/z$ by setting optional parameter `polyarg` to `z` or `inversez` respectively. By default, it is set to `inversez`. If you choose to provide the coefficients of a polynomial, enter them as a vector in ascending order of the power of the variable z or $1/z$, starting from the constant term. For example, to specify a denominator of $3z^{-2} + 4z^{-1} + 1$, use `denom=[1 4 3]`. Or to specify a denominator of $4z^2 + 3z - 2$, use `polyarg=z denom=[2 3 4]`.

To specify transfer function in terms of its zeros and poles in z -plane, give them as vectors of complex numbers. You must always give the real and imaginary portions of the root, even when the root is real. You may give either both roots of a complex-conjugate pair or only one. In the latter case the conjugate complex root will be generated automatically. The order of the roots is not important. For example, to specify poles of $z = 1$, $z = 4j$, $z = -4j$, $z = 2 + 2j$, and $z = 2 - 2j$, use `poles=[1 0 0 4 0 -4 2 2 2 -2]` or, omitting conjugate poles, `poles=[1 0 0 4 2 2]`.

Either the numerator or the denominator specification can be omitted. An omitted denominator or numerator is taken to be 1.

The parameter `gain` is interpreted either as the DC gain or, if the function has zeros or poles on the unit circle, as a constant factor.

Transition time (`tt`) is an optional parameter that at each sampling point forces linear transition of the output to a new value within the specified time range. By default, it is set to one percent of the sampling period.

The sampling delay (`td`) is another optional parameter, with the default value of 0, that lets you set asynchronous sampling rates.

To use the s to z transformation, set the optional `sxz` parameter to one of the transformation methods - forward differences, backward differences, or bilinear. When the `sxz` parameter is specified, the transfer function specification is assumed to be given in the complex variable s and it will be transformed to the complex variable z using the indicated method.

Spectre Circuit Simulator Reference

Component Statements Part III

This device is not supported within altergroup.

Sample Instance Statement

```
va (1 0) vsource type=sine freq=10K
z2 (2 0) zcccs probe=va gain=1 ts=4.9e-5 tt=1e-5 polyarg=inversez
numer=[1 -1] denom=[1 0]
```

Instance Definition

```
Name sink src zcccs parameter=value ...
```

Instance Parameters

1	probe	Device through which the controlling current flows.
2	port=0	Index of the probe port through which the controlling current flows.
3	ts=1 s	Sampling period.
4	td=0 s	Sampling delay.
5	tt=0.01 ts s	Transition time.
6	gain=1	Transfer function gain.
7	polyarg=inversez	Polynomial argument. Possible values are z or inversez.
8	szz=none	s to z transformation. Possible values are none, backward, forward, or bilinear.
9	numer=[...]	Vector of numerator coefficients.
10	denom=[...]	Vector of denominator coefficients.
11	zeros=[...]	Vector of complex zeros.
12	poles=[...]	Vector of complex poles.
13	m=1	Multiplicity factor.

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Component Statements Part III

Operating-Point Parameters

1	<code>i</code> (A)	Input current.
2	<code>v</code> (V)	Output voltage.
3	<code>pwr</code> (W)	Power dissipation.

Parameter Index

In the following index, `I` refers to instance parameters, `M` refers to the model parameters section, `O` refers to the output parameters section, and `OP` refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of `M-35` means the 35th model parameter.

<code>denom</code>	<code>I-10</code>	<code>numer</code>	<code>I-9</code>	<code>probe</code>	<code>I-1</code>	<code>ts</code>	<code>I-3</code>
<code>gain</code>	<code>I-6</code>	<code>poles</code>	<code>I-12</code>	<code>pwr</code>	<code>OP-3</code>	<code>tt</code>	<code>I-5</code>
<code>i</code>	<code>OP-1</code>	<code>polyarg</code>	<code>I-7</code>	<code>sxz</code>	<code>I-8</code>	<code>v</code>	<code>OP-2</code>
<code>m</code>	<code>I-13</code>	<code>port</code>	<code>I-2</code>	<code>td</code>	<code>I-4</code>	<code>zeros</code>	<code>I-11</code>

z-Domain Current Controlled Voltage Source (`zccvs`)

Description

The output is defined with a transfer function given as the ratio of two polynomials in the complex variable z . Each polynomial can be specified using either its coefficients or its roots. The roots of the numerator are the zeros of the transfer function and the roots of the denominator are the poles.

You may specify polynomials either in the complex variable z or $1/z$ by setting optional parameter `polyarg` to `z` or `inversez` respectively. By default, it is set to `inversez`. If you choose to provide the coefficients of a polynomial, enter them as a vector in ascending order of the power of the variable z or $1/z$, starting from the constant term. For example, to specify a denominator of $3z^{-2} + 4z^{-1} + 1$, use `denom=[1 4 3]`. Or to specify a denominator of $4z^2 + 3z - 2$, use `polyarg=z denom=[2 3 4]`.

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Component Statements Part III

To specify transfer function in terms of its zeros and poles in z-plane, give them as vectors of complex numbers. You must always give the real and imaginary portions of the root, even when the root is real. You may give either both roots of a complex-conjugate pair or only one. In the latter case the conjugate complex root will be generated automatically. The order of the roots is not important. For example, to specify poles of $z = 1$, $z = 4j$, $z = -4j$, $z = 2 + 2j$, and $z = 2 - 2j$, use `poles=[1 0 0 4 0 -4 2 2 2 -2]` or, omitting conjugate poles, `poles=[1 0 0 4 2 2]`.

Either the numerator or the denominator specification can be omitted. An omitted denominator or numerator is taken to be 1.

The parameter `gain` is interpreted either as the DC gain or, if the function has zeros or poles on the unit circle, as a constant factor.

Transition time (`tt`) is an optional parameter that at each sampling point forces linear transition of the output to a new value within the specified time range. By default, it is set to one percent of the sampling period.

The sampling delay (`td`) is another optional parameter, with the default value of 0, that lets you set asynchronous sampling rates.

To use the s to z transformation, set the optional `sxz` parameter to one of the transformation methods - forward differences, backward differences, or bilinear. When the `sxz` parameter is specified, the transfer function specification is assumed to be given in the complex variable s and it will be transformed to the complex variable z using the indicated method.

This device is not supported within altergroup.

Sample Instance Statement

```
va (1 0) vsource type=sine freq=10K
z2 2 0 zccvs probe=va gain=-2 ts=5e-5 tt=1.1e-5 numer=[1 -1]
```

Instance Definition

```
Name p n zccvs parameter=value ...
```

Instance Parameters

- | | |
|-----------------------|--|
| 1 <code>probe</code> | Device through which the controlling current flows. |
| 2 <code>port=0</code> | Index of the probe port through which the controlling current flows. |

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Component Statements Part III

3	<code>ts=1 s</code>	Sampling period.
4	<code>td=0 s</code>	Sampling delay.
5	<code>tt=0.01 ts s</code>	Transition time.
6	<code>gain=1</code>	Transfer function gain.
7	<code>polyarg=inversez</code>	Polynomial argument. Possible values are <code>z</code> or <code>inversez</code> .
8	<code>sxz=none</code>	<code>s</code> to <code>z</code> transformation. Possible values are <code>none</code> , <code>backward</code> , <code>forward</code> , or <code>bilinear</code> .
9	<code>numer=[...]</code>	Vector of numerator coefficients.
10	<code>denom=[...]</code>	Vector of denominator coefficients.
11	<code>zeros=[...]</code>	Vector of complex zeros.
12	<code>poles=[...]</code>	Vector of complex poles.
13	<code>m=1</code>	Multiplicity factor.

Operating-Point Parameters

1	<code>i (A)</code>	Output current.
2	<code>v (V)</code>	Output voltage.
3	<code>pwr (W)</code>	Power dissipation.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

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Component Statements Part III

denom	I-10	numer	I-9	probe	I-1	ts	I-3
gain	I-6	poles	I-12	pwr	OP-3	tt	I-5
i	OP-1	polyarg	I-7	sxz	I-8	v	OP-2
m	I-13	port	I-2	td	I-4	zeros	I-11

z-Domain Linear Voltage Controlled Current Source (zvccs)

Description

The output is defined with a transfer function given as the ratio of two polynomials in the complex variable z . Each polynomial can be specified using either its coefficients or its roots. The roots of the numerator are the zeros of the transfer function and the roots of the denominator are the poles.

You may specify polynomials either in the complex variable z or $1/z$ by setting optional parameter `polyarg` to `z` or `inversez` respectively. By default, it is set to `inversez`. If you choose to provide the coefficients of a polynomial, enter them as a vector in ascending order of the power of the variable z or $1/z$, starting from the constant term. For example, to specify a denominator of $3z^{-2} + 4z^{-1} + 1$, use `denom=[1 4 3]`. Or to specify a denominator of $4z^2 + 3z - 2$, use `polyarg=z denom=[2 3 4]`.

To specify transfer function in terms of its zeros and poles in z -plane, give them as vectors of complex numbers. You must always give the real and imaginary portions of the root, even when the root is real. You may give either both roots of a complex-conjugate pair or only one. In the latter case the conjugate complex root will be generated automatically. The order of the roots is not important. For example, to specify poles of $z = 1$, $z = 4j$, $z = -4j$, $z = 2 + 2j$, and $z = 2 - 2j$, use `poles=[1 0 0 4 0 -4 2 2 2 -2]` or, omitting conjugate poles, `poles=[1 0 0 4 2 2]`.

Either the numerator or the denominator specification can be omitted. An omitted denominator or numerator is taken to be 1.

The parameter `gain` is interpreted either as the DC gain or, if the function has zeros or poles on the unit circle, as a constant factor.

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Component Statements Part III

Transition time (t_t) is an optional parameter that at each sampling point forces linear transition of the output to a new value within the specified time range. By default, it is set to one percent of the sampling period.

The sampling delay (t_d) is another optional parameter, with the default value of 0, that lets you set asynchronous sampling rates.

To use the s to z transformation, set the optional `sxz` parameter to one of the transformation methods - forward differences, backward differences, or bilinear. When the `sxz` parameter is specified, the transfer function specification is assumed to be given in the complex variable s and it will be transformed to the complex variable z using the indicated method.

This device is not supported within altergroup.

Sample Instance Statement

```
va (1 0) vsource type=sine freq=10K
z1 (2 0 1 0) zvccs gain=2 ts=4.5e-5 tt=1e-5 zeros=[-1 0] poles=[0 0]
```

Instance Definition

```
Name sink src ps ns zvccs parameter=value ...
```

Instance Parameters

- | | | |
|---|-------------------------------|---|
| 1 | <code>ts=1 s</code> | Sampling period. |
| 2 | <code>td=0 s</code> | Sampling delay. |
| 3 | <code>tt=0.01 ts s</code> | Transition time. |
| 4 | <code>gain=1</code> | Transfer function gain. |
| 5 | <code>polyarg=inversez</code> | Polynomial argument.
Possible values are <code>z</code> or <code>inversez</code> . |
| 6 | <code>sxz=none</code> | s to z transformation.
Possible values are <code>none</code> , <code>backward</code> , <code>forward</code> , or <code>bilinear</code> . |
| 7 | <code>numer=[...]</code> | Vector of numerator coefficients. |
| 8 | <code>denom=[...]</code> | Vector of denominator coefficients. |

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Component Statements Part III

- 9 `zeros=[...]` Vector of complex zeros.
- 10 `poles=[...]` Vector of complex poles.
- 11 `m=1` Multiplicity factor.

Operating-Point Parameters

- 1 `i` (A) Output current.
- 2 `v` (V) Output voltage.
- 3 `pwr` (W) Power dissipation.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the description for that parameter. For example, a reference of **M-35** means the 35th model parameter.

<code>denom</code>	I-8	<code>numer</code>	I-7	<code>sxz</code>	I-6	<code>v</code>	OP-2
<code>gain</code>	I-4	<code>poles</code>	I-10	<code>td</code>	I-2	<code>zeros</code>	I-9
<code>i</code>	OP-1	<code>polyarg</code>	I-5	<code>ts</code>	I-1		
<code>m</code>	I-11	<code>pwr</code>	OP-3	<code>tt</code>	I-3		

z-Domain Voltage Controlled Voltage Source (zvcvs)

Description

The output is defined with a transfer function given as the ratio of two polynomials in the complex variable z . Each polynomial can be specified using either its coefficients or its roots. The roots of the numerator are the zeros of the transfer function and the roots of the denominator are the poles.

You may specify polynomials either in the complex variable z or $1/z$ by setting optional parameter `polyarg` to `z` or `inversez` respectively. By default, it is set to `inversez`. If you

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Component Statements Part III

choose to provide the coefficients of a polynomial, enter them as a vector in ascending order of the power of the variable z or $1/z$, starting from the constant term. For example, to specify a denominator of $3z^{-2} + 4z^{-1} + 1$, use `denom=[1 4 3]`. Or to specify a denominator of $4z^2 + 3z - 2$, use `polyarg=z denom=[2 3 4]`.

To specify transfer function in terms of its zeros and poles in z -plane, give them as vectors of complex numbers. You must always give the real and imaginary portions of the root, even when the root is real. You may give either both roots of a complex-conjugate pair or only one. In the latter case the conjugate complex root will be generated automatically. The order of the roots is not important. For example, to specify poles of $z = 1$, $z = 4j$, $z = -4j$, $z = 2 + 2j$, and $z = 2 - 2j$, use `poles=[1 0 0 4 0 -4 2 2 2 -2]` or, omitting conjugate poles, `poles=[1 0 0 4 2 2]`.

Either the numerator or the denominator specification can be omitted. An omitted denominator or numerator is taken to be 1.

The parameter `gain` is interpreted either as the DC gain or, if the function has zeros or poles on the unit circle, as a constant factor.

Transition time (`tt`) is an optional parameter that at each sampling point forces linear transition of the output to a new value within the specified time range. By default, it is set to one percent of the sampling period.

The sampling delay (`td`) is another optional parameter, with the default value of 0, that lets you set asynchronous sampling rates.

To use the s to z transformation, set the optional `sxz` parameter to one of the transformation methods - forward differences, backward differences, or bilinear. When the `sxz` parameter is specified, the transfer function specification is assumed to be given in the complex variable s and it will be transformed to the complex variable z using the indicated method.

This device is not supported within `altergroup`.

Sample Instance Statement

```
va (1 0) vsource type=sine freq=10K
z3 (3 0 1 0) zvcvs gain=-1 ts=4e-5 tt=1e-5 numer=[-1 -1]
```

Instance Definition

```
Name p n ps ns zvcvs parameter=value ...
```

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Instance Parameters

1	<code>ts=1 s</code>	Sampling period.
2	<code>td=0 s</code>	Sampling delay.
3	<code>tt=0.01 ts s</code>	Transition time.
4	<code>gain=1</code>	Transfer function gain.
5	<code>polyarg=inversez</code>	Polynomial argument. Possible values are <code>z</code> or <code>inversez</code> .
6	<code>sxz=none</code>	s to z transformation. Possible values are <code>none</code> , <code>backward</code> , <code>forward</code> , or <code>bilinear</code> .
7	<code>numer=[...]</code>	Vector of numerator coefficients.
8	<code>denom=[...]</code>	Vector of denominator coefficients.
9	<code>zeros=[...]</code>	Vector of complex zeros.
10	<code>poles=[...]</code>	Vector of complex poles.
11	<code>m=1</code>	Multiplicity factor.

Operating-Point Parameters

1	<code>i (A)</code>	Output current.
2	<code>v (V)</code>	Output voltage.
3	<code>pwr (W)</code>	Power dissipation.

Parameter Index

In the following index, **I** refers to instance parameters, **M** refers to the model parameters section, **O** refers to the output parameters section, and **OP** refers to the operating point parameters section. The number indicates where to look in the appropriate section to find the

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Component Statements Part III

description for that parameter. For example, a reference of M-35 means the 35th model parameter.

denom	I-8	numer	I-7	szx	I-6	v	OP-2
gain	I-4	poles	I-10	td	I-2	zeros	I-9
i	OP-1	polyarg	I-5	ts	I-1		
m	I-11	pwr	OP-3	tt	I-3		

Analysis Statements

This chapter discusses the following topics:

- [AC Analysis \(ac\)](#) on page 702
- [Alter a Circuit, Component, or Netlist Parameter \(alter\)](#) on page 705
- [Alter Group \(altergroup\)](#) on page 705
- [Check Parameter Values \(check\)](#) on page 707
- [Checklimit Analysis \(checklimit\)](#) on page 708
- [DC Analysis \(dc\)](#) on page 709
- [DC Device Matching Analysis \(dcmatch\)](#) on page 713
- [Envelope Following Analysis \(envlp\)](#) on page 717
- [Circuit Information \(info\)](#) on page 722
- [Monte Carlo Analysis \(montecarlo\)](#) on page 724
- [Noise Analysis \(noise\)](#) on page 735
- [Immediate Set Options \(options\)](#) on page 739
- [Periodic AC Analysis \(pac\)](#) on page 748
- [Periodic Distortion Analysis \(pdisto\)](#) on page 752
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- [Periodic Steady-State Analysis \(pss\)](#) on page 771
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Analysis Statements

- [Quasi-Periodic Noise Analysis \(qpnoise\)](#) on page 798
- [Quasi-Periodic S-Parameter Analysis \(qpssp\)](#) on page 804
- [Quasi-Periodic Steady State Analysis \(qpss\)](#) on page 810
- [Quasi-Periodic Transfer Function Analysis \(qpxf\)](#) on page 817
- [Deferred Set Options \(set\)](#) on page 821
- [Shell Command \(shell\)](#) on page 825
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- [Stability Analysis \(stb\)](#) on page 830
- [Sweep Analysis \(sweep\)](#) on page 835
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- [Transient Analysis \(tran\)](#) on page 839
- [Transfer Function Analysis \(xf\)](#) on page 848

AC Analysis (ac)

Description

The AC analysis linearizes the circuit about the DC operating point and computes the response to a given small sinusoidal stimulus.

Spectre can perform the analysis while sweeping a parameter. The parameter can be frequency, temperature, component instance parameter, component model parameter, or netlist parameter. If changing a parameter affects the DC operating point, the operating point is recomputed on each step. You can sweep the circuit temperature by giving the parameter name as `temp` with no `dev` or `mod` parameter. You can sweep a netlist parameter by giving the parameter name with no `dev`, or `mod` parameter. After the analysis has completed, the modified parameter returns to its original value.

Definition

Name `ac parameter=value ...`

Parameters

1 `prevoppoint=no` Use operating point computed on the previous analysis.
Possible values are `no` or `yes`.

Sweep interval parameters

2 `start=0` Start sweep limit.

3 `stop` Stop sweep limit.

4 `center` Center of sweep.

5 `span=0` Sweep limit span.

6 `step` Step size, linear sweep.

7 `lin=50` Number of steps, linear sweep.

8 `dec` Points per decade.

9 `log=50` Number of steps, log sweep.

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Analysis Statements

10 `values=[...]` Array of sweep values.

Sweep variable parameters

11 `dev` Device instance whose parameter value is to be swept.

12 `mod` Model whose parameter value is to be swept.

13 `param` Name of parameter to sweep.

14 `freq (Hz)` Frequency when parameter other than frequency is being swept.

State-file parameters

15 `readns` File that contains estimate of DC solution (nodeset).

Output parameters

16 `save` Signals to output.
Possible values are `all`, `lvl`, `allpub`, `lvlpub`, `selected`, or `none`.

17 `nestlvl` Levels of subcircuits to output.

18 `oppooint=no` Should operating point information be computed, and if so, where should it be sent. Operating point information would not be output if operating point is computed in the previous analysis and is unchanged.
Possible values are `no`, `screen`, `logfile`, or `rawfile`.

Convergence parameters

19 `restart=yes` Restart the DC solution from scratch if any condition has changed. If not, use the previous solution as initial guess.
Possible values are `no` or `yes`.

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Analysis Statements

Annotation parameters

- 20 `annotate=sweep` Degree of annotation.
Possible values are `no`, `title`, `sweep`, `status`, or `steps`.
- 21 `stats=no` Analysis statistics.
Possible values are `no` or `yes`.
- 22 `title` Analysis title.

You can specify sweep limits by giving the end points or by providing the center value and the span of the sweep. Steps can be linear or logarithmic, and you can specify the number of steps or the size of each step. You can give a step size parameter (`step`, `lin`, `log`, `dec`) to determine whether the sweep is linear or logarithmic. If you do not give a step size parameter, the sweep is linear when the ratio of stop to start values is less than 10, and logarithmic when this ratio is 10 or greater. All frequencies are in Hertz.

The small-signal analysis begins by linearizing the circuit about an operating-point. By default this analysis computes the operating-point if it is not known, or recomputes it if any significant component or circuit parameter has changed. However, if a previous analysis computed an operating point, you can set `prevoppoint=yes` to avoid recomputing it. For example, if you use this option when the previous analysis was a transient analysis, the operating point is the state of the circuit on the final time point.

Parameter Index

In the following index, the number following each parameter name indicates where to find the description of that parameter.

<code>annotate</code>	20	<code>log</code>	9	<code>readns</code>	15	<code>step</code>	6
<code>center</code>	4	<code>mod</code>	12	<code>restart</code>	19	<code>stop</code>	3
<code>dec</code>	8	<code>nestlvl</code>	17	<code>save</code>	16	<code>title</code>	22
<code>dev</code>	11	<code>oppoint</code>	18	<code>span</code>	5	<code>values</code>	10
<code>freq</code>	14	<code>param</code>	13	<code>start</code>	2		
<code>lin</code>	7	<code>prevoppoint</code>	1	<code>stats</code>	21		

Alter a Circuit, Component, or Netlist Parameter (alter)

Description

The `alter` statement changes the value of any modifiable component or netlist parameter for any analyses that follow. The parameter to be altered can be circuit temperature, a device instance parameter, a device model parameter, a netlist parameter, or a subcircuit parameter for a particular subcircuit instance. You can alter the circuit temperature by giving the parameter name as `param=temp` with no `dev`, `mod` or `sub` parameter. You can alter a top-level netlist parameter by giving the parameter name with no `dev`, `mod` or `sub` parameter. You can alter a subcircuit parameter for a particular subcircuit instance by specifying the subcircuit instance name with the `sub` parameter, and the subcircuit parameter name with the `param` parameter. Each `alter` statement can change only one parameter.

Definition

```
Name alter parameter=value ...
```

Parameters

1	<code>mod</code>	Device model.
2	<code>dev</code>	Device instance.
3	<code>sub</code>	Subcircuit instance.
4	<code>param</code>	Name of parameter to be altered.
5	<code>value</code>	New value for parameter.
6	<code>annotate</code>	Degree of annotation. Possible values are <code>no</code> or <code>title</code> .

Alter Group (altergroup)

Description

The `altergroup` statement changes the values of any modifiable model, instance or netlist parameter for any analyses that follow. Within an alter group, you can specify model statements, instance statements and parameter statements. These statements should be

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Analysis Statements

bound within braces. The opening brace is required at the end of the line defining the alter group. Alter groups cannot be nested or specified within subcircuits. The following statements are not allowed within altergroups (analyses, export, ic, nodeset, paramset, save, and sens).

Within an alter group, each device (instance or model) is first defaulted and then the device parameters are updated. For netlist parameters, the expressions are updated and evaluated.

For subckt within altergroup, all instances of the subckts are modified during the altergroup. There are strict checks that do not allow changes to topology.

You can include files into the alter group and can use the `simulator lang=spice` directive. See `spectre -h include` for more details. A model defined in the netlist, has to have the same model name and primitive type (`bsim2`, `bsim3`, `bjt`) in the alter group. An instance defined in the netlist, has to have the same instance name, terminal connections and primitive type. For model groups you can change the number of models in the group. There is a restriction that you cannot change from a model to a model group and vice versa. See `spectre -h bsim3v3` for details on model groups.

Definition

Name altergroup parameter=value ...

Parameters

1 annotate Degree of annotation.
Possible values are no or title.

Example:

```
FastCorner altergroup {
    parameters p2=1 p3=p1+2
    model myres resistor r1=1e3 af=1
    model mybsim bsim3v3 lmax=p1 lmin=3.5e-7
    m1 (n1 n2 n3 n4) mybsim w=0.3u l=1.2u
}
```

The list of public devices supported by altergroup:

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bjt	bjt301	bjt500	bjt503
bjt504	bjt504t	bsim1	bsim2
bsim3	bsim3v3	bsim4	bsimsoi
btasoi	capacitor	cccs	ccvs
dio500	diode	ekv	gaas
hbt	hisim	hvmos	inductor
intcap	isource	jfet	juncap
mos1	mos2	mos3	mos30
mos40	mos705	mos902	mos903
mos1000	mos1100	mos3002	mos3100
mos11010	mos11011	msline	nodcap
phy_res	psitft	rdiff	resistor
tline	tom2	tom3	vbic
vccs	vcvs	vsources	

The list of public devices not supported by altergroup:

a2d	assert	b3soipd	cktrom
core	d2a	delay	iprobe
misnan	mos0	mos15	mtline
mutual_inductor	port	pcccs	pccvs
port	pvccs	pvcvs	relay
scccs	sccvs	svccs	svcvs
switch	transformer	winding	zcccs
zccvs	zvccs	zvcvs	

Check Parameter Values (check)

Description

The `check` analysis checks the values of component parameters to assure they are reasonable. This analysis reduces the cost of data entry errors. Various filters specify which parameters are checked. You can perform checks on input, output, or operating-point parameters. Use this analysis in conjunction with the `+param` command line argument, which specifies a file that contains component parameter soft limits.

Definition

Name `check parameter=value ...`

Parameters

1 `what=all` What parameters should be checked.
Possible values are `none`, `inst`, `models`, `input`, `output`, `all`, or `oppooint`.

Checklimit Analysis (`checklimit`)

Description

A `checklimit` analysis allows the enabling or disabling of individual or group of `asserts` specified in the netlist. Use this analysis in conjunction with `assert` statements in the netlist to perform checks on parameters of device instances, models, subcircuits or expressions

Multiple `checklimit` analyses maybe defined in the netlist. The enabled checks will be applied to all subsequent analyses until the next `checklimit` analysis is encountered.

Definition

Name `checklimit` parameter=value ...

Parameters

- | | | |
|---|----------------------------------|--|
| 1 | <code>enable=[...]</code> | Array of checks to be enabled. Default is all. |
| 2 | <code>disable=[...]</code> | Array of checks to be disabled. Default is none. |
| 3 | <code>start=0 s</code> | Start(time) of the checks. |
| 4 | <code>stop (s)</code> | Stop(time) of the checks. |
| 5 | <code>severity</code> | Severity of the checks.
Possible values are <code>none</code> , <code>notice</code> , <code>warning</code> , or <code>error</code> . |
| 6 | <code>title</code> | Analysis title. |
| 7 | <code>checkallasserts=yes</code> | If all checks should be enabled or disabled. <code>CheckAllAsserts=no</code> will disable all checks.
Possible values are <code>no</code> or <code>yes</code> . |

DC Analysis (dc)

Description

The DC analysis finds the DC operating-point or DC transfer curves of the circuit. To generate transfer curves, specify a parameter and a sweep range. The swept parameter can be circuit temperature, a device instance parameter, a device model parameter, a netlist parameter, or a subcircuit parameter for a particular subcircuit instance. You can sweep the circuit temperature by giving the parameter name as `param=temp` with no `dev`, `mod` or `sub` parameter. You can sweep a top-level netlist parameter by giving the parameter name with no `dev`, `mod` or `sub` parameter. You can sweep a subcircuit parameter for a particular subcircuit instance by specifying the subcircuit instance name with the `sub` parameter, and the subcircuit parameter name with the `param` parameter. After the analysis has completed, the modified parameter returns to its original value.

Definition

Name `dc parameter=value ...`

Parameters

Sweep interval parameters

1	<code>start=0</code>	Start sweep limit.
2	<code>stop</code>	Stop sweep limit.
3	<code>center</code>	Center of sweep.
4	<code>span=0</code>	Sweep limit span.
5	<code>step</code>	Step size, linear sweep.
6	<code>lin=50</code>	Number of steps, linear sweep.
7	<code>dec</code>	Points per decade.
8	<code>log=50</code>	Number of steps, log sweep.
9	<code>values=[...]</code>	Array of sweep values.

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Analysis Statements

Sweep variable parameters

10	<code>dev</code>	Device instance whose parameter value is to be swept.
11	<code>mod</code>	Model whose parameter value is to be swept.
12	<code>param</code>	Name of parameter to sweep.

State-file parameters

13	<code>force=none</code>	What should be used to force values for DC. Uses the values from the device and node ICs. Possible values are <code>none</code> , <code>node</code> , <code>dev</code> , or <code>all</code> .
14	<code>readns</code>	File that contains estimate of DC solution (nodeset).
15	<code>readforce</code>	File that contains force values.
16	<code>write</code>	File to which solution at first step in sweep is written.
17	<code>writefinal</code>	File to which solution at last step in sweep is written.

Output parameters

18	<code>save</code>	Signals to output. Possible values are <code>all</code> , <code>lvl</code> , <code>allpub</code> , <code>lvlpub</code> , <code>selected</code> , or <code>none</code> .
19	<code>nestlvl</code>	Levels of subcircuits to output.
20	<code>print=no</code>	Print node voltages. Possible values are <code>no</code> or <code>yes</code> .
21	<code>oppoint=no</code>	Should operating point information be computed, and if so, where should it be sent. Operating point information would not be output if sweep parameter <code>param</code> is set. Possible values are <code>no</code> , <code>screen</code> , <code>logfile</code> , or <code>rawfile</code> .
22	<code>check=yes</code>	Check operating point parameters against soft limits. Possible values are <code>no</code> or <code>yes</code> .

Spectre Circuit Simulator Reference

Analysis Statements

Convergence parameters

- | | | |
|----|-----------------------------|---|
| 23 | <code>homotopy=all</code> | Method used when no convergence on initial attempt of DC analysis.
Possible values are <code>none</code> , <code>gmin</code> , <code>source</code> , <code>dptran</code> , <code>ptran</code> , <code>arclength</code> , or <code>all</code> . |
| 24 | <code>restart=yes</code> | Restart from scratch if any condition has changed. If not, use the previous solution as initial guess.
Possible values are <code>no</code> or <code>yes</code> . |
| 25 | <code>maxiters=150</code> | Maximum number of iterations. |
| 26 | <code>maxsteps=10000</code> | Maximum number of steps used in homotopy method. |

Annotation parameters

- | | | |
|----|-----------------------------|--|
| 27 | <code>annotate=sweep</code> | Degree of annotation.
Possible values are <code>no</code> , <code>title</code> , <code>sweep</code> , <code>status</code> , or <code>steps</code> . |
| 28 | <code>title</code> | Analysis title. |

You can specify sweep limits by giving the end points or by providing the center value and the span of the sweep. Steps can be linear or logarithmic, and you can specify the number of steps or the size of each step. You can give a step size parameter (`step`, `lin`, `log`, `dec`) and determine whether the sweep is linear or logarithmic. If you do not give a step size parameter, the sweep is linear when the ratio of stop to start values is less than 10, and logarithmic when this ratio is 10 or greater. If you specify the `oppoint` parameter, Spectre computes and outputs the linearized model for each nonlinear component.

Nodesets help find the DC or initial transient solution. You can supply them in the circuit description file with `nodeset` statements, or in a separate file using the `readns` parameter. When nodesets are given, Spectre computes an initial guess of the solution by performing a DC analysis while forcing the specified values onto nodes by using a voltage source in series with a resistor whose resistance is `rforce`. Spectre then removes these voltage sources and resistors and computes the true solution from this initial guess.

Nodesets have two important uses. First, if a circuit has two or more solutions, nodesets can bias the simulator towards computing the desired one. Second, they are a convergence aid. By estimating the solution of the largest possible number of nodes, you might be able to eliminate a convergence problem or dramatically speed convergence.

Spectre Circuit Simulator Reference

Analysis Statements

When you simulate the same circuit many times, we suggest that you use both the `write` and `readns` parameters and give the same file name to both parameters. The DC analysis then converges quickly even if the circuit has changed somewhat since the last simulation, and the `nodeset` file is automatically updated.

You may specify values to force for the DC analysis by setting the parameter `force`. The values used to force signals are specified by using the `force` file, the `ic` statement, or the `ic` parameter on the capacitors and inductors. The `force` parameter controls the interaction of various methods of setting the force values. The effects of individual settings are:

`force=none`: Any initial condition specifiers are ignored.

`force=node`: The `ic` statements are used, and the `ic` parameter on the capacitors and inductors are ignored.

`force=dev`: The `ic` parameters on the capacitors and inductors are used, and the `ic` statements are ignored.

`force=all`: Both the `ic` statements and the `ic` parameters are used, with the `ic` parameters overriding the `ic` statements.

If you specify a `force` file with the `readforce` parameter, force values read from the file are used, and any `ic` statements are ignored.

Once you specify the force conditions, Spectre computes the DC analysis with the specified nodes forced to the given value by using a voltage source in series with a resistor whose resistance is `rforce` (see `options`).

Parameter Index

In the following index, the number following each parameter name indicates where to find the description of that parameter.

<code>annotate</code>	27	<code>lin</code>	6	<code>param</code>	12	<code>start</code>	1
<code>center</code>	3	<code>log</code>	8	<code>print</code>	20	<code>step</code>	5
<code>check</code>	22	<code>maxiters</code>	25	<code>readforce</code>	15	<code>stop</code>	2
<code>dec</code>	7	<code>maxsteps</code>	26	<code>readns</code>	14	<code>title</code>	28
<code>dev</code>	10	<code>mod</code>	11	<code>restart</code>	24	<code>values</code>	9
<code>force</code>	13	<code>nestlvl</code>	19	<code>save</code>	18	<code>write</code>	16
<code>homotopy</code>	23	<code>oppoint</code>	21	<code>span</code>	4	<code>writefinal</code>	17

DC Device Matching Analysis (dcmatch)

Description

The DCMATCH analysis performs DC device mis-matching analysis for a given output. It computes the deviation in the DC operating point of the circuit caused by mismatch in the devices. Users need to specify mismatch parameters in their model cards for each device contributing to the deviation. The analysis uses the device mismatch models to construct equivalent mismatch current sources to all the devices that have mismatch modeled. These current sources will have zero mean and some variance. The variance of the current sources are computed according to mismatch models. It then computes the 3-sigma variance of dc voltages or currents at user specified outputs due to the mismatch current sources. The simulation results displays the devices rank ordered by their contributions to the outputs. In addition, for mosfet devices, it displays threshold voltage mismatch, current factor mismatch, gate voltage mismatch, and drain current mismatch. For bipolar devices, it displays base-emitter junction voltage mismatch. For resistors, it displays resistor mismatches.

The analysis replaces multiple simulation runs by circuit designers for accuracy vs. size analysis. It automatically identifies the set of critical matched components during circuit design. For example, when there are matched pairs in the circuit, the contribution of two matched transistors will be equal in magnitude and opposite in sign. Typical usage are to simulate the output offset voltage of operational amplifiers, estimate the variation in bandgap voltages, and predict the accuracy of current steering DACs.

Definition

Name ... dcmatch parameter=value ...

Parameters

- | | | |
|---|----------------------------|---|
| 1 | <code> mth</code> | Relative mismatch contribution threshold value. |
| 2 | <code> where=screen</code> | Where DC-Mismatch analysis results should be printed.
Possible values are <code>screen</code> , <code>logfile</code> , <code>file</code> , or <code>rawfile</code> . |
| 3 | <code> file</code> | File name for results to be printed if <code>where=file</code> is used. |

Probe parameters

- | | | |
|---|----------------------|---|
| 4 | <code> oprobe</code> | Compute mismatch at the output defined by this component. |
|---|----------------------|---|

Spectre Circuit Simulator Reference

Analysis Statements

Port parameters

5	<code>portv</code>	Voltage across this probe port is output of the analysis.
6	<code>porti</code>	Current through this probe port is output of the analysis.

Sweep interval parameters

7	<code>start=0</code>	Start sweep limit.
8	<code>stop</code>	Stop sweep limit.
9	<code>center</code>	Center of sweep.
10	<code>span=0</code>	Sweep limit span.
11	<code>step</code>	Step size, linear sweep.
12	<code>lin=50</code>	Number of steps, linear sweep.
13	<code>dec</code>	Points per decade.
14	<code>log=50</code>	Number of steps, log sweep.
15	<code>values=[...]</code>	Array of sweep values.

Sweep variable parameters

16	<code>dev</code>	Device instance whose parameter value is to be swept.
17	<code>mod</code>	Model whose parameter value is to be swept.
18	<code>param</code>	Name of parameter to sweep.

State-file parameters

19	<code>readns</code>	File that contains estimate of DC solution (nodeset).
----	---------------------	---

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Analysis Statements

Output parameters

- 20 `save` Signals to output.
Possible values are `all`, `lvl`, `allpub`, `lvlpub`, `selected`, or `none`.
- 21 `nestlvl` Levels of subcircuits to output.
- 22 `oppoint=no` Should operating point information be computed, and if so, where should it be sent. Operating point information would not be output if (1) operating point is computed in the previous analysis and is unchanged, or (2) sweep parameter `param` is set.
Possible values are `no`, `screen`, `logfile`, or `rawfile`.

Convergence parameters

- 23 `prevoppoint=no` Use operating point computed on the previous analysis.
Possible values are `no` or `yes`.
- 24 `restart=yes` Restart the DC solution from scratch if any condition has changed. If not, use the previous solution as initial guess.
Possible values are `no` or `yes`.

Annotation parameters

- 25 `annotate=sweep` Degree of annotation.
Possible values are `no`, `title`, `sweep`, `status`, or `steps`.
- 26 `stats=no` Analysis statistics.
Possible values are `no` or `yes`.
- 27 `title` Analysis title.

The `dcmatch` analysis will find a dc operating point first. If the dc analysis fails, then the `dcmatch` analysis will fail also. The parameter `mth` is a threshold value relative to maximum contribution. Any device contribution less than (`mth * maximum`) will not be reported. Where maximum is the maximum contribution among all the devices of a given type.

Examples:

```
dcmm1 dcmatch mth=1e-3 oprobe=vd porti=1
```

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```
dcmm2 dcmatch mth=1e-3 oprobe=r3 portv=1
dcmm3 n1 n2 dcmatch mth=1e-3 where=rawfile stats=yes
dcmm4 n3 0 dcmatch mth=1e-3 where=file file="%C:r.info.what"
sweep1 sweep dev=mp6 param=w start=80e-6 stop=90e-6 step=2e-6 {
dcmm5 dcmatch oprobe=vd mth=1e-3 where=rawfile }
dcmm6 n3 0 dcmatch mth=0.01 dev=x1.mp2 param=w start=15e-6 stop=20e-6 step=1e-6
dcmm7 n3 0 dcmatch mth=0.01 param=temp start=25 stop=100 step=25
```

Note:

portv allows users to select a current associated with a specific device given in oprobe as an output. This device, however, has to have its terminal currents as network variables, i.e. the device has to be an inductor, a vsource, a switch, a tline, a controlled voltage source, an iprobe, or other type of device which has current solution. Further, for inductor, vsource, switch, controlled voltage source and iprobe, portv can only be set to one, since these devices are two terminal devices (one port); and for tline portv can be set to one or two, since it is a four terminal device (two ports).

Parameter Index

In the following index, the number following each parameter name indicates where to find the description of that parameter.

annotate	25	mod	17	portv	5	stats	26
center	9	mth	1	prevoppoint	23	step	11
dec	13	nestlvl	21	readns	19	stop	8
dev	16	oppoint	22	restart	24	title	27
file	3	oprobe	4	save	20	values	15
lin	12	param	18	span	10	where	2
log	14	portv	6	start	7		

Envelope Following Analysis (envlp)

Description

This analysis computes the envelope response of a circuit. The user specifies the analysis `clockname`. The simulator automatically determines the clock period by looking through all the sources with the specified name. The envelope response is computed over the interval from `start` to `stop`. If the interval is not a multiple of the clock period, it is rounded off to the nearest multiple before the stop time. The initial condition is taken to be the DC steady-state solution if not otherwise given.

Envelope following analysis is most efficient for circuits where the modulation bandwidth is orders of magnitude lower than the clock frequency. This is typically the case, for example, in circuits where the clock is the only fast varying signal and other input signals have a spectrum whose frequency range is orders of magnitude lower than the clock frequency. For another example, the down conversion of two closely placed frequencies can also generate a slow-varying modulation envelope.

The analysis generates two types of output files, a voltage versus time (`td`) file, and an amplitude/phase versus time (`fd`) file for each of specified harmonic of the clock fundamental.

Definition

Name `envlp parameter=value ...`

Parameters

Envelope fundamental parameters

- 1 `clockname` Name of the clock fundamental.
- 2 `modulationbw (Hz)` Modulation bandwidth.

Simulation interval parameters

- 3 `stop (s)` Stop time.
- 4 `start=0 s` Start time.
- 5 `tstab=0 s` Initial stabilization time.

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Analysis Statements

6 `outputstart=start s` Output is saved only after this time is reached.

Time-step parameters

7 `maxstep (s)` Maximum time step for inner transient integration. Default derived from `errpreset`.

8 `envmaxstep (s)` Maximum outer envelope step size. Default derived from `errpreset`.

Initial-condition parameters

9 `ic=all` What should be used to set initial condition. Possible values are `dc`, `node`, `dev`, or `all`.

10 `skipdc=no` If yes, there will be no dc analysis for initial transient. Possible values are `no` or `yes`.

11 `readic` File that contains initial transient condition.

Convergence parameters

12 `readns` File that contains estimate of initial DC solution.

13 `cmin=0 F` Minimum capacitance from each node to ground.

State-file parameters

14 `write` File to which initial transient solution is to be written.

15 `writefinal` File to which final transient solution is to be written.

16 `swapfile` Temporary file that holds the matrix information used by Newtons method. Tells Spectre to use a regular file rather than virtual memory to hold the matrix information. Use this option if Spectre complains about not having enough memory to complete this analysis.

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Analysis Statements

Integration method parameters

17 `method` Integration method. Default derived from `errpreset`. Possible values are `euler`, `trap`, `traponly`, `gear2`, or `gear2only`.

Accuracy parameters

18 `errpreset=moderate` Selects a reasonable collection of parameter settings. Possible values are `liberal`, `moderate` or `conservative`.

19 `relref` Reference used for the relative convergence criteria. Default derived from `errpreset`. Possible values are `pointlocal`, `alllocal`, `sigglobal`, or `allglobal`.

20 `lteratio` Ratio used to compute LTE tolerances from Newton tolerance. Default derived from `errpreset`.

21 `steadyratio` Ratio used to compute steady state tolerances from LTE tolerance. Default derived from `errpreset`.

22 `envlteratio` Ratio used to compute envelope LTE tolerances. Default derived from `errpreset`.

Annotation parameters

23 `stats=no` Analysis statistics. Possible values are `no` or `yes`.

24 `annotate=sweep` Degree of annotation. Possible values are `no`, `title`, `sweep`, `status`, or `steps`.

25 `title` Analysis title.

Output parameters

26 `harms=1` Number of clock harmonics to output.

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Analysis Statements

- 27 `harmsvec=[...]` Array of desired clock harmonics. Alternate form of `harms` that allows selection of specific harmonics.
- 28 `outputtype=both` Output type.
Possible values are `both`, `envelope` or `spectrum`.
- 29 `save` Signals to output.
Possible values are `all`, `lvl`, `allpub`, `lvlpub`, `selected`, or `none`.
- 30 `nestlvl` Levels of subcircuits to output.
- 31 `compression=no` Do data compression on output. See full description below.
Possible values are `no` or `yes`.
- 32 `strobeperiod (s)` The output strobe interval (in seconds of envelope following time). The actual strobe interval is rounded to an integer multiple of the clock period.

Newton parameters

- 33 `maxiters=5` Maximum number of Newton iterations per transient integration time step.
- 34 `envmaxiters=3` Maximum number of Newton iterations per envelope step.
- 35 `restart=yes` Restart the DC solution from scratch if any condition has changed. If not, use the previous solution as initial guess.
Possible values are `no` or `yes`.

Circuit age

- 36 `circuitage (Years)` Stress Time. Age of the circuit used to simulate hot-electron degradation of MOSFET and BSIM circuits.

The simulator examines all the sources whose name matches the clock name specified in the analysis line by the `clockname` parameter to determine the clock frequency. If more than one frequencies are found, the least common factor of these frequencies is used as the clock frequency.

The maximum envelope step size is affected by many parameters. It can be directly limited by `envmaxstep`. It is also limited by `modulationbw`. The user gives an estimate of the

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Analysis Statements

modulation bandwidth. The simulator will put at least eight points within the modulation period.

The `harms` and `harmsvec` parameters affect the simulation time in a significant way. The spectrum is calculated for all the specified harmonics for all sampled integration cycles as the envelope following analysis marches on. For each harmonic, a file is generated. The user should refrain from specifying unnecessary harmonics. Typically, `harms` is set to 1 or 2.

Most parameters of this analysis are inherited from either transient or PSS analysis and their meanings are consistent. However, a few of them need to be clarified. The effect of `errpreset` on some particular envelope following analysis parameters is shown in the following table.

Parameter defaults as a function of `errpreset`

<code>errpreset</code>	<code>envmaxstep</code>	<code>steadyratio</code>	<code>envlteratio</code>
liberal	Interval/10	1.0	10.0
moderate	Interval/50	0.1	1.0
conservative	Interval/100	0.01	0.1

Its effect on parameters such as `reltol`, `relref`, `method`, `maxstep`, and `lteratio` are the same as defined for transient analysis, except for that the transient simulation interval is always a clock period.

The default value for `compression` is `no`. The output file stores data for every signal at every time point for which Spectre calculates a solution. Spectre saves the x axis data only once, since every signal has the same x value. If `compression=yes`, Spectre writes data to the output file only if the signal value changes by at least 2*the convergence criteria. In order to save data for each signal independently, x axis information corresponding to each signal must be saved. If the signals stay at constant values for large periods of the simulation time, setting `compression=yes` results in a smaller output data file. If the signals in your circuit move around a lot, setting `compression=yes` results in a larger output data file.

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Analysis Statements

Parameter Index

In the following index, the number following each parameter name indicates where to find the description of that parameter.

annotate	24	harms	26	outputstart	6	stats	23
circuitage	36	harmsvec	27	outputtype	28	steadyratio	21
clockname	1	ic	9	readic	11	stop	3
cmin	13	lteratio	20	readns	12	strobeperiod	32
compression	31	maxiters	33	relref	19	swapfile	16
envlteratio	22	maxstep	7	restart	35	title	25
envmaxiters	34	method	17	save	29	tstab	5
envmaxstep	8	modulationbw	2	skipdc	10	write	14
errpreset	18	nestlvl	30	start	4	writefinal	15

Circuit Information (info)

Description

The circuit information analysis outputs several kinds of information about the circuit and its components. You can use various filters to specify what information is output. You can create a listing of model, instance, temperature-dependent, input, output, and operating point parameters. You can also generate a summary of the minimum and maximum parameter values (by using `extremes=yes` or `only`). Finally, you can request that Spectre provide a node-to-terminal map (by using `what=terminals`) or a terminal-to-node map (by using `what=nodes`).

The following are brief descriptions of the types of parameters you can request with the `info` statement:

Input parameters: Parameters that you specify in the netlist, such as the given length of a MOSFET or the saturation current of a bipolar transistor (use `what=inst, models, input, or all`)

Output parameters: Parameters that are computed by Spectre, such as temperature dependent parameters and the effective length of a MOSFET after scaling (use `what=output or all`)

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Analysis Statements

Operating-point parameters: Parameters that depend on the actual solution computed (use `what=oppoint`)

Definition

Name info parameter=value ...

Parameters

- | | | |
|---|------------------------------------|---|
| 1 | <code>what=oppoint</code> | What parameters should be printed.
Possible values are none, inst, models, input, output, nodes, all, terminals, oppoint, captab, or parameters. |
| 2 | <code>where=logfile</code> | Where parameters should be printed.
Possible values are nowhere, screen, file, logfile, or rawfile. |
| 3 | <code>file="%C:r.info.what"</code> | File name when <code>where=file</code> . |
| 4 | <code>save</code> | Signals to output.
Possible values are all, lvl, allpub, lvlpub, selected, or none. |
| 5 | <code>nestlvl</code> | Levels of subcircuits to output. |
| 6 | <code>extremes=yes</code> | Print minimum and maximum values.
Possible values are no, yes or only. |
| 7 | <code>title</code> | Analysis title. |

Captab parameters

- | | | |
|----|----------------------------|--|
| 8 | <code>detail=node</code> | How detailed should the capacitance table be.
Possible values are node, nodetoground or nodetonode. |
| 9 | <code>sort=name</code> | How to sort the capacitance table.
Possible values are name or value. |
| 10 | <code>threshold=0 F</code> | Threshold value for printing capacitances (ignore capacitances smaller than this value). |

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Analysis Statements

Parameter Index

In the following index, the number following each parameter name indicates where to find the description of that parameter.

detail	8	nestlvl	5	threshold	10	where	2
extremes	6	save	4	title	7		
file	3	sort	9	what	1		

Monte Carlo Analysis (montecarlo)

Description

The `montecarlo` analysis is a swept analysis with associated child analyses similar to the sweep analysis (see `spectre -h sweep`.) The Monte Carlo analysis refers to "statistics blocks" where statistical distributions and correlations of netlist parameters are specified. (Detailed information on statistics blocks is given below.) For each iteration of the Monte Carlo analysis, new pseudo-random values are generated for the specified netlist parameters (according to their specified distributions) and the list of child analyses are then executed.

Expressions are associated with the child analyses. These expressions, which are constructed as scalar calculator expressions by the user during Monte Carlo analysis set up, can be used to measure circuit metrics, such as the slew-rate of an op-amp. During a Monte Carlo analysis, these expression results will vary as the netlist parameters vary for each Monte Carlo iteration. The Monte Carlo analysis therefore becomes a tool that allows you to examine and predict circuit performance variations, which affect yield.

The statistics blocks allow you to specify batch-to-batch (process) and per-instance (mismatch) variations for netlist parameters. These statistically-varying netlist parameters can be referenced by models or instances in the main netlist and may represent IC manufacturing process variation, or component variations for board-level designs for example. The following description gives a simplified example of the Monte Carlo analysis flow:

```
perform nominal run if requested
```

```
if any errors in nominal run then stop
```

```
foreach Monte Carlo iteration {
```

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Analysis Statements

```
if process variations specified then
    apply process variation to parameters
if mismatch variations specified then
    foreach subcircuit instance {
        apply mismatch variation to parameters
    }
foreach child analysis {
    run child analysis
    evaluate expressions
}
}
```

Definition

Name `montecarlo parameter=value ...`

Parameters

Analysis parameters

- | | | |
|---|--------------------------|--|
| 1 | <code>numruns=100</code> | Number of Monte Carlo iterations to perform (not including nominal). |
| 2 | <code>seed</code> | Optional starting seed for random number generator. |
| 3 | <code>scalarfile</code> | Output file that will contain output scalar data. |
| 4 | <code>paramfile</code> | Output file that will contain output scalar data labels. |

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Analysis Statements

Saving Process Parameters

- 5 `saveprocessparams` Whether or not to save scalar data for statistically varying process parameters which are subject to process variation. Possible values are `no` or `yes`.
- 6 `processscalarfile` Output file that will contain process parameter scalar data.
- 7 `processparamfile` Output file that will contain process parameter scalar data labels.
- 8 `saveprocessvec=[...]` Array of statistically varying process parameters (which are subject to process variation) to save as scalar data in `processscalarfile`.
- 9 `firstrun=1` Starting iteration number.
- 10 `variations=process` Level of statistical variation to apply. Possible values are `process`, `mismatch` or `all`.

Flags

- 11 `donominal=yes` Whether or not to perform nominal run. Possible values are `no` or `yes`.
- 12 `appendsd=no` Whether or not to append scalar data. Possible values are `no` or `yes`.
- 13 `savefamilyplots=no` Whether or not to save data for family plots. If yes, this could require a lot of disk space. Possible values are `no` or `yes`.
- 14 `saveprocessparams` Whether or not to save scalar data for statistically varying process parameters which are subject to process variation. Possible values are `no` or `yes`.

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Analysis Statements

Annotation parameters

- 15 `annotate=sweep` Degree of annotation.
Possible values are `no`, `title`, `sweep`, or `status`.
- 16 `title` Analysis title.

Detailed Description and Examples

`numruns:(default=100)`

The number of Monte Carlo iterations to perform. The simulator will perform a loop, running the specified child analyses and evaluating any expressions `numruns` times.

`seed:(no default)`

`seed` for the random number generator. By always specifying the same seed, you can reproduce a previous experiment. If you do not specify a seed, then each time you run the analysis, you will get different results i.e. a different stream of pseudo-random numbers will be generated.

`scalarfile="filename"`

This parameter allows an ASCII file to be specified in which scalar data (results of expressions that resolve to scalar values) will be written. The data from this file can be read and plotted in histograms by Artist. For each iteration of each Monte Carlo child analyses, Spectre (through Artil) will write a line to this ASCII file which contains scalar data (one scalar expression per column e.g. `slewrate` or `bandwidth`.) The default name for this file will be of the form `name.mcddata`, where `name` is the name of the Monte Carlo analysis instance. This file contains only the matrix of numeric values. Artist Monte Carlo users will be more familiar with the term `mcddata` file for the scalar file. Additionally, when the Analog Artist Monte Carlo tool is used to generate the spectre netlist file, Spectre will merge the values of the statistically varying process parameters into this file containing the scalar data (results of expressions). This means that Analog Artist can later read the data, and create scatterplots of the statistically varying process parameters against each other, or against the results of the expressions. In this way, the user can see correlations between process parameter variations and circuit performances variations. This data merging will take place whenever the `scalarfile` and `processscalarfile` (see below) are written in the same directory.

`paramfile="filename"`

This file contains the titles, sweep variable values and the full expression for each of the columns in the `scalarfile`. Artist Monte Carlo users will be more familiar with the term

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`mcpparam` file for the `paramfile`. This file will be created in the `psf` directory by default, unless you specify some path information in the filename.

`processscalarfile="filename"`

If `saveprocessparams` is set to `yes`, then the process (batch-to-batch) values of all statistically varying parameters are saved to this scalar data file. You can use the `saveprocessvec` to filter out a subset of parameters in which case Spectre will save only the parameters specified in `saveprocessvec` to the `processscalarfile`.) The `processscalarfile` is equivalent to the `scalarfile`, except that the data in the `scalarfile` contains the values of the scalar expressions, whereas the data in the `processscalarfile` contains the corresponding process parameter values. The default name for this file will be of the form `instname.process.mcddata`, where `instname` is the name of the Monte Carlo analysis instance. This file will be created in the `psf` directory by default, unless you specify some path information in the filename. You can load the `processscalar` file and `processparamfile` into the Artist statistical postprocessing environment to plot/verify the process parameter distributions. If you later merge the `processparamfile` with the data in the `scalarfile`, you can then plot scalar expressions values against the corresponding process parameters by loading this merged file into the Artist statistical postprocessing environment.

`processparamfile="filename"`

This file contains the titles, sweep variable values for each of the columns in the `processscalarfile`. These titles will be the names of the process parameters.

The `processparamfile` is equivalent to the `paramfile`, except that the `paramfile` contains the name of the expressions, whereas the `processparamfile` contains the names of the process parameters. The default name for this file will be of the form `instname.process.mcpparam`, where `instname` is the name of the Monte Carlo analysis instance. This file will be created in the `psf` directory by default, unless you specify some path information in the filename.

`firstrun:(default=1)`

index of first iteration. If the first iteration is specified as some number `n` greater than one, then the beginning `n-1` iterations are `skipped` i.e. the Monte Carlo analysis behaves as if the first `n-1` iterations were run, but without actually performing the child analyses for these iterations. The subsequent stream of random numbers generated for the remaining iterations will be the same as if the first `n-1` iterations were actually run. By specifying the first iteration number and the same value for `seed`, you can reproduce a particular run or sequence of runs from a previous experiment (for example to examine an outlier case in more detail.)

`variations={process,mismatch,all}` (defaults to `process`).

Whether to apply process (batch-to-batch) variations only, or `mismatch` (per-instance) variations only, or both together. This assumes that you have specified appropriate statistical

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distributions in the statistics block. You cannot request that mismatch variations be applied unless you have specified mismatch statistics in the statistics block. You cannot request that process variations be applied unless you have specified process statistics in the statistics block. More details on statistics blocks are given below.

`saveprocessvec=[rshsp TOX ...]`

If `saveprocessparams` is specified as `yes`, then save the process (batch-to-batch) values of only those parameters listed in `saveprocessvec` in the `processparamfile`. This acts as a filter so that you do not save all process parameters to the file. If you do not want to filter the list of process parameters, then do not specify this parameter.

`donominal={yes,no}`(defaults to `yes`).

This parameter controls whether or not Spectre should perform a nominal run before starting the main Monte Carlo loop of iterations. If any errors are encountered during the nominal run (e.g. convergence problems, incorrect expressions, etc.) then Spectre will issue an appropriate error message and immediately abandon the Monte Carlo analysis.

If `donominal` is set to `no` then Spectre will run the Monte Carlo iterations only, and will not perform a nominal analysis. If any errors are encountered during the Monte Carlo iterations, Spectre will issue a warning and continue with the next iteration of the Monte Carlo loop.

`appendsd={yes,no}`(defaults to `no`).

Specifies whether to append scalar data to an existing scalarfile, or to overwrite the existing scalarfile. This flag applies to both the scalar file and the `processscalarfile`.

`savefamilyplots={yes,no}`.

If `yes`, a data file (e.g. `psf`) is saved for each analysis for each Monte Carlo iteration, in addition to the expressions scalar results which are saved to the ASCII scalar data file at the end of each iteration. Saving the full data files between runs enables the cloud plotting feature (overlaid waveforms) in Artist. It also enables the user to define/evaluate new calculator measurements after the simulation has been run using the Artist calculator. This feature could result in a huge amount of data being stored to disk, and it is advised that you use this feature with care. If you do decide to use this feature, it is advisable to keep the number of saved quantities to a minimum. If this parameter is set to `no`, then data files are overwritten by each Monte Carlo iteration.

`annotate={no,title,sweep,status}`

Degree of annotation. Use the maximum value of `status` to print a summary of which runs did not converge or had problems evaluating expressions, etc.

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Examples:

```
// do a Monte Carlo analysis, with process variations only
// useful for looking at absolute performance spreads
mc1 montecarlo variations=process seed=1234 numruns=200 {
  dcop1 dc    // a child analysis
  tran1 tran start=0 stop=1u // another child analysis
  // expression calculations are sent to the scalardata file
  export slewrate=oceanEval("slewRate(v(\"vout\"),10n,t,30n,t,10,90 )")
}

// do a Monte Carlo analysis, with mismatch variations only
// useful for detecting spreads in differential circuit
// applications, etc. Do not perform a nominal run.
mc2 montecarlo donominal=no variations=mismatch seed=1234 numruns=200 {
  dcop2 dc
  tran2 tran start=0 stop=1u
  export slewrate=oceanEval("slewRate(v(\"vout\"),10n,t,30n,t,10,90 )")
}

// do both together...
mc3 montecarlo saveprocessparams=yes variations=all numruns=200 {
  dcop3 dc
  tran3 tran start=0 stop=1u
  export slewrate=oceanEval("slewRate(v(\"vout\"),10n,t,30n,t,10,90 )")
}
```

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Specifying Parameter Distributions using Statistics Blocks

The `statistics` blocks are used to specify the input statistical variations for a Monte Carlo analysis. A `statistics` block may contain one or more `process` blocks (which represents batch-to-batch type variations), and/or one or more `mismatch` blocks (which represents on-chip or device mismatch variations), in which the distributions for parameters are specified. `Statistics` blocks may also contain one or more correlation statements to specify the correlations between specified process parameters, and/or to specify correlated device instances (for example matched pairs). `Statistics` blocks may also contain a `truncate` statement which may be used for generating truncated distributions. The distributions specified in the `process` block will be sampled once per Monte Carlo iteration, and are typically used to represent batch-to-batch, or process variations, whereas the distributions specified in the `mismatch` block are sampled on a per subcircuit instance basis and are typically used to represent device-to-device mismatch for devices on the same chip. In the case where the same parameter is subject to both process and mismatch variations, then the sampled process value becomes the mean for the mismatch random number generator for that particular parameter.

NOTE: Multiple `statistics` blocks may exist, in which case they accumulate or overlay. Typically, process variations, mismatch variations and correlations between process parameters will be specified in one `statistics` block. A second `statistics` block would be specified where actual device instance correlations are specified (i.e. specification of matched pairs).

`Statistics` blocks can be specified using combinations of the Spectre keywords `statistics`, `process`, `mismatch`, `vary`, `truncate` and `correlate`. Braces `{}` are used to delimit blocks.

The following example shows some sample `statistics` blocks, which are discussed below along with syntax requirements.

```
// define some netlist parameters to represent process parameters
// such as sheet resistance and mismatch factors
parameters rshsp=200 rshpi=5k rshpi_std=0.4K xisn=1 xisp=1 xxx=20000 uuu=200
// define statistical variations, to be used
// with a MonteCarlo analysis.
statistics {
    process { // process: generate random number once per MC run
        vary rshsp dist=gauss std=12 percent=yes
```

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```
vary rshpi dist=gauss std=rshpi_std // rshpi_std is a parameter
vary xxx dist=lnorm std=12
vary uuu dist=unif N=10 percent=yes
...
}
mismatch { // mismatch: generate a random number per instance
    vary rshsp dist=gauss std=2
    vary xisn dist=gauss std=0.5
    vary xisp dist=gauss std=0.5
}
// some process parameters are correlated
correlate param=[rshsp rshpi] cc=0.6
// specify a global distribution truncation factor
truncate tr=6.0 // +/- 6 sigma
}
// a separate statistics block to specify correlated (i.e. matched) components
// where m1 and m2 are subckt instances.
statistics {
    correlate dev=[m1 m2] param=[xisn xisp] cc=0.8
}
}
```

Specifying Distributions:

Parameter variations are specified using the following syntax:

```
vary PAR_NAME dist=<type> {std=<value> | N=<value>} {percent=yes|no}
```

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Three types of parameter distributions are available: gaussian, lognormal and uniform, corresponding to the <type> keywords `gauss`, `lnorm` and `unif` respectively. For both the `gauss` and the `lnorm` distributions, you specify a standard deviation using the `std` keyword.

Gaussian Distribution:

For the gaussian distribution, the mean value is taken as the current value of the parameter being varied, giving a distribution denoted by Normal (mean,std). Using the example above, parameter `rshpi` is varied with a distribution of Normal (5k,0.4k)

Lognormal Distribution:

The lognormal distribution is denoted by

$$\log(x) = \text{Normal}(\log(\text{mean}), \text{std})$$

where `x` is the parameter being specified as having a lognormal distribution.

(NOTE: `log()` is the natural logarithm function.) For parameter `xxx` in the example above, the process variation is according to

$$\log(\text{xxx}) = \text{Normal}(\log(20000), 12)$$

Uniform Distribution:

The uniform distribution for parameter `x` is generated according to

$$x = \text{unif}(\text{mean}-N, \text{mean}+N)$$

such that the mean value is the nominal value of the parameter `x`, and the parameter is varied about the mean with a range of +/- `N`. The standard deviation is not specified for the uniform distribution, but its value can be calculated from the formula: $\text{std}=N/\text{sqrt}(3)$.

Values as percentages:

The `percent` flag indicates whether the standard deviation `std` or uniform range `N` are specified in absolute terms (`percent=no`) or as a percentage of the mean value (`percent=yes`). For parameter `uuu` in the example above, the mean value is 200, and the variation is $200 \pm 10\% \cdot (200)$ i.e. 200 ± 20 . For parameter `rshsp`, the process variation is given by Normal (200, $12\% \cdot (200)$) i.e. Normal (200, 24). It is not advised that you use `percent=yes` with the lognormal distribution.

Process and Mismatch Variations:

The statistics specified in a process block are applied at global scope, and the distributions are sampled once per Monte Carlo iteration. The statistics specified in a mismatch block are

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applied on a per-subcircuit instance basis, and are sampled once per subcircuit instance. If you place model cards and/or device instances in subcircuits, and add a mismatch block to your statistics block you can effectively model device-to-device mismatch for these devices/models.

Correlation Statements:

There are two types of correlation statements that you can use: process parameter correlation statements, and instance correlation statements.

Process Parameter Correlation:

The syntax of the process parameter correlation statement is:

```
correlate param=[list of parameters] cc=<value>
```

This allows you to specify a correlation coefficient between multiple process parameters. You can specify multiple process parameter correlation statements in a statistics block, to build a matrix of process parameter correlations. During a Monte Carlo analysis, process parameter values will be randomly generated according to the specified distributions and correlations.

Mismatch Correlation (Matched Devices):

The syntax of the instance or mismatch correlation statement is:

```
correlate dev=[list of subcircuit instances] {param=[list of parameters]} cc=<value>
```

where the device or subcircuit instances to be matched are listed in the list of subcircuit instances, and the list of parameters specifies exactly which parameters with mismatch variations are to be correlated.

The instance mismatch correlation statement is used to specify correlations for particular subcircuit instances. If a subcircuit contains a device, you can effectively use the instance correlation statements to specify that certain devices are correlated (i.e. matched) and give the correlation coefficient. You can optionally specify exactly which parameters are to be correlated by giving a list of parameters (each of which must have had distributions specified for it in a mismatch block), or specify no parameter list, in which case all parameters with mismatch statistics specified are correlated with the given correlation coefficient. The correlation coefficients are specified in the <value> field and must be between +/- 1.0, not including 1.0 or -1.0.

NOTE: correlation coefficients can be constants or expressions, as can `std` and `N` when specifying distributions.

Truncation Factor:

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The default truncation factor for gaussian distributions (and for the gaussian distribution underlying the lognormal distribution) is 4.0 sigma. Randomly generated values which are outside the range of mean +/- 4.0 sigma are automatically rejected and regenerated until they fall inside the range. You can change the truncation factor using the `truncate` statement. The syntax is:

```
truncate tr=<value>.
```

Note: The value of the truncation factor can be a constant or an expression.

Note: Parameter correlations can be affected by using small truncation factors.

Parameter Index

In the following index, the number following each parameter name indicates where to find the description of that parameter.

<code>annotate</code>	15	<code>numruns</code>	1	<code>savefamilyplots</code>	13	<code>scalarfile</code>	3
<code>appendsd</code>	12	<code>paramfile</code>	4	<code>saveprocessparams</code>	5	<code>seed</code>	2
<code>donominal</code>	11	<code>processparamfile</code>	7	<code>saveprocessparams</code>	14	<code>title</code>	16
<code>firstrun</code>	9	<code>processscalarfile</code>	6	<code>saveprocessvec</code>	8	<code>variations</code>	10

Noise Analysis (noise)

Description

The noise analysis linearizes the circuit about the operating point and computes the noise spectral density at the output. If you identify an input source, the transfer function and the input-referred noise for an equivalent noise-free network is computed. In addition, if the input source is noisy, then the noise figure is computed.

The noise is computed at the output of the circuit. The output is specified with either a pair of nodes or a probe component. To specify the output of a circuit with a probe, specify it with the `oprobe` parameter. If the output is voltage (or potential), choose a `resistor` or a `port` as the output probe. If the output is current (or flow), choose a `vsource` or `iprobe` as the output probe.

If the input-referred noise is desired, specify the input source using the `iprobe` parameter. Currently, only a `vsource`, an `isource`, or a `port` may be used as an input probe. If the

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input source is noisy, as is a `port`, the noise analysis will compute the noise factor (F) and noise figure (NF). To match the IEEE definition of noise figure, the input probe must be a port with no excess noise and its `noisetemp` must be set to 16.85C (290K). In addition, the output load must be a `resistor` or `port` and must be identified as the `oprobe`.

The noise analysis always computes the total noise at the output, which includes contributions from the input source and the output load. The amount of the output noise that is attributable to each noise source in the circuit is also computed and output individually. If the input source is identified, the input-referred noise is computed, which includes the noise from the input source itself. Finally, if the input source is identified and is noisy, the noise factor and noise figure are computed. Thus if

No = total output noise

Ns = noise at the output due to the input probe (the source)

Nl = noise at the output due to the output probe (the load)

IRN = input referred noise

G = gain of the circuit

F = noise factor

NF = noise figure

then,

$$\text{IRN} = \sqrt{\text{No}^2 / \text{G}^2}$$

$$\text{F} = (\text{No}^2 - \text{Nl}^2) / \text{Ns}^2$$

$$\text{NF} = 10 * \log_{10}(\text{F})$$

When the results are output, No is named `out`, IRN is named `in`, G is named `gain`, F is named `F`, and NF is named `NF`.

Spectre can perform the analysis while sweeping a parameter. The parameter can be frequency, temperature, component instance parameter, component model parameter, or netlist parameter. If changing a parameter affects the DC operating point, the operating point is recomputed on each step. You can sweep the circuit temperature by giving the parameter name as `temp` with no `dev` or `mod` parameter. You can sweep a netlist parameter by giving the parameter name with no `dev`, or `mod` parameter. After the analysis has completed, the modified parameter returns to its original value.

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Definition

Name [p] [n] noise parameter=value ...

The optional terminals (p and n) specify the output of the circuit. If you do not give the terminals, then you must specify the output with a probe component.

Parameters

1 `prevoppoint=no` Use operating point computed on the previous analysis.
Possible values are `no` or `yes`.

Sweep interval parameters

2 `start=0` Start sweep limit.
3 `stop` Stop sweep limit.
4 `center` Center of sweep.
5 `span=0` Sweep limit span.
6 `step` Step size, linear sweep.
7 `lin=50` Number of steps, linear sweep.
8 `dec` Points per decade.
9 `log=50` Number of steps, log sweep.
10 `values=[...]` Array of sweep values.

Sweep variable parameters

11 `dev` Device instance whose parameter value is to be swept.
12 `mod` Model whose parameter value is to be swept.
13 `param` Name of parameter to sweep.
14 `freq (Hz)` Frequency when parameter other than frequency is being swept.

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Probe parameters

- 15 `oprobe` Compute total noise at the output defined by this component.
- 16 `iprobe` Input probe. Refer the output noise to this component.

State-file parameters

- 17 `readns` File that contains estimate of DC solution (nodeset).

Output parameters

- 18 `save` Signals to output.
Possible values are `all`, `lvl`, `allpub`, `lvlpub`, `selected`, or `none`.
- 19 `nestlvl` Levels of subcircuits to output.
- 20 `oppoint=no` Should operating point information be computed, and if so, where should it be sent. Operating point information would not be output if operating point is computed in the previous analysis and is unchanged.
Possible values are `no`, `screen`, `logfile`, or `rawfile`.

Convergence parameters

- 21 `restart=yes` Restart the DC solution from scratch if any condition has changed. If not, use the previous solution as initial guess.
Possible values are `no` or `yes`.

Annotation parameters

- 22 `annotate=sweep` Degree of annotation.
Possible values are `no`, `title`, `sweep`, `status`, or `steps`.
- 23 `stats=no` Analysis statistics.
Possible values are `no` or `yes`.
- 24 `title` Analysis title.

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You can specify sweep limits by giving the end points or by providing the center value and the span of the sweep. Steps can be linear or logarithmic, and you can specify the number of steps or the size of each step. You can give a step size parameter (`step`, `lin`, `log`, `dec`) to determine whether the sweep is linear or logarithmic. If you do not give a step size parameter, the sweep is linear when the ratio of stop to start values is less than 10, and logarithmic when this ratio is 10 or greater. All frequencies are in Hertz.

The small-signal analysis begins by linearizing the circuit about an operating-point. By default this analysis computes the operating-point if it is not known, or recomputes it if any significant component or circuit parameter has changed. However, if a previous analysis computed an operating point, you can set `prevoppoint=yes` to avoid recomputing it. For example, if you use this option when the previous analysis was a transient analysis, the operating point is the state of the circuit on the final time point.

Parameter Index

In the following index, the number following each parameter name indicates where to find the description of that parameter.

<code>annotate</code>	22	<code>lin</code>	7	<code>param</code>	13	<code>start</code>	2
<code>center</code>	4	<code>log</code>	9	<code>prevoppoint</code>	1	<code>stats</code>	23
<code>dec</code>	8	<code>mod</code>	12	<code>readns</code>	17	<code>step</code>	6
<code>dev</code>	11	<code>nestlvl</code>	19	<code>restart</code>	21	<code>stop</code>	3
<code>freq</code>	14	<code>oppoint</code>	20	<code>save</code>	18	<code>title</code>	24
<code>iprobe</code>	16	<code>oprobe</code>	15	<code>span</code>	5	<code>values</code>	10

Immediate Set Options (options)

Description

The immediate set options statement sets or changes various program control options. These options take effect immediately and are set while the circuit is read. For further options, see the individual analyses.

NOTE: Options that are dependent on netlist parameter values, do not maintain their dependencies on those netlist parameters.

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Definition

Name options parameter=value ...

Parameters

Tolerance parameters

- | | | |
|---|-----------------|---|
| 1 | reltol=0.001 | Relative convergence criterion. |
| 2 | vabstol=1e-06 V | Voltage absolute tolerance convergence criterion. |
| 3 | iabstol=1e-12 A | Current absolute tolerance convergence criterion. |

Temperature parameters

- | | | |
|---|-----------------|--|
| 4 | temp=27 C | Temperature. |
| 5 | tnom=27 C | Default component parameter measurement temperature. |
| 6 | tempeffects=all | Temperature effect selector. If tempeffect = vt, only thermal voltage varies with temperature; if tempeffect = tc, parameters that start with tc are active and thermal voltage is dependent on temperature; and if tempeffect = all, all built-in temperature models are enabled.
Possible values are vt, tc or all. |

Output parameters

- | | | |
|----|-------------------|--|
| 7 | save=selected | Signals to output.
Possible values are all, lvl, allpub, lvlpub, selected, or none. |
| 8 | nestlvl= ∞ | Levels of subcircuits to output. |
| 9 | subcktprobelvl=0 | Level up to which subcircuit terminal current probes are to be set up. |
| 10 | currents=selected | Terminal currents to output. (See important note below about |

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- saving currents by using probes).
Possible values are `all`, `nonlinear` or `selected`.
- 11 `useprobes=no` Use current probes when measuring terminal currents. (See important note below about saving currents by using probes).
Possible values are `no` or `yes`.
- 12 `useterms=index` Output terminal currents by specified option.
Possible values are `name` or `index`.
- 13 `redundant_currents=no` If yes, save both currents through two terminal devices.
Possible values are `no` or `yes`.
- 14 `pwr=none` Power signals to create.
Possible values are `all`, `subckts`, `devices`, `total`, or `none`.
- 15 `saveahdlvars=selected` AHDL variables to output.
Possible values are `all` or `selected`.
- 16 `rawfmt=psfbin` Output raw data file format.
Possible values are `nutbin`, `nutascii`, `wsfbin`, `wsfascii`, `psfbin`, `psfascii`, `psfbinf`, `awb`, or `sst2`.
- 17 `rawfile="%C:r.raw"` Output raw data file name.

Convergence parameters

- 18 `homotopy=all` Method used when no convergence on initial attempt of DC analysis.
Possible values are `none`, `gmin`, `source`, `dptran`, `ptran`, `arclength`, or `all`.
- 19 `limit=dev` Limiting algorithms to aid DC convergence.
Possible values are `delta`, `log` or `dev`.

Multithreading parameters

- 20 `multithread=off` This option turns on/off multithread capability. When multithreading is turned on but the number of threads (`nThreads`)

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is not specified, Spectre will automatically detect the number of processors and select the proper number of threads to use. (See important note below about using multithreading). Possible values are `off` or `on`.

21 `nthreads` Specifies the number of threads for multithreading.

Component parameters

22 `scalem=1` Model scaling factor.

23 `scale=1` Device instance scaling factor.

24 `compatible=spectre`
Encourage device equations to be compatible with a foreign simulator. This option does not affect input syntax. Possible values are `spectre`, `spice2`, `spice3`, `cdsspice`, `hspice`, or `spiceplus`.

25 `approx=no` Use approximate models. Difference between approximate and exact models is generally very small. Possible values are `no` or `yes`.

26 `macromodels=no` Circuit contains macromodels; sometimes helps performance. Possible values are `no` or `yes`.

27 `mos_method=s` Global Mos Table Model selection flag. Possible values are `s` or `a`.

28 `maxrsd=0.0 Ω` Use approximation for drain/source parasitic resistors which are less than `maxrsd`. Applies to `bsim3v3`, `bsim4` mosfet models.

Error-checking parameters

29 `topcheck=full` Check circuit topology for errors. Possible values are `no`, `min`, `full`, or `fixall`.

30 `ignshorts=no` Silently ignore shorted components. Possible values are `no` or `yes`.

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- 31 `diagnose=no` Print additional information that might help diagnose accuracy and convergence problems.
Possible values are `no` or `yes`.
- 32 `checklimitfile` File to which assert violations will be written to.
- 33 `dochecklimit=yes` Check asserts in the netlist.
Possible values are `no` or `yes`.
- 34 `opptcheck=yes` Check operating point parameters against soft limits.
Possible values are `no` or `yes`.

Resistance parameters

- 35 `gmin=1e-12 S` Minimum conductance across each nonlinear device.
- 36 `gmin_check=max_v_only`
Specifies that effect of `gmin` should be reported if significant.
Possible values are `no`, `max_v_only`, `max_only`, or `all`.
- 37 `rforce=1 Ω` Resistance used when forcing nodesets and node-based initial conditions.

Quantity parameters

- 38 `value="V"` Default value quantity.
- 39 `flow="I"` Default flow quantity.
- 40 `quantities=no` Print quantities.
Possible values are `no`, `min` or `full`.

Annotation parameters

- 41 `audit=detailed` Print time required by various parts of the simulator.
Possible values are `no`, `brief`, `detailed`, or `full`.
- 42 `inventory=detailed`
Print summary of components used.
Possible values are `no`, `brief` or `detailed`.

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43	<code>narrate=yes</code>	Narrate the simulation. Possible values are <code>no</code> or <code>yes</code> .
44	<code>debug=no</code>	Give debugging messages. Possible values are <code>no</code> or <code>yes</code> .
45	<code>info=yes</code>	Give informational messages. Possible values are <code>no</code> or <code>yes</code> .
46	<code>note=yes</code>	Give notice messages. Possible values are <code>no</code> or <code>yes</code> .
47	<code>maxnotes=5</code>	Maximum number of times any notice will be issued per analysis.
48	<code>warn=yes</code>	Give warning messages. Possible values are <code>no</code> or <code>yes</code> .
49	<code>maxwarns=5</code>	Maximum number of times any warning message will be issued per analysis.
50	<code>maxwarnstologfile=5</code>	Maximum number of times any warning message will be printed to the log file per analysis.
51	<code>maxnotestologfile=5</code>	Maximum number of times any notice message will be printed to the log file per analysis.
52	<code>error=yes</code>	Give error messages. Possible values are <code>no</code> or <code>yes</code> .
53	<code>digits=5</code>	Number of digits used when printing numbers.
54	<code>notation=eng</code>	When printing real numbers to the screen, what notation should be used. Possible values are <code>eng</code> , <code>sci</code> or <code>float</code> .
55	<code>cols=80</code>	Width of screen in characters.
56	<code>title</code>	Circuit title.

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Matrix parameters

- 57 `pivotdc=no` Use numeric pivoting on every iteration of DC analysis.
Possible values are `no` or `yes`.
- 58 `pivrel=0.001` Relative pivot threshold.
- 59 `pivabs=0` Absolute pivot threshold.

Miscellaneous parameters

- 60 `ckptclock=1800 s` Clock time checkpoint period.

Sensitivity parameters

- 61 `sensfile` Output sensitivity data file name.
- 62 `sensformat=tabular`
 Format of sensitivity data.
Possible values are `tabular` or `list`.
- 63 `senstype=partial` Type of sensitivity being calculated.
Possible values are `partial` or `normalized`.

Important note about `currents` and `useprobes` options

Adding probes to circuits that are sensitive to numerical noise might affect the solution. In such cases accurate solution may be obtained by reducing `reltol`.

The following devices will always use probes to save currents (even with `useprobes=no`): `port`, `delay`, `switch`, `hbt`, `transformer`, `core`, `winding`, `fourier`, `d2a`, `a2d`, `a2ao`, `a2ai`.

`senstype` parameter

When `senstype` is set to `partial`, the sensitivity being calculated is the partial derivative of a differentiable output variable F with respect to a design parameter p :

$$D(F \text{ w.r.t. } p) = \frac{dF}{dp}$$

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Analysis Statements

This definition is not scale free. When `senstype` is set to `normalized`, the sensitivity being calculated is the normalized sensitivity

$$S(F \text{ w.r.t. } p) = \frac{d \ln F}{d \ln p} = \frac{p}{F} \frac{dF}{dp} = -D(F \text{ w.r.t. } p)$$

When either F or p takes a zero value, the above normalized definition no longer provides a useful measure, the following two seminormalized sensitivities are used instead:

$$S(F \text{ w.r.t. } p) = \frac{dF}{d \ln p} = p \frac{dF}{dp} = p D(F \text{ w.r.t. } p) \quad \text{if } F = 0$$

and

$$S(F \text{ w.r.t. } p) = \frac{d \ln F}{dp} = \frac{1}{F} \frac{dF}{dp} = -D(F \text{ w.r.t. } p) \quad \text{if } p = 0$$

When both F and p are zero, the partial sensitivity is used.

`topcheck` parameter:

When `topcheck=full`, the topology check is performed and `gmin` is inserted between isolated nodes and ground. A heuristic topology check is also performed to find nodes that may be isolated due to the numerical nature of the circuit. For example, nodes isolated by reverse biased diodes in MOSFETS.

Use `topcheck=fixall` to attach `gmin` to all types of isolated nodes. Including the ones found by the heuristic topology check.

Important note about using multithreading

Currently, multithreading is only available for devices evaluation for BSIM3v3, and BSIM4. Multithreading does not work with table model. If there is an instance of a primitive using table model, multithreading would not be applied to all instances of that primitive.

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Analysis Statements

Multithreading can be turned on/off by command line option, or by the multithread parameter in the options statement from the input file. If both options are specified, command line option will make the final determination on the number of threads to use.

Using multithreading on circuits that are sensitive to numerical noise might affect the solution. The solution should still be within acceptable tolerance specified by the tolerance parameters in the Spectre input file. Due to the order of evaluation of devices is different for each multithreading run of the same simulation, this could lead to different round off error in the computation. It is possible that same exact result may not be reproducible when multithreading is used.

Multithreading would work best when the following capabilities are not used: useprobes=yes, save-current/SOA/alarm for multithreaded devices.

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Periodic AC Analysis (pac)

Description

The periodic AC (PAC) analysis is used to compute transfer functions for circuits that exhibit frequency translation. Such circuits include mixers, switched-capacitor filters, samplers, phase-locked loops, and the like. It is a small-signal analysis like AC analysis, except the circuit is first linearized about a periodically varying operating point as opposed to a simple DC operating point. Linearizing about a periodically time-varying operating point allows transfer-functions that include frequency translation, whereas simply linearizing about a DC operating point could not because linear time-invariant circuits do not exhibit frequency translation. Also, the frequency of the sinusoidal stimulus is not constrained by the period of the large periodic solution.

Computing the small-signal response of a periodically varying circuit is a two step process. First, the small stimulus is ignored and the periodic steady-state response of the circuit to possibly large periodic stimulus is computed using PSS analysis. As a normal part of the PSS analysis, the periodically time-varying representation of the circuit is computed and saved for later use. The second step is applying the small stimulus to the periodically varying linear representation to compute the small signal response. This is done using the PAC analysis. A PAC analysis cannot be used alone, it must follow a PSS analysis. However, any number of periodic small-signal analyses such as PAC, PSP, PXF, PNoise, can follow a PSS analysis.

Unlike other analyses in Spectre, this analysis can only sweep frequency.

Definition

```
Name pac parameter=value ...
```

Parameters

Sweep interval parameters

- | | | |
|---|----------------------|--------------------|
| 1 | <code>start=0</code> | Start sweep limit. |
| 2 | <code>stop</code> | Stop sweep limit. |
| 3 | <code>center</code> | Center of sweep. |
| 4 | <code>span=0</code> | Sweep limit span. |

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5	<code>step</code>	Step size, linear sweep.
6	<code>lin=50</code>	Number of steps, linear sweep.
7	<code>dec</code>	Points per decade.
8	<code>log=50</code>	Number of steps, log sweep.
9	<code>values=[...]</code>	Array of sweep values.
10	<code>sweepstype</code>	Specifies if the sweep frequency range is absolute frequency of input or if it is relative to the port harmonics. Possible values are <code>absolute</code> or <code>relative</code> .
11	<code>relharmnum=1</code>	Harmonic to which relative frequency sweep should be referenced.

Output parameters

12	<code>sidebands=[...]</code>	Array of relevant sidebands for the analysis.
13	<code>maxsideband=0</code>	An alternative to the <code>sidebands</code> array specification, which automatically generates the array: [-maxsideband ... 0 ... +maxsideband].
14	<code>freqaxis</code>	Specifies whether the results should be output versus the input frequency, the output frequency, or the absolute value of the output frequency. Default is <code>in</code> for logarithmic frequency sweeps and <code>absout</code> otherwise. Possible values are <code>absout</code> , <code>out</code> or <code>in</code> .
15	<code>save</code>	Signals to output. Possible values are <code>all</code> , <code>lvl</code> , <code>allpub</code> , <code>lvlpub</code> , <code>selected</code> , or <code>none</code> .
16	<code>nestlvl</code>	Levels of subcircuits to output.
17	<code>outputperiod=0.0</code>	(no output) Time-domain output period. The time-domain small-signal response is computed for the period specified, rounded to the nearest integer multiple of the <code>pss</code> period.

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Analysis Statements

Convergence parameters

- 18 `tolerance=1e-9` Relative tolerance for linear solver.
- 19 `gear_order=2` Gear order used for small-signal integration.
- 20 `solver=turbo` Solver type.
Possible values are `std` or `turbo`.
- 21 `oscsolver=turbo` Oscillator solver type.
Possible values are `std` or `turbo`.

Annotation parameters

- 22 `annotate=sweep` Degree of annotation.
Possible values are `no`, `title`, `sweep`, `status`, or `steps`.
- 23 `stats=no` Analysis statistics.
Possible values are `no` or `yes`.
- 24 `title` Analysis title.

Modulation conversion parameters

- 25 `modulated=no` Compute transfer functions/conversion between modulated sources and outputs.
Possible values are `single`, `first`, `second`, or `no`.
- 26 `inmodharmnum=1` Harmonic for the PAC input source modulation.
- 27 `outmodharmvec=[...]` Harmonic list for the PAC output modulations.
- 28 `moduppersideband=1` Index of the upper sideband included in the modulation of an output for PAC or an input for PXF.
- 29 `modsource` Refer the output noise to this component.

You can select the set of periodic small-signal output frequencies of interest by setting either the `maxsideband` or the `sidebands` parameters. For a given set of n integer numbers representing the sidebands K_1, K_2, \dots, K_n , the output frequency at each sideband is computed as $f(\text{out}) = f(\text{in}) + K_i * f_{\text{fund}}(\text{pss})$, where $f(\text{in})$ represent the (possibly swept) input frequency, and

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`fund(pss)` represents the fundamental frequency used in the corresponding PSS analysis. Thus, when analyzing a down-converting mixer, while sweeping the RF input frequency, the most relevant sideband for IF output is $K_i = -1$. When simulating an up-converting mixer, while sweeping IF input frequency, the most relevant sideband for RF output is $K_i = 1$. By setting the `maxsideband` value to K_{max} , all $2 * K_{max} + 1$ sidebands from $-K_{max}$ to $+K_{max}$ are generated.

The number of requested sidebands does not change substantially the simulation time. However, the `maxacfreq` of the corresponding PSS analysis should be set to guarantee that $|\max\{f(\text{out})\}|$ is less than `maxacfreq`, otherwise the computed solution might be contaminated by aliasing effects. The PAC simulation is not executed for $|f(\text{in})|$ greater than `maxacfreq`. Diagnostic messages are printed for those extreme cases, indicating how `maxacfreq` should be set in the PSS analysis. In the majority of the simulations, however, this is not an issue, because `maxacfreq` is never allowed to be smaller than 40x the PSS fundamental.

With PAC the frequency of the stimulus and of the response are usually different (this is an important way in which PAC differs from AC). The `freqaxis` parameter is used to specify whether the results should be output versus the input frequency (`in`), the output frequency (`out`), or the absolute value of the output frequency (`absout`).

Modulated small signal measurements are possible using the Analog Artist(ADE) environment. The `modulated` option for PAC and other modulated parameters are set by Artist. PAC analyses with this option will produce results which could have limited use outside such environment. Direct Plot is configured to analyze these results and combine several wave forms to measure AM and PM response due to single sideband or modulated stimuli. For details, please see the SpectreRF User Guide.

Unlike AC analysis, PAC analysis can output the time-domain simulation results, by specifying the `outputperiod` parameter.

You can specify sweep limits by giving the end points or by providing the center value and the span of the sweep. Steps can be linear or logarithmic, and you can specify the number of steps or the size of each step. You can give a step size parameter (`step`, `lin`, `log`, `dec`) to determine whether the sweep is linear or logarithmic. If you do not give a step size parameter, the sweep is linear when the ratio of stop to start values is less than 10, and logarithmic when this ratio is 10 or greater. Alternatively, you may specify the particular values that the sweep parameter should take using the `values` parameter. If you give both a specific set of values and a set specified using a sweep range, the two sets are merged and collated before being used. All frequencies are in Hertz.

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Analysis Statements

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In the following index, the number following each parameter name indicates where to find the description of that parameter.

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Periodic Distortion Analysis (pdisto)

Description

This analysis computes the quasi-periodic steady-state (QPSS) response of a circuit that operates on multiple time scales. A quasi-periodic signal has dynamics in multiple fundamental frequencies. Closely spaced or incommensurate fundamentals cannot be resolved by PSS efficiently. QPSS allows you to compute a circuit responses to several moderately large input signals in addition to a strongly nonlinear tone which represents a LO or clock signal. A typical example is the intermodulation distortion measurements of a mixer with two closely spaced moderate input signals. QPSS treats one particular input signal (usually the one that causes the most nonlinearity or the largest response) as the large signal, and the others as moderate signals.

An initial transient analysis is carried out by first suppressing all moderate input signals. Then, a number of (at least 2) stabilizing iterations with all signals activated is run. They are followed by the shooting Newton method. QPSS employs the Mixed Frequency Time (MFT) algorithm extended to multiple fundamental frequencies. For details of MFT algorithm, please see *Steady-State Methods for Simulating Analog and Microwave Circuits*, by K. S. Kundert, J. K. White, and A. Sangiovanni-Vincentelli, Kluwer, Boston, 1990.

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Analysis Statements

Like PSS, QPSS uses the shooting Newton method as its backbone. However, instead of doing a single transient integration, each Newton iteration does a number of transient integrations of one large signal period.

Each of the integrations differs by a phase-shift in each moderate input signal. The number of integrations is determined by the numbers of harmonics of moderate fundamentals specified by the user. Given `maxharms=[k1 k2 ... kn]`, the total number of integrations is $(2*k2+1)*(2*k3+1)*...*(2*kn+1)$. As one consequence, the efficiency of the algorithm depends significantly on the number of harmonics required to model the responses of moderate fundamentals. As another consequence, the number of harmonics of the large fundamental does not significantly affect the efficiency of the shooting algorithm. The boundary conditions of a shooting interval are such that the time domain integrations are consistent with a frequency domain transformation with a shift of one large signal period.

QPSS inherits a majority of PSS parameters. A few new parameters are added. The most important ones are `funds` and `maxharms`. They replace PSS parameters, `fund` (or `period`) and `harms`, respectively. The `funds` parameter accepts a list of names of fundamentals that are present in the sources. These names are specified in the sources by parameter `fundname`. The first fundamental is considered as the large signal. A few heuristics can be used for picking the large fundamental.

- (1) Pick the one which is not a sinusoidal.
- (2) Pick the one which causes the most nonlinearity.
- (3) Pick the one which causes the largest response.

The `maxharms` parameter accepts a list of numbers of harmonics that are required to sufficiently model responses due to different fundamentals.

Definition

Name `pdisto parameter=value ...`

Parameters

Distortion fundamental parameters

- | | | |
|---|-----------------------------|--|
| 1 | <code>funds=[...]</code> | Array of fundamental frequency names for fundamentals to use in analysis. |
| 2 | <code>maxharms=[...]</code> | Array of number of harmonics of each fundamental to consider for each fundamental. |

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Analysis Statements

Simulation interval parameters

- 3 `tstab=0.0 s` Extra stabilization time after the onset of periodicity for independent sources.
- 4 `tstart=0.0 s` Initial transient analysis start time.

Time-step parameters

- 5 `maxstep (s)` Maximum time step. Default derived from `errpreset`.
- 6 `step=0.001 period s` Minimum time step that would be used solely to maintain the aesthetics of the results.

Initial-condition parameters

- 7 `ic=all` What should be used to set initial condition.
Possible values are `dc`, `node`, `dev`, or `all`.
- 8 `skipdc=no` If yes, there will be no dc analysis for transient.
Possible values are `no`, `yes` or `sigrampup`.
- 9 `readic` File that contains initial condition.

Convergence parameters

- 10 `readns` File that contains estimate of initial transient solution.
- 11 `cmin=0 F` Minimum capacitance from each node to ground.

Output parameters

- 12 `save` Signals to output.
Possible values are `all`, `lvl`, `allpub`, `lvlpub`, `selected`, or `none`.
- 13 `nestlvl` Levels of subcircuits to output.

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Analysis Statements

- 14 `oppoint=no` Should operating point information be computed for initial timestep, and if so, where should it be sent.
Possible values are `no`, `screen`, `logfile`, or `rawfile`.
- 15 `skipstart=starttime s` The time to start skipping output data.
- 16 `skipstop=stoptime s` The time to stop skipping output data.
- 17 `skipcount` Save only one of every `skipcount` points.
- 18 `strobeperiod (s)` The output strobe interval (in seconds of transient time).
- 19 `strobedelay=0 s` The delay (phase shift) between the `skipstart` time and the first strobe point.
- 20 `compression=no` Do data compression on output. See full description below.
Possible values are `no` or `yes`.
- 21 `saveinit=no` If set, the waveforms for the initial transient before steady state are saved.
Possible values are `no` or `yes`.

State-file parameters

- 22 `write` File to which initial transient solution (before steady-state) is to be written.
- 23 `writefinal` File to which final transient solution in steady-state is to be written.
- 24 `swapfile` Temporary file that holds steady-state information. Tells Spectre to use a regular file rather than virtual memory to hold the periodic operating point. Use this option if Spectre complains about not having enough memory to complete this analysis.

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Analysis Statements

Integration method parameters

25 `method` Integration method. Default derived from `errpreset`. Possible values are `euler`, `trap`, `traponly`, `gear2`, or `gear2only`.

Accuracy parameters

26 `errpreset` Selects a reasonable collection of parameter settings. Possible values are `liberal`, `moderate` or `conservative`.

27 `relref` Reference used for the relative convergence criteria. Default derived from `errpreset`. Possible values are `pointlocal`, `alllocal`, `sigglobal`, or `allglobal`.

28 `lteratio` Ratio used to compute LTE tolerances from Newton tolerance. Default derived from `errpreset`.

29 `steadyratio` Ratio used to compute steady state tolerances from LTE tolerance. Default derived from `errpreset`.

30 `maxperiods` Maximum number of simulated periods to reach steady-state.

31 `itres=1e-4` Relative tolerance for linear solver.

32 `finitediff` Options for finite difference method refinement after quasi-periodic shooting method. `finitediff` is changed from `no` to `samegrid` automatically when `readqpss` and `writeqpss` are used to re-use QPSS results. Possible values are `no`, `yes` or `refine`.

Annotation parameters

33 `stats=no` Analysis statistics. Possible values are `no` or `yes`.

34 `annotate=sweep` Degree of annotation. Possible values are `no`, `title`, `sweep`, `status`, or `steps`.

35 `title` Analysis title.

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Analysis Statements

Newton parameters

- 36 `maxiters=5` Maximum number of iterations per time step.
- 37 `restart=yes` Restart the DC solution from scratch if any condition has changed. If not, use the previous solution as initial guess. Possible values are `no` or `yes`.

Circuit age

- 38 `circuitage (Years)` Stress Time. Age of the circuit used to simulate hot-electron degradation of MOSFET and BSIM circuits.
- 39 `writeqpss` File to which final quasi-periodic steady-state solution is to be written. Small signal analyses such as `qpac`, `qpxf` and `qpnoise` can read in the steady-state solution from this file directly instead of running the `qpss` analysis again.
- 40 `readqpss` File from which final quasi-periodic steady-state solution is to be read. Small signal analyses such as `qpac`, `qpxf` and `qpnoise` can read in the steady-state solution from this file directly instead of running the `qpss` analysis again.

Most of QPSS analysis parameters are inherited from PSS analysis, and their meanings remain essentially unchanged. Two new important parameters are `funds` and `maxharms`. They replace and extend the role of `fund` and `harms` parameters of PSS analysis. One important difference is that `funds` accepts a list of fundamental names instead of actual frequencies. The frequencies associated with fundamentals are figured out automatically by the simulator. An important feature is that each input signal can be a composition of more than one sources. However, these sources must have the same fundamental name. For each fundamental name, its fundamental frequency is the greatest common factor of all frequencies associated with the name. Missing or not listing all fundamental names using the parameter `funds` will result in an amputation of the current simulation. However if `maxharms` is not given, a warning message will be issued, and the number of harmonics is defaulted to 1 for each of the fundamentals.

For QPSS analyses, the role of some PSS parameters is extended. The parameter `maxperiods` that controls the maximum number of shooting iterations for PSS analysis also controls the number of the maximum number of shooting iterations for QPSS analysis. Its default value is set to 50.

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The `tstab` parameter controls both the length of the initial transient integration with only the clock tone activated and the number of stable iterations with moderate tones activated. The stable iterations are run before Newton iterations.

The `errpreset` parameter lets you adjust several simulator parameters to fit your needs. In most cases, `errpreset` should be the only parameter you need to adjust. If you want a fast simulation with reasonable accuracy, you might set `errpreset` to `liberal`. If have some concern for accuracy, you might set `errpreset` to `moderate`. If accuracy is your main interest, you might set `errpreset` to `conservative`.

If users do not specify `steadyratio`, it is always 1.0, and it is not affected by `errpreset`. The following table shows the effect of `errpreset` on other parameters.

Parameter defaults as a function of `errprese`

<code>errpreset</code>	<code>reltol</code>	<code>relref</code>	<code>method</code>	<code>lteratio</code>	<code>maxstep</code>
<code>liberal</code>	1e-3	<code>sigglobal</code>	<code>gear2only</code>	3.5	clock period/80
<code>moderate</code>	1e-4	<code>siggloaal</code>	<code>gear2only</code>	3.5	clock period/100
<code>conservative</code>	1e-5	<code>sigglobal</code>	<code>gear2only</code>	*	clock period/200

* : `lteratio`=10.0 for conservative `errpreset` by default. Only if user specified `reltol` <= 1e-4*10.0/3.5, it will change to `lteratio`=3.5.

The new `errpreset` settings include a new default `reltol` which is actually an enforced upper limit for appropriate setting. An increase of `reltol` above default will be ignored by the simulator. User can decrease this value in the options statement. The only way to increase `reltol` is to `relax errpreset`. Spectre sets the value of `maxstep` so that it is no larger than the value given in the table. Except for `reltol` and `maxstep`, `errpreset` does not change the value of any parameters you have explicitly set. The actual values used for the QPSS analysis are given in the log file.

If `errpreset` is not specified in the netlist, `liberal` settings will be used.

The default value for `compression` is `no`. The output file stores data for every signal at every time point for which Spectre calculates a solution. Spectre saves the x axis data only once, since every signal has the same x value. If `compression=yes`, Spectre writes data to the output file only if the signal value changes by at least 2*the convergence criteria. In order to save data for each signal independently, x axis information corresponding to each signal must be saved. If the signals stay at constant values for large periods of the simulation time, setting `compression=yes` results in a smaller output data file. If the signals in your circuit move around a lot, setting `compression=yes` results in a larger output data file.

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Periodic Noise Analysis (pnoise)

Description

The Periodic Noise, or PNoise analysis is similar to the conventional noise analysis, except that it includes frequency conversion effects. Hence is it useful for predicting the noise behavior of mixers, switched-capacitor filters, and other periodically driven circuits. It is particularly useful for predicting the phase noise of autonomous circuits, such as oscillators.

PNoise analysis linearizes the circuit about the periodic operating point computed in the prerequisite PSS analysis. It is the periodically time-varying nature of the linearized circuit that accounts for the frequency conversion. In addition, the affect of a periodically time-varying bias point on the noise generated by the various components in the circuit is also included.

The time-average of the noise at the output of the circuit is computed in the form of a spectral density versus frequency. The output of the circuit is specified with either a pair of nodes or a probe component. To specify the output of a circuit with a probe, specify it using the `oprobe` parameter. If the output is voltage (or potential), choose a `resistor` or a `port` as the output probe. If the output is current (or flow), choose a `vsource` or `iprobe` as the output probe.

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Analysis Statements

If the input-referred noise or noise figure is desired, specify the input source using the `iprobe` parameter. For input-referred noise, use either a `vsource` or `isource` as the input probe; for noise figure, use a `port` as the probe. Currently, only a `vsource`, an `isource`, or a `port` may be used as an input probe. If the input source is noisy, as is a `port`, the noise analysis will compute the noise factor (F) and noise figure (NF). To match the IEEE definition of noise figure, the input probe must be a `port` with no excess noise and its `noisetemp` must be set to 16.85C (290K). In addition, the output load must be a `resistor` or `port` and must be identified as the `oprobe`.

The reference sideband (`refsideband`) specifies which conversion gain is used when computing input-referred noise, noise factor, and noise figure. The reference sideband specifies the input frequency relative to the output frequency with:

$$|f(\text{input})| = |f(\text{out}) + \text{refsideband} * \text{fund}(\text{pss})|$$

Use `refsideband=0` when the input and output of the circuit are at the same frequency (such as with amplifiers and filters). When `refsideband` differs from 0, the single side-band noise figure is computed.

The noise analysis always computes the total noise at the output, which includes contributions from the input source and the output load. The amount of the output noise that is attributable to each noise source in the circuit is also computed and output individually. If the input source is identified (using `iprobe`) and is a `vsource` or `isourec`, the input-referred noise is computed, which includes the noise from the input source itself. Finally, if the input source is identified (using `iprobe`) and is noisy, as is the case with `ports`, the noise factor and noise figure are computed. Thus if

No = total output noise

Ns = noise at the output due to the input probe (the source)

Nsi = noise at the output due to the image harmonic at the source

Nso = noise at the output due to harmonics other than input at the source

Nl = noise at the output due to the output probe (the load)

IRN = input referred noise

G = gain of the circuit

F = noise factor

NF = noise figure

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Fdsb = double sideband noise factor

NFdsb = double sideband noise figure

Fieeee = IEEE single sideband noise factor

NFieeee = IEEE single sideband noise figure

then,

$$\text{IRN} = \sqrt{N_o^2/G^2}$$

$$F = (N_o^2 - N_i^2)/N_s^2$$

$$\text{NF} = 10 \cdot \log_{10}(F)$$

$$\text{Fdsb} = (N_o^2 - N_i^2)/(N_s^2 + N_{si}^2)$$

$$\text{NFdsb} = 10 \cdot \log_{10}(\text{Fdsb})$$

$$\text{Fieeee} = (N_o^2 - N_i^2 - N_{so}^2)/N_s^2$$

$$\text{NFieeee} = 10 \cdot \log_{10}(\text{Fieeee}).$$

When the results are output, *No* is named *out*, *IRN* is named *in*, *G* is named *gain*, *F*, *NF*, *Fdsb*, *NFdsb*, *Fieeee*, and *NFieeee* are named *F*, *NF*, *Fdsb*, *NFdsb*, *Fieeee*, and *NFieeee* respectively.

Unlike other analyses in Spectre, this analysis can only sweep frequency.

Definition

Name [p] [n] pnoise parameter=value ...

The optional terminals (*p* and *n*) specify the output of the circuit. If you do not give the terminals, then you must specify the output with a probe component.

Parameters

Sweep interval parameters

1 *start=0* Start sweep limit.

2 *stop* Stop sweep limit.

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3	<code>center</code>	Center of sweep.
4	<code>span=0</code>	Sweep limit span.
5	<code>step</code>	Step size, linear sweep.
6	<code>lin=50</code>	Number of steps, linear sweep.
7	<code>dec</code>	Points per decade.
8	<code>log=50</code>	Number of steps, log sweep.
9	<code>values=[...]</code>	Array of sweep values.
10	<code>sweepstype</code>	Specifies if the sweep frequency range is absolute frequency of input or if it is relative to the port harmonics. Possible values are <code>absolute</code> or <code>relative</code> .
11	<code>relharmnum=1</code>	Harmonic to which relative frequency sweep should be referenced.

Probe parameters

12	<code>oprobe</code>	Compute total noise at the output defined by this component.
13	<code>iprobe</code>	Refer the output noise to this component.
14	<code>refsideband</code>	Conversion gain associated with this sideband is used when computing input-referred noise or noise figure.

Output parameters

15	<code>noisetype=sources</code>	Specifies if the noise analysis should output cross-power densities or noise source information. Possible values are <code>sources</code> , <code>correlations</code> or <code>timedomain</code> .
16	<code>maxsideband=7</code>	Maximum sideband included when computing noise either up-converted or down-converted to the output by the periodic drive signal.

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- 17 `sidebands=[...]` Array of relevant sidebands for the analysis.
- 18 `save` Signals to output.
Possible values are `all`, `lvl`, `allpub`, `lvlpub`, `selected`, or `none`.
- 19 `nestlvl` Levels of subcircuits to output.
- 20 `maxcycles=0` Maximum cycle correlation frequency included when computing noise either up-converted or down-converted to the output by the periodic drive signal.
- 21 `cycles=[...]` Array of relevant cycle frequencies. Valid only if `noisetype=correlations`.
- 22 `noiseskipcount=0` Calculate time-domain noise on only one of every `noiseskipcount` time points.
- 23 `noisetimepoints=[...]` Additional time points for time-domain noise analysis.
- 24 `numberofpoints=-1` Number of time points of interested to output. When ≥ 0 , it is effective, `noiseskipcount` is disabled; Defaultly, it is disabled and simulator uses `noiseskipcount` for ouput.
- 25 `saveallsidebands=no` Save noise contributors by sideband.
Possible values are `no` or `yes`.

Convergence parameters

- 26 `tolerance=1e-9` Relative tolerance for linear solver.
- 27 `gear_order=2` Gear order used for small-signal integration.
- 28 `solver=turbo` Solver type.
Possible values are `std` or `turbo`.
- 29 `oscsolver=turbo` Oscillator solver type.
Possible values are `std` or `turbo`.

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Annotation parameters

- 30 `annotate=sweep` Degree of annotation.
Possible values are `no`, `title`, `sweep`, `status`, or `steps`.
- 31 `stats=no` Analysis statistics.
Possible values are `no` or `yes`.
- 32 `title` Analysis title.

In practice, noise can mix with each of the harmonics of the periodic drive signal applied in the PSS analysis and end up at the output frequency. However, the PNoise analysis only includes the noise that mixes with a finite set of harmonics that are typically specified using the `maxsideband` parameter, but in special circumstances may be specified with the `sidebands` parameter. If K_i represents sideband i , then

$$f(\text{noise_source}) = f(\text{out}) + K_i * \text{fund}(\text{pss})$$

The `maxsideband` parameter specifies the maximum $|K_i|$ included in the PNoise calculation. Thus, noise at frequencies less than $f(\text{out}) - \text{maxsideband} * \text{fund}(\text{pss})$ and greater than $f(\text{out}) + \text{maxsideband} * \text{fund}(\text{pss})$ are ignored. If selected sidebands are specified using the `sidebands` parameter, then only those are included in the calculation. Care should be taken when specifying the sidebands because the results will be in error if you do not include a sideband that contributes significant noise to the output.

The number of requested sidebands does not change substantially the simulation time. However, the `maxacfreq` of the corresponding PSS analysis should be set to guarantee that $|\max\{f(\text{noise_source})\}|$ is less than `maxacfreq`, otherwise the computed solution might be contaminated by aliasing effects. The PNoise simulation is not executed for $|f(\text{out})|$ greater than `maxacfreq`. Diagnostic messages are printed for those extreme cases, indicating which `maxacfreq` should be set in the PSS analysis. In the majority of the simulations, however, this is not an issue, because `maxacfreq` is never allowed to be smaller than 40x the PSS fundamental.

Phase Noise measurements are possible using the Analog Artist(ADE) environment. Two noise analyses are preconfigured for this simulation and most of the parameters are set by Artist. First noise analysis named `mod1` is a regular noise analysis and can be used independently. The second noise correlation analysis called `mod2` has a limited use outside of the Artist environment. Direct Plot is configured to analyze these results and combine several wave forms to measure AM and PM components of the output noise. For details, please see the SpectreRF User Guide.

You can specify sweep limits by giving the end points or by providing the center value and the span of the sweep. Steps can be linear or logarithmic, and you can specify the number of

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steps or the size of each step. You can give a step size parameter (`step`, `lin`, `log`, `dec`) to determine whether the sweep is linear or logarithmic. If you do not give a step size parameter, the sweep is linear when the ratio of stop to start values is less than 10, and logarithmic when this ratio is 10 or greater. Alternatively, you may specify the particular values that the sweep parameter should take using the `values` parameter. If you give both a specific set of values and a set specified using a sweep range, the two sets are merged and collated before being used. All frequencies are in Hertz.

Parameter Index

In the following index, the number following each parameter name indicates where to find the description of that parameter.

<code>annotate</code>	30	<code>maxcycles</code>	20	<code>oscsolver</code>	29	<code>start</code>	1
<code>center</code>	3	<code>maxsideband</code>	16	<code>refsideband</code>	14	<code>stats</code>	31
<code>cycles</code>	21	<code>nestlvl</code>	19	<code>relharmnum</code>	11	<code>step</code>	5
<code>dec</code>	7	<code>noiseskipcount</code>	22	<code>save</code>	18	<code>stop</code>	2
<code>gear_order</code>	27	<code>noisetimepoints</code>	23	<code>saveallsidebands</code>	25	<code>sweeptype</code>	10
<code>iprobe</code>	13	<code>noisetype</code>	15	<code>sidebands</code>	17	<code>title</code>	32
<code>lin</code>	6	<code>numberofpoints</code>	24	<code>solver</code>	28	<code>tolerance</code>	26
<code>log</code>	8	<code>oprobe</code>	12	<code>span</code>	4	<code>values</code>	9

Periodic S-Parameter Analysis (psp)

Description

The periodic SP (PSP) analysis is used to compute scattering and noise parameters for n-port circuits that exhibit frequency translation, such as mixers. It is a small-signal analysis like SP analysis, except, as in PAC and PXF, the circuit is first linearized about a periodically varying operating point as opposed to a simple DC operating point. Linearizing about a periodically time-varying operating point allows the computation of S-parameters between circuit ports that convert signals from one frequency band to another. PSP can also calculate noise parameters in frequency-converting circuits. PSP computes noise figure (both single-sideband and double-sideband), input referred noise, equivalent noise parameters, and noise correlation matrices. As in PNoise, but unlike SP, the noise features of the PSP analysis include noise folding effects due to the periodically time-varying nature of the circuit.

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Computing the n-port S-parameters and noise parameters of a periodically varying circuit is a two step process. First, the small stimulus is ignored and the periodic steady-state response of the circuit to possibly large periodic stimulus is computed using PSS analysis. As a normal part of the PSS analysis, the periodically time-varying representation of the circuit is computed and saved for later use. The second step is applying small-signal excitations to compute the n-port S-parameters and noise parameters. This is done using the PSP analysis. A PSP analysis cannot be used alone, it must follow a PSS analysis. However, any number of periodic small-signal analyses such as PAC, PSP, PXF, PNoise, can follow a single PSS analysis.

Unlike other analyses in Spectre, this analysis can only sweep frequency.

Definition

Name `psp parameter=value ...`

Parameters

Sweep interval parameters

1	<code>start=0</code>	Start sweep limit.
2	<code>stop</code>	Stop sweep limit.
3	<code>center</code>	Center of sweep.
4	<code>span=0</code>	Sweep limit span.
5	<code>step</code>	Step size, linear sweep.
6	<code>lin=50</code>	Number of steps, linear sweep.
7	<code>dec</code>	Points per decade.
8	<code>log=50</code>	Number of steps, log sweep.
9	<code>values=[...]</code>	Array of sweep values.
10	<code>sweepstype</code>	Specifies if the sweep frequency range is absolute frequency of input or if it is relative to the port harmonics. Possible values are <code>absolute</code> or <code>relative</code> .

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Port parameters

- 11 `ports=[...]` List of active ports. Ports are numbered in the order given. For purposes of noise figure computation, the input is considered port 1 and the output is port 2.
- 12 `portharmsvec=[...]` List of harmonics active on specified list of ports. Must have a one-to-one correspondence with the ports vector.
- 13 `harmsvec=[...]` List of harmonics, in addition to ones associated with specific ports by `portharmsvec`, that are active.

Output parameters

- 14 `freqaxis` Specifies whether the results should be output versus the input frequency, the output frequency, or the absolute value of the input frequency. Default is `in`. Possible values are `absin`, `in` or `out`.

Noise parameters

- 15 `donoise=yes` Perform noise analysis. If `oprobe` is specified as a valid port, this is set to `yes`, and a detailed noise output is generated. Possible values are `no` or `yes`.

Probe parameters

- 16 `maxsideband=7` Maximum sideband included when computing noise either up-converted or down-converted to the output by the periodic drive signal.

Convergence parameters

- 17 `tolerance=1e-9` Relative tolerance for linear solver.
- 18 `gear_order=2` Gear order used for small-signal integration.
- 19 `solver=turbo` Solver type. Possible values are `std` or `turbo`.

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20 `oscsolver=turbo` Oscillator solver type.
Possible values are `std` or `turbo`.

Annotation parameters

21 `annotate=sweep` Degree of annotation.
Possible values are `no`, `title`, `sweep`, `status`, or `steps`.

22 `stats=no` Analysis statistics.
Possible values are `no` or `yes`.

23 `title` Analysis title.

To specify the PSP analysis the port and port harmonic relations must be specified. You can select the ports of interest by setting the `port` parameter and the set of periodic small-signal output frequencies of interest by setting `portharmsvec` or the `harmsvec` parameters. For a given set of n integer numbers representing the harmonics K_1, K_2, \dots, K_n , the scattering parameters at each port are computed at the frequencies $f(\text{scattered}) = f(\text{rel}) + K_i * \text{fund}(\text{pss})$, where $f(\text{rel})$ represents the relative frequency of a signal incident on a port, $f(\text{scattered})$ represents the frequency to which the relevant scattering parameter represents the conversion, and $\text{fund}(\text{pss})$ represents the fundamental frequency used in the corresponding PSS analysis.

Thus, when analyzing a down-converting mixer, with signal in the upper sideband, and sweeping the RF input frequency, the most relevant harmonic for RF input is $K_i = 1$ and for IF output $K_i = 0$. Hence we can associate $K_2 = 1$ with the IF port and $K_1 = 0$ with the RF port. S_{21} will represent the transmission of signal from the RF to IF, and S_{11} the reflection of signal back to the RF port. If the signal was in the lower sideband, then a choice of $K_1 = -1$ would be more appropriate.

Either `portharmsvec` or `harmsvec` parameters can be used to specify the harmonics of interest. If `portharmsvec` is given, the harmonics must be in one-to-one correspondence with the ports, with each harmonic associated with a single port. If harmonics are specified in the optional `harmsvec` parameter, then all possible frequency-translating scattering parameters associated with the specified harmonics are computed.

With PSP the frequency of the input and of the response are usually different (this is an important way in which PSP differs from SP). Because the PSP computation involves inputs and outputs at frequencies that are relative to multiple harmonics, the `freqaxis` and `sweepstype` parameters behave somewhat differently in PSP than in PAC and PXF.

The `sweepstype` parameter controls the way the frequencies in the PSP analysis are swept. Specifying a `relative` sweep indicates to sweep relative to the analysis harmonics (not the

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PSS fundamental) and an `absolute` sweep is a sweep of the absolute input source frequency. For example, with a PSS fundamental of 100MHz, `port harmsvec` set to [9 1] to examine a downconverting mixer, `sweep type=relative`, and a sweep range of $f(\text{rel})=0\text{-} >50\text{MHz}$, then S21 would represent the strength of signal transmitted from the input port in the range 900->950MHz to the output port at frequencies 100->150MHz. Using `sweep type=absolute` and sweeping the frequency from 900->950MHz would calculate the same quantities, since $f(\text{abs})=900\text{-} >950\text{MHz}$, and $f(\text{rel}) = f(\text{abs}) - K1 * \text{fund}(\text{pss}) = 0\text{-} >50\text{MHz}$, because $K1=9$ and $\text{fund}(\text{pss}) = 100\text{MHz}$.

Usually it is not necessary to sweep frequency in PSP over more than one fundamental PSS period.

The `freqaxis` parameter is used to specify whether the results should be output versus the scattered frequency at the input port (`in`), the scattered frequency at the output port (`out`), or the absolute value of the frequency swept at the input port (`absin`).

Unlike in PAC/PXF/PNoise, increasing the number of requested ports and harmonics will increase the simulation time substantially.

To ensure accurate results in PSP, the `maxacfreq` of the corresponding PSS analysis should be set to guarantee that $|\max\{f(\text{scattered})\}|$ is less than `maxacfreq`, otherwise the computed solution might be contaminated by aliasing effects.

PSP analysis also computes noise figures, equivalent noise sources, and noise parameters. The noise computation, which is skipped only when `donoise=no`, requires additional simulation time. If

No = total output noise at frequency f

Ns = noise at the output due to the input probe (the source)

Nsi = noise at the output due to the image harmonic at the source

Nso = noise at the output due to harmonics other than input at the source

Nl = noise at the output due to the output probe (the load)

IRN = input referred noise

G = gain of the circuit

F = noise factor (single side band)

NF = noise figure (single side band)

Fdsb = double sideband noise factor

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NFdsb = double sideband noise figure

Fieeee = IEEE single sideband noise factor

NFieeee = IEEE single sideband noise figure

then,

$$\text{IRN} = \sqrt{\text{No}^2/\text{G}^2}$$

$$\text{F} = (\text{No}^2 - \text{NI}^2)/\text{Ns}^2$$

$$\text{NF} = 10 \cdot \log_{10}(\text{F})$$

$$\text{Fdsb} = (\text{No}^2 - \text{NI}^2)/(\text{Ns}^2 + \text{Nsi}^2)$$

$$\text{NFdsb} = 10 \cdot \log_{10}(\text{Fdsb})$$

$$\text{Fieeee} = (\text{No}^2 - \text{NI}^2 - \text{Nso}^2)/\text{Ns}^2$$

$$\text{NFieeee} = 10 \cdot \log_{10}(\text{Fieeee}).$$

When the results are output, IRN is named `in`, G is named `gain`, F, NF, Fdsb, NFdsb, Fieeee, and NFieeee are named `F`, `NF`, `Fdsb`, `NFdsb`, `Fieeee`, and `NFieeee` respectively. Note that the gain computed by PSP is the voltage gain from the actual circuit input to the circuit output, not the gain from the internal port voltage source to the output.

To ensure accurate noise calculations, the `maxsideband` or `sidebands` parameters must be set to include the relevant noise folding effects. `maxsideband` is only relevant to the noise computation features of PSP.

You can specify sweep limits by giving the end points or by providing the center value and the span of the sweep. Steps can be linear or logarithmic, and you can specify the number of steps or the size of each step. You can give a step size parameter (`step`, `lin`, `log`, `dec`) to determine whether the sweep is linear or logarithmic. If you do not give a step size parameter, the sweep is linear when the ratio of stop to start values is less than 10, and logarithmic when this ratio is 10 or greater. Alternatively, you may specify the particular values that the sweep parameter should take using the `values` parameter. If you give both a specific set of values and a set specified using a sweep range, the two sets are merged and collated before being used. All frequencies are in Hertz.

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Parameter Index

In the following index, the number following each parameter name indicates where to find the description of that parameter.

annotate	21	harmsvec	13	ports	11	stop	2
center	3	lin	6	solver	19	sweeptype	10
dec	7	log	8	span	4	title	23
donoise	15	maxsideband	16	start	1	tolerance	17
freqaxis	14	oscsolver	20	stats	22	values	9
gear_order	18	portharmsvec	12	step	5		

Periodic Steady-State Analysis (pss)

Description

This analysis computes the periodic steady-state (PSS) response of a circuit, with a simulation time independent of the time-constants of the circuit. Also, it sets the circuits periodic operating point, which can then be used during a periodic time-varying small-signal analysis, such as PAC, PXF, and PNOISE.

PSS analysis is capable of handling both autonomous (non-driven) and driven (non-autonomous) circuits. Autonomous circuits are time-invariant circuits that have time-varying responses. Thus, autonomous circuits generate non-constant waveforms even though they are not driven by a time-varying stimulus. Driven circuits require some time-varying stimulus to generate a time-varying response. The most common example of an autonomous circuit is an oscillator. Common driven circuits include amplifiers, filters, mixers, etc.

With driven circuits the user specifies the analysis `period`, or its corresponding fundamental frequency `fund`. The `period` must be an integer multiple of the period of the drive signal or signals. Autonomous circuits have no drive signal and the actual period of oscillation is not known precisely by the user in advance. Instead, the user specifies an estimate of the oscillation period and PSS analysis computes the precise period along with the periodic solution waveforms.

When applied to autonomous circuits, PSS analysis requires the user to specify a pair of nodes, `p` and `n`. In fact this is how PSS analysis determines whether it is being applied to an autonomous or a driven circuit. If the pair of nodes is supplied, PSS assumes the circuit is autonomous; if not, the circuit is assumed to be driven.

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A PSS analysis consists of two phases, an initial transient phase, which allows the circuit to be initialized, and the shooting phase, which is where the periodic steady-state solution is computed. The transient phase consists of three intervals.

The first starts at `tstart`, which is normally 0, and continues through the onset of periodicity `tonset` for the independent sources. The onset of periodicity, which is automatically generated, is the minimum time for which all sources are periodic. The second is an optional user specified stabilization interval whose length is `tstab`. The final interval whose length is `period` for driven circuits, or `4xperiod` for autonomous circuits has a special use for the autonomous PSS analysis, i.e., the PSS analysis monitors the waveforms in the circuit and develops a better estimate of the oscillation period. Once the initial transient phase is complete, the shooting interval begins. In this phase, the circuit is repeatedly simulated over one period while adjusting the initial condition (and the period when applied to autonomous circuits) to find the periodic steady-state solution.

Typically the process takes three to five such iterations to reach steady-state. Upon completion, if requested by the user, the frequency-domain response is computed. For driven circuits, one can use `writepss` and `readpss` to reuse the results in a previous simulation.

Definition

Name [p] [n] pss parameter=value ...

Parameters

Simulation interval parameters

- | | | |
|---|---------------------------|---|
| 1 | <code>period</code> (s) | Steady state analysis period (or its estimate for autonomous circuits). |
| 2 | <code>fund</code> (Hz) | Alternative to period specification. Steady state analysis fundamental frequency (or its estimate for autonomous circuits). |
| 3 | <code>tstab=0.0</code> s | Extra stabilization time after the onset of periodicity for independent sources. |
| 4 | <code>tstart=0.0</code> s | Initial transient analysis start time. |

Time-step parameters

- | | | |
|---|--------------------------|--|
| 5 | <code>maxstep</code> (s) | Maximum time step. Default derived from <code>errpreset</code> . |
|---|--------------------------|--|

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- 6 `maxacfreq` Maximum frequency requested in a subsequent periodic small-signal analysis. Default derived from `errpreset` and `harms`.
- 7 `step=0.001 period s` Minimum time step that would be used solely to maintain the aesthetics of the results.

Initial-condition parameters

- 8 `ic=all` What should be used to set initial condition. Possible values are `dc`, `node`, `dev`, or `all`.
- 9 `skipdc=no` If yes, there will be no dc analysis for transient. Possible values are `no`, `yes` or `sigrampup`.
- 10 `readic` File that contains initial condition.

Convergence parameters

- 11 `readns` File that contains estimate of initial transient solution.
- 12 `cmin=0 F` Minimum capacitance from each node to ground.

Output parameters

- 13 `harms=9` Number of harmonics to output when `outputtype=freq` or `all`.
- 14 `harmsvec=[...]` Array of desired harmonics. Alternate form of `harms` that allows selection of specific harmonics.
- 15 `outputtype=time` for PSS, `freq` for QPSS
Output type.
Possible values are `all`, `time` or `freq`.
- 16 `save` Signals to output.
Possible values are `all`, `lvl`, `allpub`, `lvlpub`, `selected`, or `none`.
- 17 `nestlvl` Levels of subcircuits to output.

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- 18 `oppoint=no` Should operating point information be computed for initial timestep, and if so, where should it be sent. Possible values are `no`, `screen`, `logfile`, or `rawfile`.
- 19 `skipstart=starttime s` The time to start skipping output data.
- 20 `skipstop=stoptime s` The time to stop skipping output data.
- 21 `skipcount` Save only one of every skipcount points.
- 22 `strobeperiod (s)` The output strobe interval (in seconds of transient time).
- 23 `strobedelay=0 s` The delay (phase shift) between the skipstart time and the first strobe point.
- 24 `compression=no` Do data compression on output. See full description below. Possible values are `no` or `yes`.
- 25 `saveinit=no` If set, the waveforms for the initial transient before steady state are saved. Possible values are `no` or `yes`.

State-file parameters

- 26 `write` File to which initial transient solution (before steady-state) is to be written.
- 27 `writefinal` File to which final transient solution in steady-state is to be written.
- 28 `swapfile` Temporary file that holds steady-state information. Tells Spectre to use a regular file rather than virtual memory to hold the periodic operating point. Use this option if Spectre complains about not having enough memory to complete this analysis.
- 29 `writepss` File to which the converged steady-state solution is to be written. `finitediff` is set to `yes` automatically to improve PSS results.
- 30 `readpss` File from which a previously converged steady-state solution is to be read. PSS loads the solution and checks the residue of the

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circuit equations only. The solution is re-used if the residue is satisfying. Otherwise, the solution is re-converged using the finite difference method.

Integration method parameters

- 31 `method` Integration method. Default derived from `errpreset`. Possible values are `euler`, `trap`, `traponly`, `gear2`, or `gear2only`.
- 32 `tstabmethod` Integration method used in stabilization time. Default is `traponly` for autonomous circuits, or is derived from `errpreset` for driven circuits. Possible values are `euler`, `trap`, `traponly`, `gear2`, or `gear2only`.

Accuracy parameters

- 33 `errpreset` Selects a reasonable collection of parameter settings. Possible values are `liberal`, `moderate` or `conservative`.
- 34 `relref` Reference used for the relative convergence criteria. Default derived from `errpreset`. Possible values are `pointlocal`, `alllocal`, `sigglobal`, or `allglobal`.
- 35 `lteratio` Ratio used to compute LTE tolerances from Newton tolerance. Default derived from `errpreset`.
- 36 `steadyratio` Ratio used to compute steady state tolerances from LTE tolerance. Default derived from `errpreset`.
- 37 `maxperiods` Maximum number of simulated periods to reach steady-state.
- 38 `itres=1e-4` Relative tolerance for linear solver.
- 39 `finitediff` Options for finite difference method refinement after shooting method for driven circuits. Possible values are `no`, `yes` or `refine`.

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- 40 `highorder` Perform a high-order refinement after low-order convergence. The Multi-Interval Chebyshev polynomial spectral algorithm will be used. Possible values are `no` or `yes`.
- 41 `psaratio=1` Ratio used to compute high-order polynomial spectral accuracy from Newton tolerance.
- 42 `maxorder=16` The maximum order of the Chebyshev polynomials used in waveform approximation. Possible values are from 2 to 16.
- 43 `fullpssvec` Use the full vector containing solutions at all PSS time steps in the linear solver. Default derived from the size of the equation and the property of the PSS time steps. Possible values are `no` or `yes`.

Annotation parameters

- 44 `stats=no` Analysis statistics. Possible values are `no` or `yes`.
- 45 `annotate=sweep` Degree of annotation. Possible values are `no`, `title`, `sweep`, `status`, or `steps`.
- 46 `title` Analysis title.

Newton parameters

- 47 `maxiters=5` Maximum number of iterations per time step.
- 48 `restart=yes` Restart the DC solution from scratch if any condition has changed. If not, use the previous solution as initial guess. Possible values are `no` or `yes`.

Circuit age

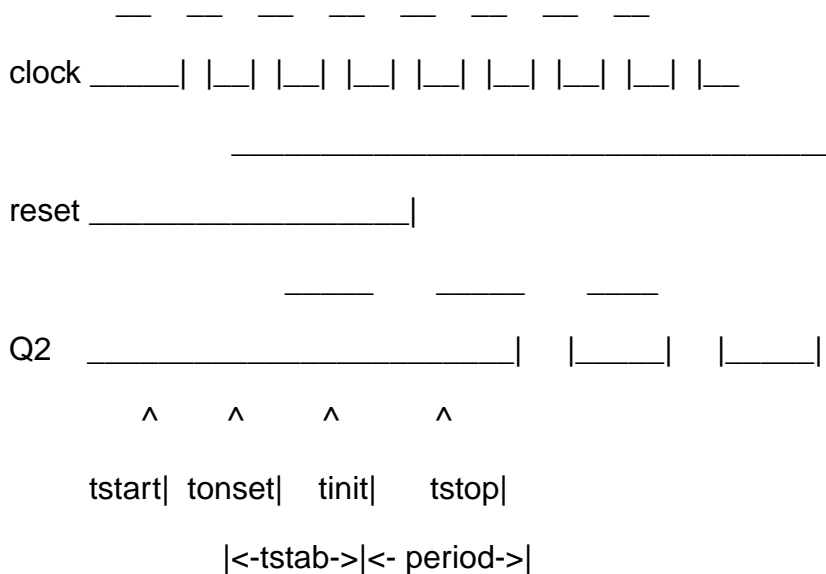
- 49 `circuitage (Years)` Stress Time. Age of the circuit used to simulate hot-electron degradation of MOSFET and BSIM circuits.

The initial transient analysis provides a flexible mechanism to direct the circuit to a particular steady-state solution of interest, and to avoid undesired solutions. Another usage of the initial

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transient simulation is helping convergence by eliminating large but fast decaying modes that are present in many circuits. For example, in case of driven circuits, consider the reset signal in the figure below.



In the figure above, the initial transient analysis is executed from t_{start} to t_{stop} . If initial transient results are relevant, you can output them by setting `saveinit` to `yes`. The steady-state results are always computed for the specified `period`, from t_{init} to t_{stop} . By default, t_{start} and t_{stab} are set to zero, while t_{init} , t_{onset} and t_{stop} are always automatically generated.

It happens in some circuits that the linearity of the relationship between the initial and final state depends on when the shooting interval begins. Conceptually, when the shooting interval begins should not matter, as long as it is after the time when the stimuli have become periodic, because the periodic response repeats endlessly. However in practice, one can improve the convergence by starting at a good point, and degrade the convergence, which slows the analysis, by starting in a bad spot. In general, it is best to try to avoid starting the shooting interval at a point where the circuit is undergoing strong nonlinear behavior. For example, in switch-capacitor filters it is best if t_{init} falls at the beginning of a clock transition, preferably a transition that follows a relatively long period of settling. If instead t_{init} occurred during a clock transition or soon after one, then it is likely the opamps would be undergoing slew-rate limiting at the start of the shooting interval, which would act to slow convergence. Switching mixers follow similar rules.

When applying PSS analysis to oscillators, it is necessary to start the oscillator, just as you would if you were simulating the turn-on transient of the oscillator using transient analysis. The Designers Guide to Spice and Spectre [K. S. Kundert, Kluwer Academic Publishers, 1995] describes techniques for starting oscillators in some depth. In summary, there are two

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techniques for starting oscillators, using initial conditions, or using a brief impulsive stimulus. Initial conditions would be provided for the components of the oscillators resonator. If an impulsive stimulus is used, it should be applied so as to couple strongly into the oscillatory mode of the circuit, and poorly into any other long-lasting modes, such as those associated with bias circuitry. Either way, once the trigger is applied to start the oscillator, it is important to allow the oscillator to run for a while before the shooting methods

are applied to compute the steady-state result. To do so, specify an additional stabilization interval using the `tstab` parameter. In practice, an additional stabilization interval often improves convergence.

By default, only the time-domain results are computed. If you specify either `harms` or `harmsvec` or set `outputtype` to `freq` or `all`, the frequency-domain results will also be computed. If frequency-domain results are requested, but the desired harmonics are not specified, its default value is 10. The time-domain output waveform generation can be inhibited by setting `outputtype` to `freq`.

The accuracy of the results does not depend on the number of harmonics that are requested, only on the accuracy parameters, which are set in the same fashion as in the transient analysis. Besides a few new parameters, like `steadyratio` and `maxacfreq`, all the others parameters work in PSS analysis in the exact same fashion as they work on transient analysis.

Several parameters determine the accuracy of the PSS analysis. `reltol` and `abstol` control the accuracy of the discretized equation solution. These parameters determine how well charge is conserved and how accurately steady-state or equilibrium points are computed. You can set the integration error, or the errors in the computation of the circuit dynamics (such as time constants), relative to `reltol` and `abstol` by setting the `lteratio` parameter.

The `steadyratio` parameter adjusts the maximum allowed mismatch in node voltages or current branches from the beginning to the end of the steady-state period. This value is multiplied by the `lteratio` and `reltol` to determine the convergence criterion. The relative convergence norm is printed out along with the actual mismatch value at the end of each iteration, thus indicating the progress of the steady-state iteration.

The parameter `maxperiods` controls the maximum number of shooting iterations for PSS analysis. Its default value is set to 20 for driven PSS and 50 for autonomous PSS.

The `finitediff` parameter allows the use of finite difference (FD) after shooting. Usually this will eliminate the above mismatch in node voltages or current branches. It can also refine the grid of time steps. In some cases, numerical error of the linear solver still introduces a mismatch. One can set `steadyratio` to a smaller value to activate a tighter tolerance for the iterative linear solver. If `finitediff` is set to `no`, FD method is turned off. If it is set to

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yes, pss applies FD method and trying to improve the beginning small time steps if necessary. If it is set to refine, pss applies FD method and tries to refine the time steps. When the simulation uses 2nd-order method, uniform 2nd order gear is used. `finitediff` is changed from no to yes automatically when `readpss` and `writepss` are used to re-use PSS results

The `maxacfreq` parameter is used to automatically adjust the `maxstep` to reduce errors due to aliasing in frequency-domain results. By default, the `maxacfreq` is set to 4x the frequency of the largest requested harmonic, but is never set to less than 40x the fundamental.

The parameter `relref` determines how the relative error is treated. The `relref` options are:

`relref=pointlocal`: Compares the relative errors in quantities at each node to that node alone.

`relref=alllocal`: Compares the relative errors at each node to the largest values found for that node alone for all past time.

`relref=sigglobal`: Compares relative errors in each of the circuit signals to the maximum for all signals at any previous point in time.

`relref=allglobal`: Same as `relref=sigglobal` except that it also compares the residues (KCL error) for each node to the maximum of that nodes past history.

The `errpreset` parameter lets you adjust the simulator parameters to fit your needs quickly. In most cases, it should also be the only parameter you need to adjust.

Guidelines for using `errpreset` in driven circuits are described in the following. If the circuit contains only one periodic tone and you are only interested in obtaining the periodic operating point, you might set `errpreset` to `liberal`, which gives a reasonably accurate result and the fastest simulation speed. If the circuit contains more than one periodic tone and you are interested in intermodulation results, you might set `errpreset` to `moderate`, which gives a very accurate result. If you want a very low noise floor in your simulation result and accuracy is your main interest, you might set `errpreset` to `conservative`. Multi-interval Chebyshev (MIC) is activated automatically for both moderate and conservative settings unless you explicitly set `highorder=no`. MIC falls back to the original method if it encounters difficulty converging. If you sets `highorder=yes`, MIC will try harder to converge.

The effect of `errpreset` on other parameters for driven circuits is shown in the following table.

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Parameter defaults and estimated numerical noise floor in simulation result as a function of `errpreset`

<code>errpreset</code>	<code>reltol</code>	<code>relref</code>	<code>method</code>	<code>lteratio</code>	<code>steadyratio</code>	<code>noisefloor</code>
<code>liberal</code>	1e-3	<code>sigglobal</code>	<code>traonly</code>	3.5	0.001	-70dB
<code>moderate</code>	1e-3	<code>alllocal</code>	<code>gear2only</code> <code>+mic</code>	3.5	0.001	-120d
<code>conservative</code>	1e-4	<code>alllocal</code>	<code>gear2only</code> <code>+mic</code>	*	0.01	-200dB

* : `lteratio`=10.0 for conservative `errpreset`. Only if user specified `reltol` <= 1e-4 * 10.0/3.5, it will change to `lteratio`=3.5.

The new `errpreset` settings include a new default `reltol` which is actually an upper limit. An increase of `reltol` above default will be ignored by the simulator. User can decrease this value in the options statement. The only way to increase `reltol` is to relax `errpreset`.

Estimated numerical noise floor is for a weakly nonlinear circuit with a successful MIC simulation. For a linear circuit, the noise floor is even lower and for a very nonlinear circuit, you might need to tighten `psaratio` or increase `maxacfreq` to achieve this noise floor. The actual values used for the PSS analysis are given in the log file.

Guidelines for using `errpreset` in autonomous circuits are described in the following. If you want a fast simulation with reasonable accuracy, you might set `errpreset` to `liberal`. If you have some concern for accuracy, you might set `errpreset` to `moderate`. If accuracy is your main interest, you might set `errpreset` to `conservative`.

The effect of `errpreset` on other parameters for autonomous circuits is shown in the following table.

Parameter defaults as a function of `errpreset`

<code>errpreset</code>	<code>reltol</code>	<code>relref</code>	<code>method</code>	<code>lteratio</code>	<code>steadyratio</code>	<code>maxstep</code>
<code>liberal</code>	1e-3	<code>sigglobal</code>	<code>traonly</code>	3.5	0.001	period/200
<code>moderate</code>	1e-4	<code>alllocal</code>	<code>gear2only</code>	3.5	0.1	period/200
<code>conservative</code>	1e-5	<code>alllocal</code>	<code>gear2only</code>	*	0.1	period/400

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* : Iteratio=10.0 for conservative `errpreset` by default. Only if user specified `reltol <= 1e-4*10.0/3.5`, it will change to `Iteratio=3.5`.

The value of `reltol` can be decreased from default in the options statement. The only way to increase `reltol` is to relax `errpreset`. Spectre sets the value of `maxstep` so that it is no larger than the value given in the table. Except for `reltol` and `maxstep`, `errpreset` does not change the value of any parameters you have explicitly set. The actual values used for the PSS analysis are given in the log file.

If `errpreset` is not specified in the netlist, `liberal` settings will be used.

If the circuit you are simulating can have infinitely fast transitions (for example, a circuit that contains nodes with no capacitance), Spectre might have convergence problems. To avoid this, you must prevent the circuit from responding instantaneously. You can accomplish this by setting `cmin`, the minimum capacitance to ground at each node, to a physically reasonable nonzero value. This often significantly improves Spectre convergence.

You may specify the initial condition for the transient analysis by using the `ic` statement or the `ic` parameter on the capacitors and inductors. If you do not specify the initial condition, the DC solution is used as the initial condition. The `ic` parameter on the transient analysis controls the interaction of various methods of setting the initial conditions. The effects of individual settings are:

`ic=dc`: Any initial condition specifiers are ignored, and the DC solution is used.

`ic=node`: The `ic` statements are used, and the `ic` parameter on the capacitors and inductors are ignored.

`ic=dev`: The `ic` parameters on the capacitors and inductors are used, and the `ic` statements are ignored.

`ic=all`: Both the `ic` statements and the `ic` parameters are used, and the `ic` parameters override the `ic` statements.

If you specify an initial condition file with the `readic` parameter, initial conditions from the file are used, and any `ic` statements are ignored.

Once you specify the initial conditions, Spectre computes the actual initial state of the circuit by performing a DC analysis. During this analysis, Spectre forces the initial conditions on nodes by using a voltage source in series with a resistor whose resistance is `rforce` (see `options`).

With the `ic` statement it is possible to specify an inconsistent initial condition (one that cannot be sustained by the reactive elements). Examples of inconsistent initial conditions include setting the voltage on a node with no path of capacitors to ground or setting the current

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through a branch that is not an inductor. If you initialize Spectre inconsistently, its solution jumps; that is, it changes instantly at the beginning of the simulation interval. You should avoid such changes if possible because Spectre can have convergence problems while trying to make the jump.

You can skip the DC analysis entirely by using the parameter `skipdc`. If the DC analysis is skipped, the initial solution will be either trivial, or given in the file you specified by the `readic` parameter, or, if the `readic` parameter is not given, the values specified on the `ic` statements. Device based initial conditions are not used for `skipdc`. Nodes that you do not specify with the `ic` file or `ic` statements will start at zero. You should not use this parameter unless you are generating a nodeset file for circuits that have trouble in the DC solution; it usually takes longer to follow the initial transient spikes that occur when the DC analysis is skipped than it takes to find the real DC solution. The `skipdc` parameter might also cause convergence problems in the transient analysis.

The possible settings of parameter `skipdc` and their meanings are:

`skipdc=no`: Initial solution is calculated using the normal DC analysis (default).

`skipdc=yes`: Initial solution is given in the file specified by the `readic` parameter or the values specified on the `ic` statements.

`skipdc=sigrampup`: Independent source values start at 0 and ramp up to their initial values in the first phase of the simulation. The waveform production in the time-varying independent source is enabled after the rampup phase. The rampup simulation is from `tstart` to `time=0` s, and the main simulation is from `time=0` s to `tstab`. If the `tstart` parameter is not specified, the default `tstart` time is set to $-0.1*tstab$.

Nodesets help find the DC or initial transient solution. You can supply them in the circuit description file with `nodeset` statements, or in a separate file using the `readns` parameter. When nodesets are given, Spectre computes an initial guess of the solution by performing a DC analysis while forcing the specified values onto nodes by using a voltage source in series with a resistor whose resistance is `rforce`. Spectre then removes these voltage sources and resistors and computes the true solution from this initial guess.

Nodesets have two important uses. First, if a circuit has two or more solutions, nodesets can bias the simulator towards computing the desired one. Second, they are a convergence aid. By estimating the solution of the largest possible number of nodes, you might be able to eliminate a convergence problem or dramatically speed convergence.

When you simulate the same circuit many times, we suggest that you use both the `write` and `readns` parameters and give the same file name to both parameters. The DC analysis then converges quickly even if the circuit has changed somewhat since the last simulation, and the nodeset file is automatically updated.

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Nodesets and initial conditions have similar implementation but produce different effects. Initial conditions actually define the solution, whereas nodesets only influence it. When you simulate a circuit with a transient analysis, Spectre forms and solves a set of differential equations. However, differential equations have an infinite number of solutions, and a complete set of initial conditions must be specified in order to identify the desired solution. Any initial conditions you do not specify are computed by the simulator to be consistent. The transient waveforms then start from initial conditions. Nodesets are usually used as a convergence aid and do not affect the final results. However, in a circuit with more than one solution, such as a latch, nodesets bias the simulator towards finding the solution closest to the nodeset values.

The `method` parameter specifies the integration method. The possible settings and their meanings are:

`method=euler:` Backward-Euler is used exclusively.

`method=traponly:` Trapezoidal rule is used almost exclusively.

`method=trap:` Backward-Euler and the trapezoidal rule are used.

`method=gear2only:` Gears second-order backward-difference method is used almost exclusively.

`method=gear2:` Backward-Euler and second-order Gear are used.

The trapezoidal rule is usually the most efficient when you want high accuracy. This method can exhibit point-to-point ringing, but you can control this by tightening the error tolerances. For this reason, though, if you choose very loose tolerances to get a quick answer, either backward-Euler or second-order Gear will probably give better results than the trapezoidal rule. Second-order Gear and backward-Euler can make systems appear more stable than they really are. This effect is less pronounced with second-order Gear or when you request high accuracy.

Spectre provides two methods for reducing the number of output data points saved: `strobing`, based on the simulation time, and `skipping` time points, which saves only every Nth point.

The parameters `strobeperiod` and `strobedelay` control the strobing method. `strobeperiod` sets the interval between points that you want to save, and `strobedelay` sets the offset within the period relative to `skipstart`. The simulator forces a time step on each point to be saved, so the data is computed, not interpolated.

The skipping method is controlled by `skipcount`. If this is set to N, then only every Nth point is saved.

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The parameters `skipstart` and `skipstop` apply to both data reduction methods. Before `skipstart` and after `skipstop`, Spectre saves all computed data.

The default value for `compression` is `no`. The output file stores data for every signal at every time point for which Spectre calculates a solution. Spectre saves the x axis data only once, since every signal has the same x value. If `compression=yes`, Spectre writes data to the output file only if the signal value changes by at least 2*the convergence criteria. In order to save data for each signal independently, x axis information corresponding to each signal must be saved. If the signals stay at constant values for large periods of the simulation time, setting `compression=yes` results in a smaller output data file. If the signals in your circuit move around a lot, setting `compression=yes` results in a larger output data file.

Parameter Index

In the following index, the number following each parameter name indicates where to find the description of that parameter.

<code>annotate</code>	45	<code>lteratio</code>	35	<code>readns</code>	11	<code>strobedelay</code>	23
<code>circuitage</code>	49	<code>maxacfreq</code>	6	<code>readpss</code>	30	<code>strobeperiod</code>	22
<code>cmin</code>	12	<code>maxiters</code>	47	<code>relref</code>	34	<code>swapfile</code>	28
<code>compression</code>	24	<code>maxorder</code>	42	<code>restart</code>	48	<code>title</code>	46
<code>errpreset</code>	33	<code>maxperiods</code>	37	<code>save</code>	16	<code>tstab</code>	3
<code>finitediff</code>	39	<code>maxstep</code>	5	<code>saveinit</code>	25	<code>tstabmethod</code>	32
<code>fullpssvec</code>	43	<code>method</code>	31	<code>skipcount</code>	21	<code>tstart</code>	4
<code>fund</code>	2	<code>nestlvl</code>	17	<code>skipdc</code>	9	<code>write</code>	26
<code>harms</code>	13	<code>oppoint</code>	18	<code>skipstart</code>	19	<code>writefinal</code>	27
<code>harmsvec</code>	14	<code>outputtype</code>	15	<code>skipstop</code>	20	<code>writepss</code>	29
<code>highorder</code>	40	<code>period</code>	1	<code>stats</code>	44		
<code>ic</code>	8	<code>psaratio</code>	41	<code>steadyratio</code>	36		
<code>itres</code>	38	<code>readic</code>	10	<code>step</code>	7		

Periodic Transfer Function Analysis (pxf)

Description

A conventional transfer function analysis computes the transfer function from every source in the circuit to a single output. It differs from a conventional AC analysis in that the AC analysis computes the response from a single stimulus to every node in the circuit. The difference between PAC and PXF analysis are similar. The Periodic Transfer Function or PXF analysis computes the transfer functions from any source at any frequency to a single output at a single frequency. Thus, like PAC analysis, PXF analysis includes frequency conversion effects.

The PXF analysis directly computes such useful quantities as conversion efficiency (transfer function from input to output at desired frequency), image and sideband rejection (input to output at undesired frequency), and LO feed-through and power supply rejection (undesired input to output at all frequencies).

As with a PAC, PSP, and PNoise analyses, a PXF analysis must follow a PSS analysis.

Unlike other analyses in Spectre, this analysis can only sweep frequency.

Definition

Name [p] [n] pxf parameter=value ...

The optional terminals (p and n) specify the output of the circuit. If you do not give the terminals, then you must specify the output with a probe component.

Parameters

Sweep interval parameters

1	start=0	Start sweep limit.
2	stop	Stop sweep limit.
3	center	Center of sweep.
4	span=0	Sweep limit span.
5	step	Step size, linear sweep.

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6	<code>lin=50</code>	Number of steps, linear sweep.
7	<code>dec</code>	Points per decade.
8	<code>log=50</code>	Number of steps, log sweep.
9	<code>values=[...]</code>	Array of sweep values.
10	<code>sweepstype</code>	Specifies if the sweep frequency range is absolute frequency of input or if it is relative to the port harmonics. Possible values are <code>absolute</code> or <code>relative</code> .
11	<code>relharmnum=1</code>	Harmonic to which relative frequency sweep should be referenced.

Probe parameters

12	<code>probe</code>	Compute every transfer function to this probe component.
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Output parameters

13	<code>stimuli=sources</code>	Stimuli used for xf analysis. Possible values are <code>sources</code> or <code>nodes_and_terminals</code> .
14	<code>sidebands=[...]</code>	Array of relevant sidebands for the analysis.
15	<code>maxsideband=0</code>	An alternative to the <code>sidebands</code> array specification, which automatically generates the array: <code>[-maxsideband ... 0 ... +maxsideband]</code> .
16	<code>freqaxis</code>	Specifies whether the results should be output versus the input frequency, the output frequency, or the absolute value of the input frequency. Default is <code>out</code> for logarithmic frequency sweeps and <code>absin</code> otherwise. Possible values are <code>absin</code> , <code>in</code> or <code>out</code> .
17	<code>save</code>	Signals to output. Possible values are <code>all</code> , <code>lvl</code> , <code>allpub</code> , <code>lvlpub</code> , <code>selected</code> , or <code>none</code> .
18	<code>nestlvl</code>	Levels of subcircuits to output.

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Convergence parameters

- 19 `tolerance=1e-9` Relative tolerance for linear solver.
- 20 `gear_order=2` Gear order used for small-signal integration.
- 21 `solver=turbo` Solver type.
Possible values are `std` or `turbo`.
- 22 `oscsolver=turbo` Oscillator solver type.
Possible values are `std` or `turbo`.

Annotation parameters

- 23 `annotate=sweep` Degree of annotation.
Possible values are `no`, `title`, `sweep`, `status`, or `steps`.
- 24 `stats=no` Analysis statistics.
Possible values are `no` or `yes`.
- 25 `title` Analysis title.

Modulation conversion parameters

- 26 `modulated=no` Compute transfer functions/conversion between modulated sources and outputs.
Possible values are `single`, `first`, `second`, or `no`.
- 27 `outmodharmnum=1` Harmonic for the PXF output modulation.
- 28 `inmodharmvec=[...]` Harmonic list for the PXF modulated sources..
- 29 `moduppersideband=1` Index of the upper sideband included in the modulation of an output for PAC or an input for PXF.

The variable of interest at the output can be voltage or current, and its frequency is not constrained by the period of the large periodic solution. While sweeping the selected output frequency, you can select the periodic small-signal input frequencies of interest by setting either the `maxsideband` or the `sidebands` parameters. For a given set of n integer numbers representing the sidebands K_1, K_2, \dots, K_n , the input signal frequency at each sideband is computed as $f(\text{in}) = f(\text{out}) + K_i * f_{\text{fund}}(\text{pss})$, where $f(\text{out})$ represent the (possibly swept) output

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signal frequency, and `fund(pss)` represents the fundamental frequency used in the corresponding PSS analysis. Thus, when analyzing a down-converting mixer, and sweeping the IF output frequency, `Ki= +1` for the RF input represents the first upper-sideband, while `Ki=-1` for the RF input represents the first lower-sideband. By setting the `maxsideband` value to `Kmax`, all $2 * Kmax + 1$ sidebands from `-Kmax` to `+Kmax` are be selected.

The number of requested sidebands does not change substantially the simulation time. However, the `maxacfreq` of the corresponding PSS analysis should be set to guarantee that $| \max\{f(in)\} |$ is less than `maxacfreq`, otherwise the computed solution might be contaminated by aliasing effects. The PXF simulation is not executed for $| f(out) |$ greater than `maxacfreq`. Diagnostic messages are printed for those extreme cases, indicating how `maxacfreq` should be set in the PSS analysis. In the majority of the simulations, however, this is not an issue, because `maxacfreq` is never allowed to be smaller than 40x the PSS fundamental.

With PXF the frequency of the stimulus and of the response are usually different (this is an important way in which PXF differs from XF). The `freqaxis` parameter is used to specify whether the results should be output versus the input frequency (`in`), the output frequency (`out`), or the absolute value of the input frequency (`absin`).

You can specify the output with a pair of nodes or a probe component. Any component with two or more terminals can be a voltage probe. When there are more than two terminals, they are grouped in pairs; and you use the `portv` parameter to select the appropriate pair of terminals. Alternatively, you can simply specify a voltage to be the output by giving a pair of nodes on the PXF analysis statement.

Any component that naturally computes current as an internal variable can be a current probe. If the probe component computes more than one current, you use the `porti` parameter to select the appropriate current. It is an error to specify both `portv` and `porti`. If neither is specified, the probe component provides a reasonable default.

The `stimuli` parameter specifies what is used for the inputs for the transfer functions. There are two choices. `stimuli=sources` indicates that the sources present in the circuit should be used. The `xfmag` parameters provided by the sources may be used to adjust the computed gain to compensate for gains or losses in a test fixture. One can limit the number of sources in hierarchical netlists by using the `save` and `nestlvl` parameters.

`stimuli=nodes_and_terminals` indicates that all possible transfer functions should be computed.

This is useful when it is not known in advance which transfer functions are interesting. Transfer functions for nodes are computed assuming that a unit magnitude flow (current) source is connected from the node to ground. Transfer functions for terminals are computed assuming that a unit magnitude value (voltage) source is connected in series with the terminal. By default, the transfer functions from a small set of terminals are computed. If

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transfer functions from specific terminals are desired, specify the terminals in the `save` statement. You must use the `:probe` modifier (ex. `Rout:1:probe`) or specify `useprobes=yes` on the options statement. If transfer functions from all terminals are desired, specify `currents=all` and `useprobes=yes` on the options statement.

Modulated small signal measurements are possible using the Analog Artist(ADE) environment. The `modulated` option for PXF and other modulated parameters are set by Artist. PXF analyses with this option will produce results which could have limited use outside such environment. Direct Plot is configured to analyze these results and combine several wave forms to measure AM and PM transfer function from single sideband or modulated stimuli to the specified output. For details, please see the SpectreRF User Guide.

You can specify sweep limits by giving the end points or by providing the center value and the span of the sweep. Steps can be linear or logarithmic, and you can specify the number of steps or the size of each step. You can give a step size parameter (`step`, `lin`, `log`, `dec`) to determine whether the sweep is linear or logarithmic. If you do not give a step size parameter, the sweep is linear when the ratio of stop to start values is less than 10, and logarithmic when this ratio is 10 or greater. Alternatively, you may specify the particular values that the sweep parameter should take using the `values` parameter. If you give both a specific set of values and a set specified using a sweep range, the two sets are merged and collated before being used. All frequencies are in Hertz.

Parameter Index

In the following index, the number following each parameter name indicates where to find the description of that parameter.

<code>annotate</code>	23	<code>maxsideband</code>	15	<code>save</code>	17	<code>stop</code>	2
<code>center</code>	3	<code>modulated</code>	26	<code>sidebands</code>	14	<code>sweeptype</code>	10
<code>dec</code>	7	<code>moduppersideband</code>	29	<code>solver</code>	21	<code>title</code>	25
<code>freqaxis</code>	16	<code>nestlvl</code>	18	<code>span</code>	4	<code>tolerance</code>	19
<code>gear_order</code>	20	<code>oscsolver</code>	22	<code>start</code>	1	<code>values</code>	9
<code>inmodharmvec</code>	28	<code>outmodharmnum</code>	27	<code>stats</code>	24		
<code>lin</code>	6	<code>probe</code>	12	<code>step</code>	5		
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PZ Analysis (pz)

Description

The PZ analysis linearizes the circuit about the DC operating point and computes the poles and zeros of the linearized network. To compute zeros, users need to specify input sources and output voltages or currents. If no input or output are given, then only poles are computed. In case there are frequency dependent components, poles and zeros are computed by approximating those components as equivalent conductances and capacitances evaluated at 1Hz. The PZ analysis currently uses direct solver for better accuracy. Small to medium circuit size will achieve better performance.

(Note: A frequency dependent component means the capacitance or conductance equivalent representation of the component is frequency varying. Examples are transmission lines or bjts with excess phases. A linear capacitor is not a frequency dependent component.)

Spectre can perform the analysis while sweeping a parameter. The parameter can be temperature, component instance parameter, component model parameter, or netlist parameter. If changing a parameter affects the DC operating point, the operating point is recomputed on each step. You can sweep the parameter `temp` or a netlist parameter by giving the parameter name with `no dev`, or `mod` parameter. After the analysis has completed, the modified parameter returns to its original value.

Definition

Name ... pz parameter=value ...

Parameters

Probe parameters

- | | | |
|---|---------------------|--|
| 1 | <code>iprobe</code> | Input probe for zeros of the transfer function. |
| 2 | <code>oprobe</code> | Output probe for zeros of the transfer function. |

Port parameters

- | | | |
|---|--------------------|---|
| 3 | <code>portv</code> | Voltage across this <code>oprobe</code> port is output of the analysis. |
|---|--------------------|---|

Spectre Circuit Simulator Reference

Analysis Statements

4 `port i` Current through this oprobe port is output of the analysis. Should be used when oprobe is a voltage source or a current probe.

Sweep interval parameters

5 `start=0` Start sweep limit.

6 `stop` Stop sweep limit.

7 `center` Center of sweep.

8 `span=0` Sweep limit span.

9 `step` Step size, linear sweep.

10 `lin=50` Number of steps, linear sweep.

11 `dec` Points per decade.

12 `log=50` Number of steps, log sweep.

13 `values=[...]` Array of sweep values.

Sweep variable parameters

14 `dev` Device instance whose parameter value is to be swept.

15 `mod` Model whose parameter value is to be swept.

16 `param` Name of parameter to sweep.

17 `freq (Hz)` Frequency at which components will be evaluated in setting up the linearized network.

State-file parameters

18 `readns` File that contains estimate of DC solution (nodeset).

Spectre Circuit Simulator Reference

Analysis Statements

Output parameters

- 19 `oppoint=no` Should operating point information be computed, and if so, where should it be sent.
Possible values are `no`, `screen`, `logfile`, or `rawfile`.
- 20 `fmax` Maximum pole and zero frequency value to filter out spurious poles and zeros. This parameter is passed to `psf` outputs for plotting filtering.
- 21 `zeroonly=no` If set, only zeros are requested.
Possible values are `no` or `yes`.

Convergence parameters

- 22 `prevoppoint=no` Use operating point computed on the previous analysis.
Possible values are `no` or `yes`.
- 23 `restart=yes` Restart the DC solution from scratch if any condition has changed. If not, use the previous solution as initial guess.
Possible values are `no` or `yes`.

Annotation parameters

- 24 `stats=no` Analysis statistics.
Possible values are `no` or `yes`.
- 25 `annotate=sweep` Degree of annotation.
Possible values are `no`, `title`, `sweep`, `status`, or `steps`.
- 26 `title` Analysis title.

Examples

```
mypz pz
```

Pole analysis will be performed.

```
mypz2 (n1 n2) pz iprobe=VIN
```

- Input is `VIN`, output is voltage difference between nodes `n1` and `n2`.
- Both pole and zero analyses will be performed.

Spectre Circuit Simulator Reference Analysis Statements

```
mypz3 (n1 n2) pz iprobe=I1
```

- Input is I1, output is voltage difference between n1 and n2.
- Both pole and zero analyses will be performed.

```
mypz4 pz iprobe=VIN oprobe=IP1 porti=1
```

- Input is VIN, output is current through IP1, where IP1 is an iprobe.
- Both pole and zero analyses will be performed.

```
mypz5 pz iprobe=VIN oprobe=V3 porti=1
```

- Input is VIN, output is current through voltage source V3.
- Both pole and zero analyses will be performed.

```
mypz6 pz iprobe=VIN oprobe=R3 portv=1
```

- Input is VIN, output is the voltage across the resistor R3.
- Both pole and zero analyses will be performed.

```
mypz7 (n1 n2) pz iprobe=I1 param=temp start=25 stop=100 step=25
```

Sweep temperature from 25 C to 100 C with increment of 25 C.

```
parameters rval=2.0  
R2 3 4 resistor r=rval  
...  
sweep1 sweep param=rval start=1 stop=10 step=1 {  
  mypz8 (n1 n2) iprobe=VIN  
}
```

External sweep parameter rval from 1 to 10 with increment of 1.

Note: porti allows users to select a current associated with a specific device given in oprobe as an output. This device, however, has to have its terminal currents as network variables. Thus, to avoid confusion, porti should be used with voltage sources and current probes, and other components that have voltage-defined branches exclusively.

Parameter Index

In the following index, the number following each parameter name indicates where to find the description of that parameter.

annotate	25	lin	10	portv	3	step	9
center	7	log	12	prevoppoint	22	stop	6
dec	11	mod	15	readns	18	title	26

Spectre Circuit Simulator Reference Analysis Statements

dev	14	oppoint	19	restart	23	values	13
fmax	20	oprobe	2	span	8	zeroonly	21
freq	17	param	16	start	5		
iprobe	1	porti	4	stats	24		

Quasi-Periodic AC Analysis (qpac)

Description

The quasi periodic AC (QPAC) analysis is used to compute transfer functions for circuits that exhibit multitone frequency translation. Such circuits include mixers, switched-capacitor filters, samplers, phase-locked loops, and the like. It is a small-signal analysis like AC analysis, except the circuit is first linearized about a quasiperiodically varying operating point as opposed to a simple DC operating point. Linearizing about a quasiperiodically time-varying operating point allows transfer-functions that include frequency translation, whereas simply linearizing about a DC operating point could not because linear time-invariant circuits do not exhibit frequency translation. Also, the frequency of the sinusoidal stimulus is not constrained by the period of the large periodic solution.

Computing the small-signal response of a quasiperiodically varying circuit is a two step process. First, the small stimulus is ignored and the quasiperiodic steady-state response of the circuit to possibly large periodic stimuli is computed using QPSS analysis. As a normal part of the QPSS analysis, the quasiperiodically time-varying representation of the circuit is computed and saved for later use. The second step is to apply the small stimulus to the periodically varying linear representation to compute the small signal response. This is done using the QPAC analysis.

A QPAC analysis cannot be used alone, it must follow a QPSS analysis. However, any number of quasiperiodic small-signal analyses such as QPAC, QPSP, QPXF, QPNOISE, can follow a QPSS analysis.

Unlike other analyses in Spectre, this analysis can only sweep frequency.

Definition

Name `qpac parameter=value ...`

Spectre Circuit Simulator Reference Analysis Statements

Parameters

Sweep interval parameters

1	<code>start=0</code>	Start sweep limit.
2	<code>stop</code>	Stop sweep limit.
3	<code>center</code>	Center of sweep.
4	<code>span=0</code>	Sweep limit span.
5	<code>step</code>	Step size, linear sweep.
6	<code>lin=50</code>	Number of steps, linear sweep.
7	<code>dec</code>	Points per decade.
8	<code>log=50</code>	Number of steps, log sweep.
9	<code>values=[...]</code>	Array of sweep values.
10	<code>sweeptype</code>	Specifies if the sweep frequency range is an absolute frequency, i.e. actual frequency; or if it is relative to the "relharmvec" sideband frequency. In QPSP, relative means relative to the input port frequency. Possible values are <code>absolute</code> or <code>relative</code> .
11	<code>relharmvec=[...]</code>	Sideband - vector of QPSS harmonics - to which relative frequency sweep should be referenced.

Output parameters

12	<code>sidevec=[...]</code>	Array of relevant sidebands for the analysis.
13	<code>clockmaxharm=0</code>	An alternative to the <code>sidevec</code> array specification, which automatically generates the array: <code>[-clockmaxharm ... 0 ... +clockmaxharms][maxharms(QPSS)[2]...0...maxharms(QPSS)[2]] [...]</code> .

Spectre Circuit Simulator Reference

Analysis Statements

- 14 `freqaxis` Specifies whether the results should be output versus the input frequency, the output frequency, or the absolute value of the output frequency. Default is `in` for logarithmic frequency sweeps and `absout` otherwise.
Possible values are `absout`, `out` or `in`.
- 15 `save` Signals to output.
Possible values are `all`, `lvl`, `allpub`, `lvlpub`, `selected`, or `none`.
- 16 `nestlvl` Levels of subcircuits to output.

Convergence parameters

- 17 `tolerance=1e-9` Relative tolerance for linear solver.
- 18 `gear_order=2` Gear order used for small-signal integration, 1 or 2.
- 19 `solver=turbo` Solver type.
Possible values are `std` or `turbo`.

Annotation parameters

- 20 `annotate=sweep` Degree of annotation.
Possible values are `no`, `title`, `sweep`, `status`, or `steps`.
- 21 `stats=no` Analysis statistics.
Possible values are `no` or `yes`.
- 22 `title` Analysis title.

User can select the set of periodic small-signal output frequencies of interest by setting either the `clockmaxharm` or the `sidevec` parameters. Sidebands are vectors in QPAC. Assume we have one large tone and one moderate tone in QPSS. A sideband K1 will be represented as $[K1_1 \ K1_2]$. Corresponding frequency is

$$K1_1 * \text{fund}(\text{large tone of QPSS}) + K1_2 * \text{fund}(\text{moderate tone of QPSS})$$

We assume that there are L large and moderate tones in QPSS analysis and a given set of n integer vectors representing the sidebands

Spectre Circuit Simulator Reference

Analysis Statements

$K1 = \{ K1_1, \dots, K1_j, \dots, K1_L \}$, $K2$, ... Kn . The output frequency at each sideband is computed as

$$f(\text{out}) = f(\text{in}) + \text{SUM}_{j=1_to_L} \{ K_{i_j} * \text{fund}_j(\text{qpss}) \},$$

where $f(\text{in})$ represents the (possibly swept) input frequency, and $\text{fund}_j(\text{qpss})$ represents the fundamental frequency used in the corresponding QPSS analysis. Thus, when analyzing a down-converting mixer, while sweeping the RF input frequency, the most relevant sideband for IF output is $\{-1, 0\}$. When simulating an up-converting mixer, while sweeping IF input frequency, the most relevant sideband for RF output is $\{1, 0\}$. User would enter `sidevec` as a sequence of integer numbers, separated by spaces. The set of vectors $\{1\ 1\ 0\}$ $\{1\ -1\ 0\}$ $\{1\ 1\ 1\}$ becomes `sidevec=[1 1 0 1 -1 0 1 1 1]`. For `clockmaxharm`, only the large tone - first fundamental will be affected by this entry, all the rest - moderate tones - will be limited by `maxharms`, specified for a QPSS analysis. Given `maxharms=[k1max k2max ... knmax]` in QPSS and `clockmaxharm=Kmax` all $(2 * K_{\text{max}} + 1) * (2 * k_{2\text{max}} + 1) * (2 * k_{3\text{max}} + 1) * \dots * (2 * k_{n\text{max}} + 1)$ sidebands are generated.

The number of requested sidebands changes substantially the simulation time.

With QPAC the frequency of the stimulus and of the response are usually different (this is an important way in which QPAC differs from AC). The `freqaxis` parameter is used to specify whether the results should be output versus the input frequency (`in`), the output frequency (`out`), or the absolute value of the output frequency (`absout`).

You can specify sweep limits by giving the end points or by providing the center value and the span of the sweep. Steps can be linear or logarithmic, and you can specify the number of steps or the size of each step. You can give a step size parameter (`step`, `lin`, `log`, `dec`) to determine whether the sweep is linear or logarithmic. If you do not give a step size parameter, the sweep is linear when the ratio of stop to start values is less than 10, and logarithmic when this ratio is 10 or greater. Alternatively, you may specify the particular values that the sweep parameter should take using the `values` parameter. If you give both a specific set of values and a set specified using a sweep range, the two sets are merged and collated before being used. All frequencies are in Hertz.

Parameter Index

In the following index, the number following each parameter name indicates where to find the description of that parameter.

<code>annotate</code>	20	<code>lin</code>	6	<code>solver</code>	19	<code>sweepstype</code>	10
<code>center</code>	3	<code>log</code>	8	<code>span</code>	4	<code>title</code>	22

Spectre Circuit Simulator Reference Analysis Statements

<code>clockmaxharm</code>	<code>13</code>	<code>nestlvl</code>	<code>16</code>	<code>start</code>	<code>1</code>	<code>tolerance</code>	<code>17</code>
<code>dec</code>	<code>7</code>	<code>relharmvec</code>	<code>11</code>	<code>stats</code>	<code>21</code>	<code>values</code>	<code>9</code>
<code>freqaxis</code>	<code>14</code>	<code>save</code>	<code>15</code>	<code>step</code>	<code>5</code>		
<code>gear_order</code>	<code>18</code>	<code>sidevec</code>	<code>12</code>	<code>stop</code>	<code>2</code>		

Quasi-Periodic Noise Analysis (qpnoise)

Description

The Quasi-Periodic Noise, or QPNOISE analysis is similar to the conventional noise analysis, except that it includes frequency conversion and intermodulation effects. Hence is it useful for predicting the noise behavior of mixers, switched-capacitor filters, and other periodically or quasi-periodically driven circuits.

QPNOISE analysis linearizes the circuit about the quasi-periodic operating point computed in the prerequisite QPSS analysis. It is the quasiperiodically time-varying nature of the linearized circuit that accounts for the frequency conversion and intermodulation. In addition, the affect of a quasi-periodically time-varying bias point on the noise generated by the various components in the circuit is also included.

The time-average of the noise at the output of the circuit is computed in the form of a spectral density versus frequency. The output of the circuit is specified with either a pair of nodes or a probe component. To specify the output of a circuit with a probe, specify it using the `oprobe` parameter. If the output is voltage (or potential), choose a `resistor` or a `port` as the output probe. If the output is current (or flow), choose a `vsource` or `iprobe` as the output probe.

If the input-referred noise is desired, specify the input source using the `iprobe` parameter. Currently, only a `vsource`, an `isource`, or a `port` may be used as an input probe. If the input source is noisy, as is a `port`, the noise analysis will compute the noise factor (F) and noise figure (NF). To match the IEEE definition of noise figure, the input probe must be a port with no excess noise and its `noisetemp` must be set to 16.85C (290K). In addition, the output load must be a `resistor` or `port` and must be identified as the `oprobe`.

The reference sideband (`refsideband`) specifies which conversion gain is used when computing input-referred noise, noise factor, and noise figure. The reference sideband satisfies:

$$|f(\text{input})| = |f(\text{out}) + \text{refsideband frequency shift}|.$$

Spectre Circuit Simulator Reference

Analysis Statements

The reference sideband option (`refsidebandoption`) specifies whether to consider the input at the frequency or the input at the individual quasi-periodic sideband specified. Note that Different sidebands can lead to the same frequency.

Sidebands are vectors in QPNOISE. Assume we have one large tone and one moderate tone in QPSS. A sideband K_i will be a vector $[K_{i_1} K_{i_2}]$. It gives the frequency at

$$K_{i_1} * \text{fund}(\text{large tone of QPSS}) + K_{i_2} * \text{fund}(\text{moderate tone of QPSS})$$

Use `refsideband=[0 0 ...]` when the input and output of the circuit are at the same frequency (such as with amplifiers and filters).

The noise analysis always computes the total noise at the output, which includes contributions from the input source and the output load. The amount of the output noise that is attributable to each noise source in the circuit is also computed and output individually. If the input source is identified (using `iprobe`) and is a `vsource` or `isourec`, the input-referred noise is computed, which includes the noise from the input source itself. Finally, if the input source is identified (using `iprobe`) and is noisy, as is the case with ports, the noise factor and noise figure are computed. Thus if

N_o = total output noise

N_s = noise at the output due to the input probe (the source)

N_{si} = noise at the output due to the image harmonic at the source

N_{so} = noise at the output due to harmonics other than input at the source

N_l = noise at the output due to the output probe (the load)

IRN = input referred noise

G = gain of the circuit

F = noise factor

NF = noise figure

F_{dsb} = double sideband noise factor

NF_{dsb} = double sideband noise figure

F_{ieee} = IEEE single sideband noise factor

NF_{ieee} = IEEE single sideband noise figure

Spectre Circuit Simulator Reference

Analysis Statements

then,

$$\text{IRN} = \sqrt{\text{No}^2 / \text{G}^2}$$

$$\text{F} = (\text{No}^2 - \text{NI}^2) / \text{Ns}^2$$

$$\text{NF} = 10 * \log_{10}(\text{F})$$

$$\text{Fdsb} = (\text{No}^2 - \text{NI}^2) / (\text{Ns}^2 + \text{Nsi}^2)$$

$$\text{NFdsb} = 10 * \log_{10}(\text{Fdsb})$$

$$\text{Fieeee} = (\text{No}^2 - \text{NI}^2 - \text{Nso}^2) / \text{Ns}^2$$

$$\text{NFieeee} = 10 * \log_{10}(\text{Fieeee}).$$

When the results are output, No is named `out`, IRN is named `in`, G is named `gain`, F, NF, Fdsb, NFdsb, Fieeee, and NFieeee are named `F`, `NF`, `Fdsb`, `NFdsb`, `Fieeee`, and `NFieeee` respectively.

The computation of gain and IRN in QPNOISE assumes that the circuit under test is impedance-matched to the input source. This can introduce inaccuracy into the gain and IRN computation.

Unlike other analyses in Spectre, this analysis can only sweep frequency.

Definition

Name [p] [n] `qpnoise parameter=value ...`

The optional terminals (p and n) specify the output of the circuit. If you do not give the terminals, then you must specify the output with a probe component.

Parameters

Sweep interval parameters

- | | | |
|---|----------------------|--------------------|
| 1 | <code>start=0</code> | Start sweep limit. |
| 2 | <code>stop</code> | Stop sweep limit. |
| 3 | <code>center</code> | Center of sweep. |

Spectre Circuit Simulator Reference

Analysis Statements

4	<code>span=0</code>	Sweep limit span.
5	<code>step</code>	Step size, linear sweep.
6	<code>lin=50</code>	Number of steps, linear sweep.
7	<code>dec</code>	Points per decade.
8	<code>log=50</code>	Number of steps, log sweep.
9	<code>values=[...]</code>	Array of sweep values.
10	<code>sweepstype</code>	Specifies if the sweep frequency range is an absolute frequency, i.e. actual frequency; or if it is relative to the "relharmvec" sideband frequency. In QPSP, relative means relative to the input port frequency. Possible values are <code>absolute</code> or <code>relative</code> .
11	<code>relharmvec=[...]</code>	Sideband - vector of QPSS harmonics - to which relative frequency sweep should be referenced.

Probe parameters

12	<code>oprobe</code>	Compute total noise at the output defined by this component.
13	<code>iprobe</code>	Refer the output noise to this component.
14	<code>refsideband=[...]</code>	Conversion gain associated with this sideband is used when computing input-referred noise or noise figure.
15	<code>refsidebandoption=individual</code>	Whether to view the sideband as a specification of a frequency or a specification of an individual sideband. Possible values are <code>freq</code> or <code>individual</code> .

Output parameters

16	<code>clockmaxharm=7</code>	Maximum large tone harmonics included when computing noise either up-converted or down-converted to the output by that large signal..
----	-----------------------------	---

Spectre Circuit Simulator Reference

Analysis Statements

- 17 `sidevec=[...]` Array of relevant sidebands for the analysis.
- 18 `save` Signals to output.
Possible values are `all`, `lvl`, `allpub`, `lvlpub`, `selected`, or `none`.
- 19 `nestlvl` Levels of subcircuits to output.
- 20 `saveallsidebands=no` Save noise contributors by sideband.
Possible values are `no` or `yes`.

Convergence parameters

- 21 `tolerance=1e-9` Relative tolerance for linear solver.
- 22 `gear_order=2` Gear order used for small-signal integration, 1 or 2.
- 23 `solver=turbo` Solver type.
Possible values are `std` or `turbo`.

Annotation parameters

- 24 `annotate=sweep` Degree of annotation.
Possible values are `no`, `title`, `sweep`, `status`, or `steps`.
- 25 `stats=no` Analysis statistics.
Possible values are `no` or `yes`.
- 26 `title` Analysis title.

In practice, noise can mix with each of the harmonics of the quasi-periodic drive signal applied in the QPSS analysis and end up at the output frequency. The QPNOISE analysis only includes the noise that mixes with a finite set of harmonics that are specified using the `clockmaxharm` and `sidevec` parameters. Sidebands are vectors in quasi-periodic analyses. For one large tone and one moderate tone in QPSS, a sideband K1 will be represented as [K1_1 K1_2]. Corresponding frequency shift is

$$K1_1 * \text{fund}(\text{large tone of QPSS}) + K1_2 * \text{fund}(\text{moderate tone of QPSS})$$

We assume that there are L large and moderate tones in QPSS analysis and a given set of n integer vectors representing the sidebands

Spectre Circuit Simulator Reference Analysis Statements

$$K1 = \{ K1_1, \dots, K1_j, \dots, K1_L \},$$

K2, ... Kn.

If K_i represents sideband i , then

$$f(\text{noise_source}) = f(\text{out}) + \text{SUM}_{j=1_to_L} \{ K_{i_j} * \text{fund}_{j}(\text{qpss}) \},$$

The `clockmaxharm` parameter only affects clock frequency. It can be less or more than `maxharms[1]` in QPSS. Moderate tones are limited by `maxharms` specified in QPSS. If selected sidebands are specified using the `sidevec` parameter, then only those are included in the calculation. Care should be taken when specifying the `sidevec` or `clockmaxharm` QPNOISE and `maxharms` in QPSS. Noise results will be in error if you do not include the sidebands that contribute significant noise to the output.

The number of requested sidebands will change substantially the simulation time.

You can specify sweep limits by giving the end points or by providing the center value and the span of the sweep. Steps can be linear or logarithmic, and you can specify the number of steps or the size of each step. You can give a step size parameter (`step`, `lin`, `log`, `dec`) to determine whether the sweep is linear or logarithmic. If you do not give a step size parameter, the sweep is linear when the ratio of stop to start values is less than 10, and logarithmic when this ratio is 10 or greater. Alternatively, you may specify the particular values that the sweep parameter should take using the `values` parameter. If you give both a specific set of values and a set specified using a sweep range, the two sets are merged and collated before being used. All frequencies are in Hertz.

Parameter Index

In the following index, the number following each parameter name indicates where to find the description of that parameter.

<code>annotate</code> 24	<code>log</code> 8	<code>saveallsidebands</code> 20	<code>stop</code> 2
<code>center</code> 3	<code>nestlvl</code> 19	<code>sidevec</code> 17	<code>sweepstype</code> 10
<code>clockmaxharm</code> 16	<code>oprobe</code> 12	<code>solver</code> 23	<code>title</code> 26
<code>dec</code> 7	<code>refsideband</code> 14	<code>span</code> 4	<code>tolerance</code> 21
<code>gear_order</code> 22	<code>refsidebandoption</code> 15	<code>start</code> 1	<code>values</code> 9
<code>iprobe</code> 13	<code>relharmvec</code> 11	<code>stats</code> 25	

lin 6 save 18 step 5

Quasi-Periodic S-Parameter Analysis (qpsp)

Description

The quasi-periodic SP (QPSP) analysis is used to compute scattering and noise parameters for n-port circuits that exhibit frequency translation. Such circuits include mixers, switched-capacitor filters, samplers, phase-locked loops, and the like. It is a small-signal analysis like SP analysis, except, as in QPAC and QPXF, the circuit is first linearized about a quasiperiodically varying operating point as opposed to a simple DC operating point. Linearizing about a quasiperiodically time-varying operating point allows the computation of S-parameters between circuit ports that convert signals from one frequency band to another. QPSP can also calculate noise parameters in frequency-converting circuits. QPSP computes noise figure (both single-sideband and double-sideband), input referred noise, equivalent noise parameters, and noise correlation matrices. As in QPNOISE, but unlike SP, the noise features of the QPSP analysis include noise folding effects due to the periodically time-varying nature of the circuit.

Computing the n-port S-parameters and noise parameters of a quasiperiodically varying circuit is a two step process. First, the small stimulus is ignored and the quasiperiodic steady-state response of the circuit to possibly large periodic stimulus is computed using QPSS analysis. As a normal part of the QPSS analysis, the quasiperiodically time-varying representation of the circuit is computed and saved for later use. The second step is applying small-signal excitations to compute the n-port S-parameters and noise parameters. This is done using the QPSP analysis. A QPSP analysis cannot be used alone, it must follow a QPSS analysis. However, any number of periodic small-signal analyses such as QPAC, QPSP, QPXF, QPNOISE, can follow a single QPSS analysis.

Unlike other analyses in Spectre, this analysis can only sweep frequency.

Definition

Name `qpsp parameter=value ...`

Parameters

Sweep interval parameters

1 `start=0` Start sweep limit.

Spectre Circuit Simulator Reference

Analysis Statements

2	<code>stop</code>	Stop sweep limit.
3	<code>center</code>	Center of sweep.
4	<code>span=0</code>	Sweep limit span.
5	<code>step</code>	Step size, linear sweep.
6	<code>lin=50</code>	Number of steps, linear sweep.
7	<code>dec</code>	Points per decade.
8	<code>log=50</code>	Number of steps, log sweep.
9	<code>values=[...]</code>	Array of sweep values.
10	<code>sweepstype</code>	Specifies if the sweep frequency range is an absolute frequency, i.e. actual frequency; or if it is relative to the "relharmvec" sideband frequency. In QPSP, relative means relative to the input port frequency. Possible values are <code>absolute</code> or <code>relative</code> .

Port parameters

11	<code>ports=[...]</code>	List of active ports. Ports are numbered in the order given. For purposes of noise figure computation, the input is considered port 1 and the output is port 2.
12	<code>portharmsvec=[...]</code>	List of the reference sidebands for the specified list of ports. Must have a one-to-one correspondence with the ports vector.
13	<code>harmsvec=[...]</code>	List of sidebands, in addition to ones associated with specific ports by <code>portharmsvec</code> , that are active. Call them secondary.

Output parameters

14	<code>freqaxis</code>	Specifies whether the results should be output versus the input port frequency, the output port frequency, or the absolute value of the input frequency. Default is <code>in</code> . Possible values are <code>absin</code> , <code>in</code> or <code>out</code> .
----	-----------------------	---

Spectre Circuit Simulator Reference

Analysis Statements

Noise parameters

15 `donoise=yes` Perform noise analysis. If `oprobe` is specified as a valid port, this is set to `yes`, and a detailed noise output is generated. Possible values are `no` or `yes`.

Probe parameters

16 `clockmaxharm=7` Maximum large tone harmonics included when computing noise either up-converted or down-converted to the output by that large signal..

Convergence parameters

17 `tolerance=1e-9` Relative tolerance for linear solver.

18 `gear_order=2` Gear order used for small-signal integration, 1 or 2.

19 `solver=turbo` Solver type.
Possible values are `std` or `turbo`.

Annotation parameters

20 `annotate=sweep` Degree of annotation.
Possible values are `no`, `title`, `sweep`, `status`, or `steps`.

21 `stats=no` Analysis statistics.
Possible values are `no` or `yes`.

22 `title` Analysis title.

To specify the QPSP analysis the port and sideband combinations must be specified. You can select the ports of interest by setting the `port` parameter and the set of periodic small-signal output frequencies of interest by setting `port harmsvec` or `harmsvec` parameters. Sidebands are vectors in QPSP. Assume we have one large tone and one moderate tone in QPSS. A sideband K1 will be represented as [K1_1 K1_2]. Corresponding frequency is

$$K1_1 * \text{fund}(\text{large tone of QPSS}) + K1_2 * \text{fund}(\text{moderate tone of QPSS}) = \text{SUM}_{j=1_to_L}\{Ki_j * \text{fund}_j(\text{qpss})\}$$

Spectre Circuit Simulator Reference

Analysis Statements

We assume that there are L (1 large plus $L-1$ moderate) tones in QPSS analysis and a given set of n integer vectors representing the sidebands

$$K1 = \{ K1_1, \dots, K1_j, \dots, K1_L \}, K2, \dots, Kn.$$

If we specify the relative frequency then the scattering parameters at each port are computed at the frequencies

$$f(\text{scattered}) = f(\text{rel}) + \text{SUM}_{j=1_to_L} \{ K1_j * \text{fund}_j(\text{qpss}) \},$$

where $f(\text{rel})$ represents the relative frequency of a signal incident on a port, $f(\text{scattered})$ represents the frequency to which the relevant scattering parameter represents the conversion, and $\text{fund}_j(\text{qpss})$ represents the fundamental frequency used in the corresponding QPSS analysis.

In analysis of a down-converting mixer with a blocker and the signal in the upper sideband, we sweep the input frequency of the signal coming into RF port. The most relevant sideband for this input is $K1 = \{ 1, 0 \}$ - and for IF output $K2 = \{ 0, 0 \}$. Hence we can associate $K1 = \{ 1, 0 \}$ with the RF port and $K2 = \{ 0, 0 \}$ with the IF port. $S21$ will represent the transmission of signal from the RF to IF, and $S11$ the reflection of signal back to the RF port. If the signal was in the lower sideband, then a choice of $K1 = \{ -1, 0 \}$ would be more appropriate.

Either `port harmsvec` or `harmsvec` parameters can be used to specify the sidebands of the interest. If `port harmsvec` is given, the sidebands must be in one-to-one correspondence with the ports, with each sideband associated with a single port. If sidebands are specified in the optional `harmsvec` parameter, then all possible frequency-translating scattering parameters associated with the specified sidebands on each port are computed.

With QPSP the frequency of the input and of the response are usually different (this is an important way in which QPSP differs from SP). Because the QPSP computation involves inputs and outputs at frequencies that are relative to multiple sidebands, the `freqaxis` and `sweepstype` parameters behave somewhat differently in QPSP than in QPAC and QPXF.

The `sweepstype` parameter controls the way the frequencies in the QPSP analysis are swept. Specifying a `relative` sweep indicates to sweep relative to the port sideband (not the QPSS fundamental) and an `absolute` sweep is a sweep of the absolute input source frequency. For example, with a QPSS fundamentals of 1000MHz (LO) and 966MHz (blocker in RF channel), `port harmsvec` could be set to `[0 1 -1 1]` to examine a downconverting mixer. Lets set `sweepstype=relative` and a sweep range of $f(\text{rel}) = -10\text{MHz} \leftrightarrow 10\text{MHz}$. Then $S21$ would represent the strength of the signal transmitted from the input port in the range 956->976MHz to the output port to the frequencies 24->44MHz. Using `sweepstype=absolute` and sweeping the frequency from 966->976MHz would calculate the same quantities, since $f(\text{abs}) = 956 \leftrightarrow 976\text{MHz}$, and $f(\text{rel}) = f(\text{abs}) - (K1_1 * \text{fund}_1(\text{qpss}) + K1_2 * \text{fund}_2(\text{qpss})) = -10\text{MHz} \leftrightarrow 10\text{MHz}$, because $K1_1 = 0$, $K1_2 = 1$ and $\text{fund}_1(\text{qpss}) = 1000\text{MHz}$, $\text{fund}_2(\text{qpss}) = 966\text{MHz}$.

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The `freqaxis` parameter is used to specify whether the results should be output versus the scattered frequency at the input port (`in`), the scattered frequency at the output port (`out`), or the absolute value of the frequency swept at the input port (`absin`).

An increase in the number of requested ports will increase the simulation time substantially. The same will happen if we increase the number of sidebands to be included into the noise computations.

QPSP analysis also computes noise figures, equivalent noise sources, and noise parameters. The noise computation, which is skipped only when `donoise=no`, requires additional simulation time. If

`No` = total output noise at frequency `f`

`Ns` = noise at the output due to the input probe (the source)

`Nsi` = noise at the output due to the image harmonic at the source

`Nso` = noise at the output due to harmonics other than input at the source

`NI` = noise at the output due to the output probe (the load)

`IRN` = input referred noise

`G` = gain of the circuit

`F` = noise factor (single side band)

`NF` = noise figure (single side band)

`Fdsb` = double sideband noise factor

`NFdsb` = double sideband noise figure

`Fieee` = IEEE single sideband noise factor

`NFieee` = IEEE single sideband noise figure

then,

$$IRN = \sqrt{No^2 / G^2}$$

$$F = (No^2 - NI^2) / Ns^2$$

$$NF = 10 * \log_{10}(F)$$

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$$F_{dsb} = (N_o^2 - N_I^2)/(N_s^2 + N_{si}^2)$$

$$NF_{dsb} = 10 \cdot \log_{10}(F_{dsb})$$

$$F_{ieee} = (N_o^2 - N_I^2 - N_{so}^2)/N_s^2$$

$$NF_{ieee} = 10 \cdot \log_{10}(F_{ieee}).$$

When the results are output, IRN is named `in`, G is named `gain`, F, NF, F_{dsb}, NF_{dsb}, F_{ieee}, and NF_{ieee} are named `F`, `NF`, `Fdsb`, `NFdsb`, `Fieee`, and `NFieee` respectively. Note that the gain computed by QPSP is the voltage gain from the actual circuit input to the circuit output, not the gain from the internal port voltage source to the output.

To ensure accurate noise calculations, the `clockmaxharm` parameters must be set to include the relevant noise folding effects. `clockmaxharm` is only relevant to the noise computation features of QPSP.

You can specify sweep limits by giving the end points or by providing the center value and the span of the sweep. Steps can be linear or logarithmic, and you can specify the number of steps or the size of each step. You can give a step size parameter (`step`, `lin`, `log`, `dec`) to determine whether the sweep is linear or logarithmic. If you do not give a step size parameter, the sweep is linear when the ratio of stop to start values is less than 10, and logarithmic when this ratio is 10 or greater. Alternatively, you may specify the particular values that the sweep parameter should take using the `values` parameter. If you give both a specific set of values and a set specified using a sweep range, the two sets are merged and collated before being used. All frequencies are in Hertz.

Parameter Index

In the following index, the number following each parameter name indicates where to find the description of that parameter.

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<code>clockmaxharm</code>	16	<code>lin</code>	6	<code>start</code>	1	<code>tolerance</code>	17
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Quasi-Periodic Steady State Analysis (qpss)

Description

This analysis computes the quasi-periodic steady-state (QPSS) response of a circuit that operates on multiple time scales. A quasi-periodic signal has dynamics in multiple fundamental frequencies. Closely spaced or incommensurate fundamentals cannot be resolved by PSS efficiently. QPSS allows you to compute a circuit responses to several moderately large input signals in addition to a strongly nonlinear tone which represents a LO or clock signal. A typical example is the intermodulation distortion measurements of a mixer with two closely spaced moderate input signals. QPSS treats one particular input signal (usually the one that causes the most nonlinearity or the largest response) as the large signal, and the others as moderate signals.

An initial transient analysis is carried out by first suppressing all moderate input signals. Then, a number of (at least 2) stabilizing iterations with all signals activated is run. They are followed by the shooting Newton method. QPSS employs the Mixed Frequency Time (MFT) algorithm extended to multiple fundamental frequencies. For details of MFT algorithm, please see *Steady-State Methods for Simulating Analog and Microwave Circuits*, by K. S. Kundert, J. K. White, and A. Sangiovanni-Vincentelli, Kluwer, Boston, 1990.

Like PSS, QPSS uses the shooting Newton method as its backbone. However, instead of doing a single transient integration, each Newton iteration does a number of transient integrations of one large signal period.

Each of the integrations differs by a phase-shift in each moderate input signal. The number of integrations is determined by the numbers of harmonics of moderate fundamentals specified by the user. Given $\text{maxharms}=[k_1 k_2 \dots k_n]$, the total number of integrations is $(2k_1+1)(2k_2+1)\dots(2k_n+1)$. As one consequence, the efficiency of the algorithm depends significantly on the number of harmonics required to model the responses of moderate fundamentals. As another consequence, the number of harmonics of the large fundamental does not significantly affect the efficiency of the shooting algorithm. The boundary conditions of a shooting interval are such that the time domain integrations are consistent with a frequency domain transformation with a shift of one large signal period.

QPSS inherits a majority of PSS parameters. A few new parameters are added. The most important ones are `funds` and `maxharms`. They replace PSS parameters, `fund` (or `period`) and `harms`, respectively. The `funds` parameter accepts a list of names of fundamentals that are present in the sources. These names are specified in the sources by parameter `fundname`. The first fundamental is considered as the large signal. A few heuristics can be used for picking the large fundamental.

(1) Pick the one which is not a sinusoidal.

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(2) Pick the one which causes the most nonlinearity.

(3) Pick the one which causes the largest response.

The `maxharms` parameter accepts a list of numbers of harmonics that are required to sufficiently model responses due to different fundamentals.

Definition

Name `qpss parameter=value ...`

Parameters

Distortion fundamental parameters

- | | | |
|---|-----------------------------|--|
| 1 | <code>funds=[...]</code> | Array of fundamental frequency names for fundamentals to use in analysis. |
| 2 | <code>maxharms=[...]</code> | Array of number of harmonics of each fundamental to consider for each fundamental. |

Simulation interval parameters

- | | | |
|---|---------------------------|--|
| 3 | <code>tstab=0.0 s</code> | Extra stabilization time after the onset of periodicity for independent sources. |
| 4 | <code>tstart=0.0 s</code> | Initial transient analysis start time. |

Time-step parameters

- | | | |
|---|----------------------------------|--|
| 5 | <code>maxstep (s)</code> | Maximum time step. Default derived from <code>errpreset</code> . |
| 6 | <code>step=0.001 period s</code> | Minimum time step that would be used solely to maintain the aesthetics of the results. |

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Initial-condition parameters

- 7 `ic=all` What should be used to set initial condition.
Possible values are `dc`, `node`, `dev`, or `all`.
- 8 `skipdc=no` If yes, there will be no dc analysis for transient.
Possible values are `no`, `yes` or `sigrampup`.
- 9 `readic` File that contains initial condition.

Convergence parameters

- 10 `readns` File that contains estimate of initial transient solution.
- 11 `cmin=0 F` Minimum capacitance from each node to ground.

Output parameters

- 12 `save` Signals to output.
Possible values are `all`, `lvl`, `allpub`, `lvlpub`, `selected`, or `none`.
- 13 `nestlvl` Levels of subcircuits to output.
- 14 `oppoint=no` Should operating point information be computed for initial timestep, and if so, where should it be sent.
Possible values are `no`, `screen`, `logfile`, or `rawfile`.
- 15 `skipstart=starttime s` The time to start skipping output data.
- 16 `skipstop=stoptime s` The time to stop skipping output data.
- 17 `skipcount` Save only one of every skipcount points.
- 18 `strobeperiod (s)` The output strobe interval (in seconds of transient time).
- 19 `strobedelay=0 s` The delay (phase shift) between the skipstart time and the first strobe point.

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- 20 `compression=no` Do data compression on output. See full description below. Possible values are `no` or `yes`.
- 21 `saveinit=no` If set, the waveforms for the initial transient before steady state are saved. Possible values are `no` or `yes`.

State-file parameters

- 22 `write` File to which initial transient solution (before steady-state) is to be written.
- 23 `writefinal` File to which final transient solution in steady-state is to be written.
- 24 `swapfile` Temporary file that holds steady-state information. Tells Spectre to use a regular file rather than virtual memory to hold the periodic operating point. Use this option if Spectre complains about not having enough memory to complete this analysis.

Integration method parameters

- 25 `method` Integration method. Default derived from `errpreset`. Possible values are `euler`, `trap`, `traponly`, `gear2`, or `gear2only`.

Accuracy parameters

- 26 `errpreset` Selects a reasonable collection of parameter settings. Possible values are `liberal`, `moderate` or `conservative`.
- 27 `relref` Reference used for the relative convergence criteria. Default derived from `errpreset`. Possible values are `pointlocal`, `alllocal`, `sigglobal`, or `allglobal`.
- 28 `lteratio` Ratio used to compute LTE tolerances from Newton tolerance. Default derived from `errpreset`.

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- 29 `steadyratio` Ratio used to compute steady state tolerances from LTE tolerance. Default derived from `errpreset`.
- 30 `maxperiods` Maximum number of simulated periods to reach steady-state.
- 31 `itres=1e-4` Relative tolerance for linear solver.
- 32 `finitediff` Options for finite difference method refinement after quasi-periodic shooting method. `finitediff` is changed from `no` to `samegrid` automatically when `readqpss` and `writeqpss` are used to re-use QPSS results. Possible values are `no`, `yes` or `refine`.

Annotation parameters

- 33 `stats=no` Analysis statistics. Possible values are `no` or `yes`.
- 34 `annotate=sweep` Degree of annotation. Possible values are `no`, `title`, `sweep`, `status`, or `steps`.
- 35 `title` Analysis title.

Newton parameters

- 36 `maxiters=5` Maximum number of iterations per time step.
- 37 `restart=yes` Restart the DC solution from scratch if any condition has changed. If not, use the previous solution as initial guess. Possible values are `no` or `yes`.

Circuit age

- 38 `circuitage (Years)` Stress Time. Age of the circuit used to simulate hot-electron degradation of MOSFET and BSIM circuits.
- 39 `writeqpss` File to which final quasi-periodic steady-state solution is to be written. Small signal analyses such as `qpac`, `qpxf` and `qpnoise` can read in the steady-state solution from this file directly instead of running the `qpss` analysis again.

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Analysis Statements

40 `readqpss` File from which final quasi-periodic steady-state solution is to be read. Small signal analyses such as `qpac`, `qpxf` and `qpnoise` can read in the steady-state solution from this file directly instead of running the `qpss` analysis again.

Most of QPSS analysis parameters are inherited from PSS analysis, and their meanings remain essentially unchanged. Two new important parameters are `funds` and `maxharms`. They replace and extend the role of `fund` and `harms` parameters of PSS analysis. One important difference is that `funds` accepts a list of fundamental names instead of actual frequencies. The frequencies associated with fundamentals are figured out automatically by the simulator. An important feature is that each input signal can be a composition of more than one sources. However, these sources must have the same fundamental name. For each fundamental name, its fundamental frequency is the greatest common factor of all frequencies associated with the name. Missing or not listing all fundamental names using the parameter `funds` will result in an amputation of the current simulation. However if `maxharms` is not given, a warning message will be issued, and the number of harmonics is defaulted to 1 for each of the fundamentals.

For QPSS analyses, the role of some PSS parameters is extended. The parameter `maxperiods` that controls the maximum number of shooting iterations for PSS analysis also controls the number of the maximum number of shooting iterations for QPSS analysis. Its default value is set to 50.

The `tstab` parameter controls both the length of the initial transient integration with only the clock tone activated and the number of stable iterations with moderate tones activated. The stable iterations are run before Newton iterations.

The `errpreset` parameter lets you adjust several simulator parameters to fit your needs. In most cases, `errpreset` should be the only parameter you need to adjust. If you want a fast simulation with reasonable accuracy, you might set `errpreset` to `liberal`. If have some concern for accuracy, you might set `errpreset` to `moderate`. If accuracy is your main interest, you might set `errpreset` to `conservative`.

If users do not specify `steadyratio`, it is always 1.0, and it is not affected by `errpreset`. The following table shows the effect of `errpreset` on other parameters.

Parameter defaults as a function of `errpreset`

<code>errpreset</code>	<code>reltol</code>	<code>relref</code>	<code>method</code>	<code>lteratio</code>	<code>maxstep</code>
<code>liberal</code>	1e-3	<code>sigglobal</code>	<code>gear2only</code>	3.5	clock period/80
<code>moderate</code>	1e-4	<code>siggloaal</code>	<code>gear2only</code>	3.5	clock period/80

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conservative 1e-5 sigglobal gear2only * clock period/80

* : Iteratio=10.0 for conservative `errpreset` by default. Only if user specified `reltol` $\leq 1e-4 \cdot 10.0 / 3.5$, it will change to `Iteratio=3.5`.

The new `errpreset` settings include a new default `reltol` which is actually an enforced upper limit for appropriate setting. An increase of `reltol` above default will be ignored by the simulator. User can decrease this value in the options statement. The only way to increase `reltol` is to relax `errpreset`. Spectre sets the value of `maxstep` so that it is no larger than the value given in the table. Except for `reltol` and `maxstep`, `errpreset` does not change the value of any parameters you have explicitly set. The actual values used for the QPSS analysis are given in the log file.

If `errpreset` is not specified in the netlist, `liberal` settings will be used.

The default value for `compression` is `no`. The output file stores data for every signal at every time point for which Spectre calculates a solution. Spectre saves the x axis data only once, since every signal has the same x value. If `compression=yes`, Spectre writes data to the output file only if the signal value changes by at least 2*the convergence criteria. In order to save data for each signal independently, x axis information corresponding to each signal must be saved. If the signals stay at constant values for large periods of the simulation time, setting `compression=yes` results in a smaller output data file. If the signals in your circuit move around a lot, setting `compression=yes` results in a larger output data file.

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funds 1	oppoint 14	skipstart 15	tstart 4
ic 7	readic 9	skipstop 16	write 22
itres 31	readns 10	stats 33	writefinal 23

Quasi-Periodic Transfer Function Analysis (qpxf)

Description

A conventional transfer function analysis computes the transfer function from every source in the circuit to a single output. It differs from a conventional AC analysis in that the AC analysis computes the response from a single stimulus to every node in the circuit. The difference between QPAC and QPXF analysis are similar. The Quasi Periodic Transfer Function or QPXF analysis computes the transfer functions from any source at any frequency to a single output at a single frequency. Thus, like QPAC analysis, QPXF analysis includes frequency conversion effects.

The QPXF analysis directly computes such useful quantities as conversion efficiency (transfer function from input to output at desired frequency), image and sideband rejection (input to output at undesired frequency), and LO feed-through and power supply rejection (undesired input to output at all frequencies).

As with a QPAC, QPSP, and QPNOISE analyses, a QPXF analysis must follow a QPSS analysis.

Unlike other analyses in Spectre, this analysis can only sweep frequency.

Definition

Name [p] [n] qpxf parameter=value ...

The optional terminals (p and n) specify the output of the circuit. If you do not give the terminals, then you must specify the output with a probe component.

Parameters

Sweep interval parameters

- | | | |
|---|---------|--------------------|
| 1 | start=0 | Start sweep limit. |
| 2 | stop | Stop sweep limit. |
| 3 | center | Center of sweep. |

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4	<code>span=0</code>	Sweep limit span.
5	<code>step</code>	Step size, linear sweep.
6	<code>lin=50</code>	Number of steps, linear sweep.
7	<code>dec</code>	Points per decade.
8	<code>log=50</code>	Number of steps, log sweep.
9	<code>values=[...]</code>	Array of sweep values.
10	<code>sweepstype</code>	Specifies if the sweep frequency range is an absolute frequency, i.e. actual frequency; or if it is relative to the "relharmvec" sideband frequency. In QPSP, relative means relative to the input port frequency. Possible values are <code>absolute</code> or <code>relative</code> .
11	<code>relharmvec=[...]</code>	Sideband - vector of QPSS harmonics - to which relative frequency sweep should be referenced.

Probe parameters

12	<code>probe</code>	Compute every transfer function to this probe component.
----	--------------------	--

Output parameters

13	<code>stimuli=sources</code>	Stimuli used for xf analysis. Possible values are <code>sources</code> or <code>nodes_and_terminals</code> .
14	<code>sidevec=[...]</code>	Array of relevant sidebands for the analysis.
15	<code>clockmaxharm=0</code>	An alternative to the <code>sidevec</code> array specification, which automatically generates the array: <code>[-clockmaxharm ... 0 ... +clockmaxharms][maxharms(QPSS)[2]...0...maxharms(QPSS)[2]][...]</code> .
16	<code>freqaxis</code>	Specifies whether the results should be output versus the input frequency, the output frequency, or the absolute value of the input frequency. Default is <code>out</code> for logarithmic frequency sweeps

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and `absin` otherwise.
Possible values are `absin`, `in` or `out`.

17 `save` Signals to output.
Possible values are `all`, `lvl`, `allpub`, `lvlpub`, `selected`, or `none`.

18 `nestlvl` Levels of subcircuits to output.

Convergence parameters

19 `tolerance=1e-9` Relative tolerance for linear solver.

20 `gear_order=2` Gear order used for small-signal integration, 1 or 2.

21 `solver=turbo` Solver type.
Possible values are `std` or `turbo`.

Annotation parameters

22 `annotate=sweep` Degree of annotation.
Possible values are `no`, `title`, `sweep`, `status`, or `steps`.

23 `stats=no` Analysis statistics.
Possible values are `no` or `yes`.

24 `title` Analysis title.

The variable of interest at the output can be voltage or current, and its frequency is not constrained by the period of the large periodic solution. While sweeping the selected output frequency, you can select the periodic small-signal input frequencies of interest by setting either the `clockmaxharm` or the `sidevec` parameters. Sidebands are vectors in QPXF. Assume we have one large tone and one moderate tone in QPSS. A sideband `K1` will be represented as `[K1_1 K1_2]`. Corresponding frequency is

$$K1_1 * \text{fund}(\text{large tone of QPSS}) + K1_2 * \text{fund}(\text{moderate tone of QPSS})$$

We assume that there are `L` (1 large plus `L-1` moderate) tones in QPSS analysis and a given set of `n` integer vectors representing the sidebands

$$K1 = \{ K1_1, \dots, K1_j, \dots, K1_L \}, K2, \dots, Kn.$$

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The input signal frequency at each sideband is computed as

$$f(\text{in}) = f(\text{out}) + \text{SUM}_{j=1_to_L} \{K_i_j * \text{fund}_j(\text{qpss})\},$$

where $f(\text{out})$ represent the (possibly swept) output signal frequency, and $\text{fund}_j(\text{pss})$ represents the fundamental frequency used in the corresponding QPSS analysis. Thus, when analyzing a down-converting mixer, and sweeping the IF output frequency, $K_i = \{1, 0\}$ for the RF input represents the first upper-sideband, while $K_i = \{-1, 0\}$ for the RF input represents the first lower-sideband.

User would enter `sidevec` as a sequence of integer numbers, separated by spaces. The set of vectors $\{1\ 1\ 0\}$ $\{1\ -1\ 0\}$ $\{1\ 1\ 1\}$ becomes `sidevec=[1 1 0 1 -1 0 1 1 1]`. For `clockmaxharm`, only the large tone - first fundamental will be affected by this entry, all the rest - moderate tones - will be limited by `maxharms`, specified for a QPSS analysis. Given `maxharms=[k1 max k2max ... knmax]` in QPSS and `clockmaxharm=Kmax all (2*Kmax + 1)*(2*k2max+1)*(2*k3max+1)*...*(2*knmax+1)` sidebands are generated.

The number of requested sidebands changes substantially the simulation time.

With QPXF the frequency of the stimulus and of the response are usually different (this is an important way in which QPXF differs from XF). The `freqaxis` parameter is used to specify whether the results should be output versus the input frequency (`in`), the output frequency (`out`), or the absolute value of the input frequency (`absin`).

You can specify the output with a pair of nodes or a probe component. Any component with two or more terminals can be a voltage probe. When there are more than two terminals, they are grouped in pairs; and you use the `portv` parameter to select the appropriate pair of terminals. Alternatively, you can simply specify a voltage to be the output by giving a pair of nodes on the QPXF analysis statement.

Any component that naturally computes current as an internal variable can be a current probe. If the probe component computes more than one current, you use the `porti` parameter to select the appropriate current. It is an error to specify both `portv` and `porti`. If neither is specified, the probe component provides a reasonable default.

The `stimuli` parameter specifies what is used for the inputs for the transfer functions. There are two choices. `stimuli=sources` indicates that the sources present in the circuit should be used. The `xfmag` parameters provided by the sources may be used to adjust the computed gain to compensate for gains or losses in a test fixture. One can limit the number of sources in hierarchical netlists by using the `save` and `nestlvl` parameters. `stimuli=nodes_and_terminals` indicates that all possible transfer functions should be computed.

This is useful when it is not known in advance which transfer functions are interesting. Transfer functions for nodes are computed assuming that a unit magnitude flow (current)

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source is connected from the node to ground. Transfer functions for terminals are computed assuming that a unit magnitude value (voltage) source is connected in series with the terminal. By default, the transfer functions from a small set of terminals are computed. If transfer functions from specific terminals are desired, specify the terminals in the save statement. You must use the `:probe` modifier (ex. `Rout:1:probe`) or specify `useprobes=yes` on the options statement. If transfer functions from all terminals are desired, specify `currents=all` and `useprobes=yes` on the options statement.

You can specify sweep limits by giving the end points or by providing the center value and the span of the sweep. Steps can be linear or logarithmic, and you can specify the number of steps or the size of each step. You can give a step size parameter (`step`, `lin`, `log`, `dec`) to determine whether the sweep is linear or logarithmic. If you do not give a step size parameter, the sweep is linear when the ratio of stop to start values is less than 10, and logarithmic when this ratio is 10 or greater. Alternatively, you may specify the particular values that the sweep parameter should take using the `values` parameter. If you give both a specific set of values and a set specified using a sweep range, the two sets are merged and collated before being used. All frequencies are in Hertz.

Parameter Index

In the following index, the number following each parameter name indicates where to find the description of that parameter.

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Deferred Set Options (set)

Description

The deferred set options statement sets or changes various program control options. You can set the options in any order and, once set, the options retain their value until reset. The set statement is queued with all analyses and is executed sequentially (The changes made to

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Analysis Statements

these options are deferred until the statement setting them is encountered). To set `temp`, `tnom`, `scalem`, or `scale`, use the `alter` statement. For further options, see individual analyses.

Definition

Name set parameter=value ...

Parameters

Tolerance parameters

- | | | |
|---|------------------------------|---|
| 1 | <code>reltol=0.001</code> | Relative convergence criterion. |
| 2 | <code>vabstol=1e-06 V</code> | Voltage absolute tolerance convergence criterion. |
| 3 | <code>iabstol=1e-12 A</code> | Current absolute tolerance convergence criterion. |

Temperature parameters

- | | | |
|---|------------------------------|--|
| 4 | <code>tempeffects=all</code> | Temperature effect selector. If <code>tempeffect = vt</code> , only thermal voltage varies with temperature; if <code>tempeffect = tc</code> , parameters that start with <code>tc</code> are active and thermal voltage is dependent on temperature; and if <code>tempeffect = all</code> , all built-in temperature models are enabled.
Possible values are <code>vt</code> , <code>tc</code> or <code>all</code> . |
|---|------------------------------|--|

Convergence parameters

- | | | |
|---|---------------------------|---|
| 5 | <code>homotopy=all</code> | Method used when no convergence on initial attempt of DC analysis.
Possible values are <code>none</code> , <code>gmin</code> , <code>source</code> , <code>dptran</code> , <code>ptran</code> , <code>arclength</code> , or <code>all</code> . |
| 6 | <code>limit=dev</code> | Limiting algorithms to aid DC convergence.
Possible values are <code>delta</code> , <code>log</code> or <code>dev</code> . |

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Component parameters

- 7 `compatible=spectre` Encourage device equations to be compatible with a foreign simulator. This option does not affect input syntax. Possible values are `spectre`, `spice2`, `spice3`, `cdsspice`, `hspice`, or `spiceplus`.
- 8 `approx=no` Use approximate models. Difference between approximate and exact models is generally very small. Possible values are `no` or `yes`.

Error-checking parameters

- 9 `diagnose=no` Print additional information that might help diagnose accuracy and convergence problems. Possible values are `no` or `yes`.
- 10 `opptcheck=yes` Check operating point parameters against soft limits. Possible values are `no` or `yes`.

Resistance parameters

- 11 `gmin=1e-12 S` Minimum conductance across each nonlinear device.
- 12 `gmin_check=max_v_only` Specifies that effect of `gmin` should be reported if significant. Possible values are `no`, `max_v_only`, `max_only`, or `all`.
- 13 `rforce=1 Ω` Resistance used when forcing nodesets and node-based initial conditions.

Quantity parameters

- 14 `quantities=no` Print quantities. Possible values are `no`, `min` or `full`.

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Annotation parameters

- | | | |
|----|----------------------------------|--|
| 15 | <code>narrate=yes</code> | Narrate the simulation.
Possible values are <code>no</code> or <code>yes</code> . |
| 16 | <code>debug=no</code> | Give debugging messages.
Possible values are <code>no</code> or <code>yes</code> . |
| 17 | <code>info=yes</code> | Give informational messages.
Possible values are <code>no</code> or <code>yes</code> . |
| 18 | <code>note=yes</code> | Give notice messages.
Possible values are <code>no</code> or <code>yes</code> . |
| 19 | <code>maxnotes=5</code> | Maximum number of times any notice will be issued per analysis. |
| 20 | <code>warn=yes</code> | Give warning messages.
Possible values are <code>no</code> or <code>yes</code> . |
| 21 | <code>maxwarns=5</code> | Maximum number of times any warning message will be issued per analysis. |
| 22 | <code>maxwarnstologfile=5</code> | Maximum number of times any warning message will be printed to the log file per analysis. |
| 23 | <code>maxnotestologfile=5</code> | Maximum number of times any notice message will be printed to the log file per analysis. |
| 24 | <code>error=yes</code> | Give error messages.
Possible values are <code>no</code> or <code>yes</code> . |
| 25 | <code>digits=5</code> | Number of digits used when printing numbers. |
| 26 | <code>notation=eng</code> | When printing real numbers to the screen, what notation should be used.
Possible values are <code>eng</code> , <code>sci</code> or <code>float</code> . |
| 27 | <code>annotate=no</code> | Degree of annotation.
Possible values are <code>no</code> or <code>title</code> . |

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Matrix parameters

- 28 `pivotdc=no` Use numeric pivoting on every iteration of DC analysis.
Possible values are `no` or `yes`.
- 29 `pivrel=0.001` Relative pivot threshold.
- 30 `pivabs=0` Absolute pivot threshold.

Parameter Index

In the following index, the number following each parameter name indicates where to find the description of that parameter.

<code>annotate</code> 27	<code>gmin_check</code> 12	<code>maxwarnstologfile</code> 22	<code>quantities</code> 14
<code>approx</code> 8	<code>homotopy</code> 5	<code>narrate</code> 15	<code>reltol</code> 1
<code>compatible</code> 7	<code>iabstol</code> 3	<code>notation</code> 26	<code>rforce</code> 13
<code>debug</code> 16	<code>info</code> 17	<code>note</code> 18	<code>tempeffects</code> 4
<code>diagnose</code> 9	<code>limit</code> 6	<code>opptcheck</code> 10	<code>vabstol</code> 2
<code>digits</code> 25	<code>maxnotes</code> 19	<code>pivabs</code> 30	<code>warn</code> 20
<code>error</code> 24	<code>maxnotestologfile</code> 23	<code>pivotdc</code> 28	
<code>gmin</code> 11	<code>maxwarns</code> 21	<code>pivrel</code> 29	

Shell Command (shell)

Description

The shell analysis passes a command to the operating system command interpreter given in the `SHELL` environment variable. The command behaves as if it were typed into the command interpreter, except that any `%X` codes in the command are expanded first.

The default action of the shell analysis is to terminate the simulation.

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Analysis Statements

Definition

Name shell parameter=value ...

Parameters

- | | | |
|---|---------------|---|
| 1 | cmd="kill %P" | Shell command. |
| 2 | iferror=quit | What to do if command returns nonzero error status.
Possible values are <code>quit</code> or <code>continue</code> . |
| 3 | annotate | Degree of annotation.
Possible values are <code>no</code> , <code>title</code> or <code>yes</code> . |

S-Parameter Analysis (sp)

Description

The S-parameter analysis linearizes the circuit about the DC operating point and computes S-parameters of the circuit taken as an N-port. The port statements define the ports of the circuit. Each active port is turned on sequentially, and a linear small-signal analysis is performed. Spectre converts the response of the circuit at each active port into S-parameters and outputs these parameters. There must be at least one active port statement in the circuit.

If a filename is specified using the `file` parameter, the S-parameter analysis generates an ASCII file containing the S-parameters of the circuit that can later be read-in by the `nport` component. The generated file can be in either Spectres native format or in Touchstone format.

Spectre can perform the analysis while sweeping a parameter. The parameter can be frequency, temperature, component instance parameter, component model parameter, or netlist parameter. If changing a parameter affects the DC operating point, the operating point is recomputed on each step. You can sweep the circuit temperature by giving the parameter name as `temp` with `no dev` or `mod` parameter. You can sweep a netlist parameter by giving the parameter name with `no dev`, or `mod` parameter. After the analysis has completed, the modified parameter returns to its original value.

Definition

Name sp parameter=value ...

Spectre Circuit Simulator Reference

Analysis Statements

Parameters

1 `prevoppoint=no` Use operating point computed on the previous analysis.
Possible values are `no` or `yes`.

Sweep interval parameters

2 `start=0` Start sweep limit.

3 `stop` Stop sweep limit.

4 `center` Center of sweep.

5 `span=0` Sweep limit span.

6 `step` Step size, linear sweep.

7 `lin=50` Number of steps, linear sweep.

8 `dec` Points per decade.

9 `log=50` Number of steps, log sweep.

10 `values=[...]` Array of sweep values.

Sweep variable parameters

11 `dev` Device instance whose parameter value is to be swept.

12 `mod` Model whose parameter value is to be swept.

13 `param` Name of parameter to sweep.

14 `freq (Hz)` Frequency when parameter other than frequency is being swept.

Port parameters

15 `ports=[...]` List of active ports. Ports are numbered in the order given.

Spectre Circuit Simulator Reference

Analysis Statements

State-file parameters

16 `readns` File that contains estimate of DC solution (nodeset).

Output parameters

17 `file` S-parameters output file name.

18 `datafmt=spectre` Data format of the S-parameter output file.
Possible values are `spectre` or `touchstone`.

19 `datatype=realimag` Data type of the S-parameter output file.
Possible values are `realimag`, `magphase` or `dbphase`.

20 `noisedata=no` Should noise data be saved to the S-parameter output file, and if so, in what format.
Possible values are `no`, `twoport` or `cy`.

21 `oppoint=no` Should operating point information be computed, and if so, where should it be sent. Operating point information would not be output if operating point is computed in the previous analysis and is unchanged.
Possible values are `no`, `screen`, `logfile`, or `rawfile`.

Noise parameters

22 `donoise=no` Perform noise analysis. If `oprobe` is specified as a valid port, this is set to `yes`, and a detailed noise output is generated.
Possible values are `no` or `yes`.

23 `oprobe` Compute total noise at the output defined by this component.

24 `iprobe` Input probe. Refer the output noise to this component.

Convergence parameters

25 `restart=yes` Restart the DC solution from scratch if any condition has changed. If not, use the previous solution as initial guess.
Possible values are `no` or `yes`.

Spectre Circuit Simulator Reference

Analysis Statements

Annotation parameters

- 26 `annotate=sweep` Degree of annotation.
Possible values are `no`, `title`, `sweep`, `status`, or `steps`.
- 27 `stats=no` Analysis statistics.
Possible values are `no` or `yes`.
- 28 `title` Analysis title.

If the list of active ports is specified with the `ports` parameter, then the ports are numbered sequentially from one in the order given. Otherwise, all ports present in the circuit are active, and the port numbers used are those that were assigned on the port statements. If `donoise=yes` is specified, then the noise correlation matrix is computed. If in addition, the output is specified using `oprobe`, the amount that each noise source contributes to the output is computed. Finally, if an input is also specified (using `iprobe`), the two-port noise parameters are computed (F, Fmin, NF, NFmin, Gopt, Bopt, and Rn).

You can specify sweep limits by giving the end points or by providing the center value and the span of the sweep. Steps can be linear or logarithmic, and you can specify the number of steps or the size of each step. You can give a step size parameter (`step`, `lin`, `log`, `dec`) to determine whether the sweep is linear or logarithmic. If you do not give a step size parameter, the sweep is linear when the ratio of stop to start values is less than 10, and logarithmic when this ratio is 10 or greater. All frequencies are in Hertz.

The small-signal analysis begins by linearizing the circuit about an operating-point. By default this analysis computes the operating-point if it is not known, or recomputes it if any significant component or circuit parameter has changed. However, if a previous analysis computed an operating point, you can set `prevoppoint=yes` to avoid recomputing it. For example, if you use this option when the previous analysis was a transient analysis, the operating point is the state of the circuit on the final time point.

Parameter Index

In the following index, the number following each parameter name indicates where to find the description of that parameter.

<code>annotate</code> 26	<code>file</code> 17	<code>oppoint</code> 21	<code>span</code> 5
<code>center</code> 4	<code>freq</code> 14	<code>oprobe</code> 23	<code>start</code> 2
<code>datafmt</code> 18	<code>iprobe</code> 24	<code>param</code> 13	<code>stats</code> 27

Spectre Circuit Simulator Reference Analysis Statements

datatype 19	lin 7	ports 15	step 6
dec 8	log 9	prevoppoint 1	stop 3
dev 11	mod 12	readns 16	title 28
donoise 22	noisedata 20	restart 25	values 10

Stability Analysis (stb)

Description

The STB analysis linearizes the circuit about the DC operating point and computes the loop gain, gain and phase margins (if the sweep variable is frequency), for a feedback loop or a gain device.

Spectre can perform the analysis while sweeping a parameter. The parameter can be frequency, temperature, component instance parameter, component model parameter, or netlist parameter. If changing a parameter affects the DC operating point, the operating point is recomputed on each step. You can sweep the circuit temperature by giving the parameter name as `temp` with no `dev` or `mod` parameter. You can sweep a netlist parameter by giving the parameter name with no `dev`, or `mod` parameter. After the analysis has completed, the modified parameter returns to its original value.

Definition

Name `stb parameter=value ...`

Parameters

1 `prevoppoint=no` Use operating point computed on the previous analysis.
Possible values are `no` or `yes`.

Sweep interval parameters

2 `start=0` Start sweep limit.

3 `stop` Stop sweep limit.

4 `center` Center of sweep.

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Analysis Statements

5	<code>span=0</code>	Sweep limit span.
6	<code>step</code>	Step size, linear sweep.
7	<code>lin=50</code>	Number of steps, linear sweep.
8	<code>dec</code>	Points per decade.
9	<code>log=50</code>	Number of steps, log sweep.
10	<code>values=[...]</code>	Array of sweep values.

Sweep variable parameters

11	<code>dev</code>	Device instance whose parameter value is to be swept.
12	<code>mod</code>	Model whose parameter value is to be swept.
13	<code>param</code>	Name of parameter to sweep.
14	<code>freq (Hz)</code>	Frequency when parameter other than frequency is being swept.

Probe parameters

15	<code>probe</code>	Probe instance around which the loop gain is calculated.
----	--------------------	--

State-file parameters

16	<code>readns</code>	File that contains estimate of DC solution (nodeset).
----	---------------------	---

Output parameters

17	<code>save</code>	Signals to output. Possible values are <code>all</code> , <code>lvl</code> , <code>allpub</code> , <code>lvlpub</code> , <code>selected</code> , or <code>none</code> .
18	<code>nestlvl</code>	Levels of subcircuits to output.

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Analysis Statements

- 19 `oppooint=no` Should operating point information be computed, and if so, where should it be sent. Operating point information would not be output if operating point is computed in the previous analysis and is unchanged.
Possible values are `no`, `screen`, `logfile`, or `rawfile`.

Convergence parameters

- 20 `restart=yes` Restart the DC solution from scratch if any condition has changed. If not, use the previous solution as initial guess.
Possible values are `no` or `yes`.

Annotation parameters

- 21 `annotate=sweep` Degree of annotation.
Possible values are `no`, `title`, `sweep`, `status`, or `steps`.
- 22 `stats=no` Analysis statistics.
Possible values are `no` or `yes`.
- 23 `title` Analysis title.

You can specify sweep limits by giving the end points or by providing the center value and the span of the sweep. Steps can be linear or logarithmic, and you can specify the number of steps or the size of each step. You can give a step size parameter (`step`, `lin`, `log`, `dec`) to determine whether the sweep is linear or logarithmic. If you do not give a step size parameter, the sweep is linear when the ratio of stop to start values is less than 10, and logarithmic when this ratio is 10 or greater. All frequencies are in Hertz.

The small-signal analysis begins by linearizing the circuit about an operating-point. By default this analysis computes the operating-point if it is not known, or recomputes it if any significant component or circuit parameter has changed. However, if a previous analysis computed an operating point, you can set `prevoppooint=yes` to avoid recomputing it. For example, if you use this option when the previous analysis was a transient analysis, the operating point is the state of the circuit on the final time point.

Understanding Loop based and Device Based Algorithms

Two algorithms--the loop based and the device based, are available for small-signal stability analysis. Both algorithms are based on the calculation of Bodes return ratio. Loop gain waveform, gain margin, and phase margin are the analysis output.

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Analysis Statements

The `probe` parameter must be specified to perform stability analysis. When it points to a current probe or voltage source instance, the loop based algorithm will be invoked; when it points to a supported active device instance, the device based algorithm will be invoked.

Loop Based Algorithm

The loop based algorithm calculates the true loop gain that consists of normal loop gain and reverse loop gain. The loop based algorithm requires the `probe` being placed on the feedback loop to identify and characterize the particular loop of interest. The introduction of the probe component should not change any of the circuit characteristics.

The loop based algorithm provides accurate stability information for single loop circuits, and multiloop circuits in which a `probe` component can be placed on a critical wire to break all loops. For a general multiloop circuit, such a critical wire may not be available. The loop based algorithm can only be performed on individual feedback loops to ensure they are stable. Although the stability of all feedback loops is only a necessary condition for the whole circuit to be stable, the multiloop circuit tends to be stable if all individual loops are associated with reasonable stability margins.

Device Based Algorithm

The device based algorithm calculates the loop gain around a particular active device. This algorithm is often applied to assess the stability of circuit design in which local feedback loops cannot be neglected; the loop based algorithm cannot be performed for these applications since the local feedback loops are inside the devices, they are not accessible from the schematic level or netlist level to insert the `probe` component.

With the `probe` parameter points to a particular active device, the dominant controlled source in the device will be nulled during the analysis. The dominant controlled source is defined as by nulling this source renders the active device to be passive. The device based algorithm produces accurate stability information for a circuit in which a critical active device can be identified such that nulling the dominant gain source of this device renders the whole network to be passive.

Stability Analysis of Differential Feedback Circuits

A balanced fully differential feedback circuit is illustrated below:

|-----[ZF]-----|

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```

      |           |
      |           X1
      | |-----| |
      | |           | |
... ---[ ZS ]---|--|in+   out- |--|---[ ZL ]--- ...
      |   OPAMP   |
... ---[ ZS ]---|--|in-   out+ |--|---[ ZL ]--- ...
      | |           | |
      | |-----| |
      |           X2
      |           |
      |-----[ ZF ]-----|

```

The feedback loops are broken at X1 and X2, with x1in and x2in being the input side nodes, x1out and x2out being the output side nodes. The following subcircuit connects these four nodes together:

```

subckt diffprobe x1in x2in x1out x2out

  ibranh inout x1out iprobe

  vinj inout x1in iprobe

  evinj x2in x2out x1in x1out vcvs gain=0

  fiinj 0 x2out pcccs probes=[ibranch vinj] coeffs=[0 1 1] gain=0

ends diffprobe

```

Let `diffprobe_inst` be the instance of subcircuit `diffprobe`, the following analysis measures the differential-mode loop gain:

```

DMalterv alter dev=diffprobe_inst.evinj param=gain value=-1

```

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Analysis Statements

```
DMalteri alter dev=diffprobe_inst.fiinj param=gain value=-1
```

```
DMloopgain stb probe=diffprobe_inst.vinj
```

and the following analysis measures the common-mode loop gain:

```
CMalterv alter dev=diffprobe_inst.evinj param=gain value=1
```

```
CMalteri alter dev=diffprobe_inst.fiinj param=gain value=1
```

```
CMloopgain stb probe=diffprobe_inst.vinj
```

Parameter Index

In the following index, the number following each parameter name indicates where to find the description of that parameter.

annotate	21	log	9	probe	15	stats	22
center	4	mod	12	readns	16	step	6
dec	8	nestlvl	18	restart	20	stop	3
dev	11	oppoint	19	save	17	title	23
freq	14	param	13	span	5	values	10
lin	7	prevoppoint	1	start	2		

Sweep Analysis (sweep)

Description

The `sweep` analysis sweeps a parameter executing the list of analyses (or multiple analyses) for each value of the parameter. The swept parameter can be circuit temperature, a device instance parameter, a device model parameter, a netlist parameter, or a subcircuit parameter for a particular subcircuit instance.

A set of parameters can be swept simultaneously, using the `paramset` parameter. The other sweep interval or variable parameters cannot be specified with the `paramset` parameter. Do `spectre -h paramset` for information on defining a `paramset`.

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Analysis Statements

Within a sweep statement, you can specify analyses statements. These statements should be bound within braces. The opening brace is required at the end of the line defining the sweep. Sweep statements can be nested.

You can sweep the circuit temperature by giving the parameter name as `param=temp` with no `dev`, `mod`, or `sub` parameter. You can sweep a top-level netlist parameter by giving the parameter name with no `dev`, `mod`, or `sub` parameter. You can sweep a subcircuit parameter for a particular subcircuit instance by specifying the subcircuit instance name with the `sub` parameter and the subcircuit parameter name with the `param` parameter. The same can be done using `dev` for the device instance name or `mod` for the device model name.

After the analysis has completed, the modified parameter returns to its original value.

Definition

Name sweep parameter=value ...

Parameters

Sweep interval parameters

1	<code>start=0</code>	Start sweep limit.
2	<code>stop</code>	Stop sweep limit.
3	<code>center</code>	Center of sweep.
4	<code>span=0</code>	Sweep limit span.
5	<code>step</code>	Step size, linear sweep.
6	<code>lin=50</code>	Number of steps, linear sweep.
7	<code>dec</code>	Points per decade.
8	<code>log=50</code>	Number of steps, log sweep.
9	<code>values=[...]</code>	Array of sweep values.

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Analysis Statements

Sweep variable parameters

10	<code>dev</code>	Device instance whose parameter value is to be swept.
11	<code>sub</code>	Subcircuit instance whose parameter value is to be swept.
12	<code>mod</code>	Model whose parameter value is to be swept.
13	<code>param</code>	Name of parameter to sweep.
14	<code>paramset</code>	Name of parameter set to sweep.

Annotation parameters

15	<code>annotate=sweep</code>	Degree of annotation. Possible values are <code>no</code> , <code>title</code> or <code>sweep</code> .
16	<code>title</code>	Analysis title.

You can specify sweep limits by giving the end points or by providing the center value and the span of the sweep. Steps can be linear or logarithmic, and you can specify the number of steps or the size of each step. You can give a step size parameter (`step`, `lin`, `log`, or `dec`) and determine whether the sweep is linear or logarithmic. If you do not give a step size parameter, the sweep is linear when the ratio of the stop-to-start values is less than 10 and logarithmic when this ratio is 10 or greater.

Example:

```
swp sweep param=temp values=[-50 0 50 100 125] {  
    oppoint dc oppoint=logfile  
}
```

Parameter Index

In the following index, the number following each parameter name indicates where to find the description of that parameter.

```
annotate 15          lin 6          paramset 14        stop 2
```

Spectre Circuit Simulator Reference Analysis Statements

center 3	log 8	span 4	sub 11
dec 7	mod 12	start 1	title 16
dev 10	param 13	step 5	values 9

Time-Domain Reflectometer Analysis (tdr)

Description

The time-domain reflectometer analysis linearizes the circuit about the DC operating point and computes the reflection coefficients versus time, looking from the active ports into the circuit.

Definition

Name `tdr parameter=value ...`

Parameters

- | | | |
|----|------------------------------|--|
| 1 | <code>stop</code> | Stop time. |
| 2 | <code>settling=stop</code> | Time required for circuit to settle. |
| 3 | <code>start=-0.1 stop</code> | Time output waveforms begin. |
| 4 | <code>smoothing=2</code> | Window smoothing parameter (useful range is 0 to 15). |
| 5 | <code>vel=1</code> | Propagation velocity of medium normalized to <i>c</i> . |
| 6 | <code>points=64</code> | Number of time points. |
| 7 | <code>ports=[...]</code> | List of active ports. If not given, all ports are used. |
| 8 | <code>readns</code> | File that contains estimate of DC solution (nodeset). |
| 9 | <code>restart=yes</code> | Restart the DC solution from scratch if any condition has changed. If not, use the previous solution as initial guess. Possible values are <code>no</code> or <code>yes</code> . |
| 10 | <code>annotate=sweep</code> | Degree of annotation. Possible values are <code>no</code> , <code>title</code> , <code>sweep</code> , <code>status</code> , or <code>steps</code> . |

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Analysis Statements

- 11 `title` Analysis title.
- 12 `oppooint=no` Should operating point information be computed, and if so, where should it be sent. Operating point information would not be output if operating point is computed in the previous analysis and is unchanged.
Possible values are `no`, `screen`, `logfile`, or `rawfile`.
- 13 `prevoppooint=yes` Use operating point computed on the previous analysis.
Possible values are `no` or `yes`.

Such a small-signal analysis begins by linearizing the circuit about an operating point. By default, this analysis computes the operating point, if it is not yet known, or recomputes it, if any significant component or circuit parameter has changed. However, if a previous analysis computed an operating point, you can set `prevoppooint=yes` to avoid recomputing it. For example, if you use this command when the previous analysis was a transient analysis, the operating point is the state of the circuit on the final time point.

Parameter Index

In the following index, the number following each parameter name indicates where to find the description of that parameter.

<code>annotate</code>	10	<code>prevoppooint</code>	13	<code>smoothing</code>	4	<code>vel</code>	5
<code>oppooint</code>	12	<code>readns</code>	8	<code>start</code>	3		
<code>points</code>	6	<code>restart</code>	9	<code>stop</code>	1		
<code>ports</code>	7	<code>settling</code>	2	<code>title</code>	11		

Transient Analysis (`tran`)

Description

This analysis computes the transient response of a circuit over the interval from `start` to `stop`. The initial condition is taken to be the DC steady-state solution if not otherwise given.

Definition

Name `tran parameter=value ...`

Spectre Circuit Simulator Reference

Analysis Statements

Parameters

Simulation interval parameters

- | | | |
|---|----------------------------------|---|
| 1 | <code>stop (s)</code> | Stop time. |
| 2 | <code>start=0 s</code> | Start time. |
| 3 | <code>outputstart=start s</code> | Output is saved only after this time is reached. |
| 4 | <code>autostop=no</code> | If yes, the analysis is terminated when all event-type measurement expressions have been evaluated. Event-type expressions use thresholding, event or delay type functions. Possible values are <code>no</code> or <code>yes</code> . |

Time-step parameters

- | | | |
|---|--|--|
| 5 | <code>maxstep (s)</code> | Maximum time step. Default derived from <code>errpreset</code> . |
| 6 | <code>step=0.001 (stop-start) s</code> | Minimum time step used by the simulator solely to maintain the aesthetics of the computed waveforms. |

Initial-condition parameters

- | | | |
|---|------------------------|--|
| 7 | <code>ic=all</code> | What should be used to set initial condition. Possible values are <code>dc</code> , <code>node</code> , <code>dev</code> , or <code>all</code> . |
| 8 | <code>skipdc=no</code> | If yes, there will be no dc analysis for transient. Possible values are <code>no</code> , <code>yes</code> , <code>waveless</code> , <code>rampup</code> , <code>autodc</code> , or <code>sigrampup</code> . |
| 9 | <code>readic</code> | File that contains initial condition. |

Convergence parameters

- | | | |
|----|---------------------|--|
| 10 | <code>readns</code> | File that contains estimate of initial transient solution. |
|----|---------------------|--|

Spectre Circuit Simulator Reference

Analysis Statements

11 `cmin=0 F` Minimum capacitance from each node to ground.

State-file parameters

12 `write` File to which initial transient solution is to be written.

13 `writefinal` File to which final transient solution is to be written.

14 `ckptperiod` Checkpoint the analysis periodically using the specified period.

Integration method parameters

15 `method` Integration method. Default derived from `errpreset`. Possible values are `euler`, `trap`, `traponly`, `gear2`, `gear2only`, or `trapgear2`.

Accuracy parameters

16 `errpreset` Selects a reasonable collection of parameter settings. Possible values are `liberal`, `moderate` or `conservative`.

17 `relref` Reference used for the relative convergence criteria. Default derived from `errpreset`. Possible values are `pointlocal`, `alllocal`, `sigglobal`, or `allglobal`.

18 `lteratio` Ratio used to compute LTE tolerances from Newton tolerance. Default derived from `errpreset`.

Annotation parameters

19 `stats=no` Analysis statistics. Possible values are `no` or `yes`.

20 `annotate=sweep` Degree of annotation. Possible values are `no`, `title`, `sweep`, `status`, or `steps`.

21 `title` Analysis title.

Spectre Circuit Simulator Reference Analysis Statements

Output parameters

- 22 `save` Signals to output.
Possible values are `all`, `lvl`, `allpub`, `lvlpub`, `selected`, or `none`.
- 23 `nestlvl` Levels of subcircuits to output.
- 24 `oppooint=no` Should operating point information be computed for initial timestep, and if so, where should it be sent.
Possible values are `no`, `screen`, `logfile`, or `rawfile`.
- 25 `skipstart=starttime s` The time to start skipping output data.
- 26 `skipstop=stoptime s` The time to stop skipping output data.
- 27 `skipcount` Save only one of every `skipcount` points.
- 28 `strobeperiod (s)` The output strobe interval (in seconds of transient time).
- 29 `strobedelay=0 s` The delay (phase shift) between the `skipstart` time and the first strobe point.
- 30 `compression=no` Do data compression on output. See full description below.
Possible values are `no` or `yes`.
- 31 `flushpoints` Flush outputs after number of calculated points.
- 32 `flushtime (s)` Flush outputs after real time has elapsed.
- 33 `flushofftime (s)` Time to stop flushing outputs.
- 34 `infoname` Name of info analysis to be performed at each time point in the `infotimes` array.
- 35 `infotimes=[...] s` Times when info analysis specified by `infoname` is performed.

Newton parameters

- 36 `maxiters=5` Maximum number of iterations per time step.

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Analysis Statements

37 `restart=yes` Restart the DC solution from scratch if any condition has changed. If not, use the previous solution as initial guess. Possible values are `no` or `yes`.

Circuit age

38 `circuitage (Years)` Stress Time. Age of the circuit used to simulate hot-electron degradation of MOSFET and BSIM circuits.

You may specify the initial condition for the transient analysis by using the `ic` statement or the `ic` parameter on the capacitors and inductors. If you do not specify the initial condition, the DC solution is used as the initial condition. The `ic` parameter on the transient analysis controls the interaction of various methods of setting the initial conditions. The effects of individual settings are:

`ic=dc`: Any initial condition specifiers are ignored, and the DC solution is used.

`ic=node`: The `ic` statements are used, and the `ic` parameter on the capacitors and inductors are ignored.

`ic=dev`: The `ic` parameters on the capacitors and inductors are used, and the `ic` statements are ignored.

`ic=all`: Both the `ic` statements and the `ic` parameters are used, and the `ic` parameters override the `ic` statements.

If you specify an initial condition file with the `readic` parameter, initial conditions from the file are used, and any `ic` statements are ignored.

Once you specify the initial conditions, Spectre computes the actual initial state of the circuit by performing a DC analysis. During this analysis, Spectre forces the initial conditions on nodes by using a voltage source in series with a resistor whose resistance is `rforce` (see `options`).

With the `ic` statement it is possible to specify an inconsistent initial condition (one that cannot be sustained by the reactive elements). Examples of inconsistent initial conditions include setting the voltage on a node with no path of capacitors to ground or setting the current through a branch that is not an inductor. If you initialize Spectre inconsistently, its solution jumps; that is, it changes instantly at the beginning of the simulation interval. You should avoid such changes if possible because Spectre can have convergence problems while trying to make the jump.

You can skip the DC analysis entirely by using the parameter `skipdc`. If the DC analysis is skipped, the initial solution will be either trivial, or given in the file you specified by the `readic`

Spectre Circuit Simulator Reference

Analysis Statements

parameter, or, if the `readic` parameter is not given, the values specified on the `ic` statements. Device based initial conditions are not used for `skipdc`. Nodes that you do not specify with the `ic` file or `ic` statements will start at zero. You should not use this parameter unless you are generating a nodeset file for circuits that have trouble in the DC solution; it usually takes longer to follow the initial transient spikes that occur when the DC analysis is skipped than it takes to find the real DC solution. The `skipdc` parameter might also cause convergence problems in the transient analysis.

The possible settings of parameter `skipdc` and their meanings are:

`skipdc=no`: Initial solution is calculated using the normal DC analysis (default).

`skipdc=yes`: Initial solution is given in the file specified by the `readic` parameter or the values specified on the `ic` statements.

`skipdc=waveless`: Same initial solution as `skipdc=yes`, but the waveform production in the time-varying independent sources is disabled during the transient analysis. Independent source values are fixed to their initial values (not their DC values).

`skipdc=rampup`: Independent source values start at 0 and ramp up to their initial values in the first 10% of the analysis interval. After that their values remain constant. Zero initial solution is used.

`skipdc=autodc`: Same as `skipdc=waveless` if a nonzero initial condition is specified. Otherwise, same as `skipdc=rampup`.

`skipdc=sigrampup`: Independent source values start at 0 and ramp up to their initial values in the first phase of the simulation. Unlike `skipdc=rampup`, the waveform production in the time-varying independent source is enabled after the rampup phase. The rampup simulation is from `start` to `stop`. If the `start` parameter is not specified, the default `start` time is set to $0.1 * stop$.

Nodesets help find the DC or initial transient solution. You can supply them in the circuit description file with `nodeset` statements, or in a separate file using the `readns` parameter. When nodesets are given, Spectre computes an initial guess of the solution by performing a DC analysis while forcing the specified values onto nodes by using a voltage source in series with a resistor whose resistance is `rforce`. Spectre then removes these voltage sources and resistors and computes the true solution from this initial guess.

Nodesets have two important uses. First, if a circuit has two or more solutions, nodesets can bias the simulator towards computing the desired one. Second, they are a convergence aid. By estimating the solution of the largest possible number of nodes, you might be able to eliminate a convergence problem or dramatically speed convergence.

Spectre Circuit Simulator Reference

Analysis Statements

When you simulate the same circuit many times, we suggest that you use both the `write` and `readns` parameters and give the same file name to both parameters. The DC analysis then converges quickly even if the circuit has changed somewhat since the last simulation, and the nodeset file is automatically updated.

Nodesets and initial conditions have similar implementation but produce different effects. Initial conditions actually define the solution, whereas nodesets only influence it. When you simulate a circuit with a transient analysis, Spectre forms and solves a set of differential equations. However, differential equations have an infinite number of solutions, and a complete set of initial conditions must be specified in order to identify the desired solution. Any initial conditions you do not specify are computed by the simulator to be consistent. The transient waveforms then start from initial conditions. Nodesets are usually used as a convergence aid and do not affect the final results. However, in a circuit with more than one solution, such as a latch, nodesets bias the simulator towards finding the solution closest to the nodeset values.

The `method` parameter specifies the integration method. The possible settings and their meanings are:

`method=euler:` Backward-Euler is used exclusively.

`method=traponly:` Trapezoidal rule is used almost exclusively.

`method=trap:` Backward-Euler and the trapezoidal rule are used.

`method=gear2only:` Gears second-order backward-difference method is used almost exclusively.

`method=gear2:` Backward-Euler and second-order Gear are used.

`method=trapgear2:` Allows all three integration methods to be used.

The trapezoidal rule is usually the most efficient when you want high accuracy. This method can exhibit point-to-point ringing, but you can control this by tightening the error tolerances. For this reason, though, if you choose very loose tolerances to get a quick answer, either backward-Euler or second-order Gear will probably give better results than the trapezoidal rule. Second-order Gear and backward-Euler can make systems appear more stable than they really are. This effect is less pronounced with second-order Gear or when you request high accuracy.

Several parameters determine the accuracy of the transient analysis. `reltol` and `abstol` control the accuracy of the discretized equation solution. These parameters determine how well charge is conserved and how accurately steady-state or equilibrium points are computed. You can set the integration error, or the errors in the computation of the circuit

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Analysis Statements

dynamics (such as time constants), relative to `reltol` and `abstol` by setting the `lteratio` parameter.

The parameter `relref` determines how the relative error is treated. The `relref` options are:

`relref=pointlocal`: Compares the relative errors in quantities at each node to that node alone.

`relref=alllocal`: Compares the relative errors at each node to the largest values found for that node alone for all past time.

`relref=sigglobal`: Compares relative errors in each of the circuit signals to the maximum for all signals at any previous point in time.

`relref=allglobal`: Same as `relref=sigglobal` except that it also compares the residues (KCL error) for each node to the maximum of that nodes past history.

The `errpreset` parameter lets you adjust the simulator parameters to fit your needs quickly. You can set `errpreset` to `conservative` if the circuit is very sensitive, or you can set it to `liberal` for a fast, but possibly inaccurate, simulation. The setting `errpreset=moderate` suits most needs.

The effect of `errpreset` on other parameters is shown in the following table. In this table, $T = \text{stop} - \text{start}$.

<code>errpreset</code>	<code>reltol</code>	<code>relref</code>	<code>method</code>	<code>maxstep</code>	<code>lteratio</code>
liberal	* 10	allglobal	gear2	Interval/10	3.5
moderate		sigglobal	traonly	Interval/50	3.5
conservative	* 0.1	alllocal	gear2only	Interval/100	10.0

The default value for `errpreset` is `moderate`.

The value of `reltol` is increased or decreased from its value in the options statement, but it is not allowed to be larger than 0.01. Spectre sets the value of `maxstep` so that it is no larger than the value given in the table. Except for `reltol` and `maxstep`, `errpreset` does not change the value of any parameters you have explicitly set. The actual values used for the transient analysis are given in the log file.

If the circuit you are simulating can have infinitely fast transitions (for example, a circuit that contains nodes with no capacitance), Spectre might have convergence problems. To avoid this, you must prevent the circuit from responding instantaneously. You can accomplish this

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by setting `cmin`, the minimum capacitance to ground at each node, to a physically reasonable nonzero value. This often significantly improves Spectre convergence.

Spectre provides two methods for reducing the number of output data points saved: `strobing`, based on the simulation time, and `skipping` time points, which saves only every Nth point.

The parameters `strobeperiod` and `strobedelay` control the strobing method. `strobeperiod` sets the interval between points that you want to save, and `strobedelay` sets the offset within the period relative to `skipstart`. The simulator forces a time step on each point to be saved, so the data is computed, not interpolated.

The skipping method is controlled by `skipcount`. If this is set to N, then only every Nth point is saved.

The parameters `skipstart` and `skipstop` apply to both data reduction methods. Before `skipstart` and after `skipstop`, Spectre saves all computed data.

If you do not want any data saved before a given time, use `outputstart`. If you do not want any data saved after a given time, change the `stop` time.

The default value for `compression` is `no`. The output file stores data for every signal at every time point for which Spectre calculates a solution. Spectre saves the x axis data only once, since every signal has the same x value. If `compression=yes`, Spectre writes data to the output file only if the signal value changes by at least 2*the convergence criteria. In order to save data for each signal independently, x axis information corresponding to each signal must be saved. If the signals stay at constant values for large periods of the simulation time, setting `compression=yes` results in a smaller output data file. If the signals in your circuit move around a lot, setting `compression=yes` results in a larger output data file.

Parameter Index

In the following index, the number following each parameter name indicates where to find the description of that parameter.

<code>annotate</code>	20	<code>ic</code>	7	<code>readic</code>	9	<code>stats</code>	19
<code>autostop</code>	4	<code>infoname</code>	34	<code>readns</code>	10	<code>step</code>	6
<code>circuitage</code>	38	<code>infotimes</code>	35	<code>relref</code>	17	<code>stop</code>	1
<code>ckptperiod</code>	14	<code>lteratio</code>	18	<code>restart</code>	37	<code>strobedelay</code>	29
<code>cmin</code>	11	<code>maxiters</code>	36	<code>save</code>	22	<code>strobeperiod</code>	28

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compression	30	maxstep	5	skipcount	27	title	21
errpreset	16	method	15	skipdc	8	write	12
flushofftime	33	nestlvl	23	skipstart	25	writeln	13
flushpoints	31	oppoint	24	skipstop	26		
flushtime	32	outputstart	3	start	2		

Transfer Function Analysis (xf)

Description

The transfer function analysis linearizes the circuit about the DC operating point and performs a small-signal analysis that calculates the transfer function from every independent source in the circuit to a designated output. The variable of interest at the output can be voltage or current.

You can specify the output with a pair of nodes or a probe component. Any component with two or more terminals can be a voltage probe. When there are more than two terminals, they are grouped in pairs; and you use the `portv` parameter to select the appropriate pair of terminals. Alternatively, you can simply specify a voltage to be the output by giving a pair of nodes on the `xf` analysis statement.

Any component that naturally computes current as an internal variable can be a current probe. If the probe component computes more than one current (as transmission lines, microstrip lines, and N-ports do), you use the `porti` parameter to select the appropriate current. It is an error to specify both `portv` and `porti`. If neither is specified, the probe component provides a reasonable default.

The `stimuli` parameter specifies what is used for the inputs for the transfer functions. There are two choices. `stimuli=sources` indicates that the sources present in the circuit should be used. The `xfmag` parameters provided by the sources may be used to adjust the computed gain to compensate for gains or losses in a test fixture. One can limit the number of sources in hierarchical netlists by using the `save` and `nestlvl` parameters.

The transfer functions computed versus output and source types are:

Source		Output Type		Source		
Type		voltage		current		Amplitude
-----+-----+-----+-----						

Spectre Circuit Simulator Reference

Analysis Statements

`vsource | V(out)/V(src) | I(out)/V(src) | V(src)=xfmag`

`isource | V(out)/I(src) | I(out)/I(src) | I(src)=xfmag`

`port | 2*V(out)/V(src) | 2*I(out)/V(src) | V(src)=2*xfmag`

where `xfmag` defaults to 1 for each source type. For the port, `V(src)` is the internal source voltage.

Specifying `stimuli=nodes_and_terminals` indicates that all possible transfer functions should be computed. This is useful when it is not known in advance which transfer functions are interesting. Transfer functions for nodes are computed assuming that a unit magnitude flow (current) source is connected from the node to ground. Transfer functions for terminals are computed assuming that a unit magnitude potential (voltage) source is connected in series with the terminal. By default, the transfer functions from a small set of terminals are computed. If transfer functions from specific terminals are desired, specify the terminals in the save statement. You must use the `:probe` modifier (ex. `Rout:1:probe`) or specify `useprobes=yes` on the options statement. If transfer functions from all terminals are desired, specify `currents=all` and `useprobes=yes` on the options statement.

Spectre can perform the analysis while sweeping a parameter. The parameter can be frequency, temperature, component instance parameter, component model parameter, or netlist parameter. If changing a parameter affects the DC operating point, the operating point is recomputed on each step. You can sweep the circuit temperature by giving the parameter name as `temp` with no `dev` or `mod` parameter. You can sweep a netlist parameter by giving the parameter name with no `dev`, or `mod` parameter. After the analysis has completed, the modified parameter returns to its original value.

Definition

Name [p] [n] xf parameter=value ...

The optional terminals (p and n) specify the output of the circuit. If you do not give the terminals, then you must specify the output with a probe component.

Parameters

1 `prevoppoint=no` Use operating point computed on the previous analysis.
Possible values are `no` or `yes`.

Sweep interval parameters

2 `start=0` Start sweep limit.

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3	<code>stop</code>	Stop sweep limit.
4	<code>center</code>	Center of sweep.
5	<code>span=0</code>	Sweep limit span.
6	<code>step</code>	Step size, linear sweep.
7	<code>lin=50</code>	Number of steps, linear sweep.
8	<code>dec</code>	Points per decade.
9	<code>log=50</code>	Number of steps, log sweep.
10	<code>values=[...]</code>	Array of sweep values.

Sweep variable parameters

11	<code>dev</code>	Device instance whose parameter value is to be swept.
12	<code>mod</code>	Model whose parameter value is to be swept.
13	<code>param</code>	Name of parameter to sweep.
14	<code>freq (Hz)</code>	Frequency when parameter other than frequency is being swept.

Probe parameters

15	<code>probe</code>	Compute every transfer function to this probe component.
----	--------------------	--

State-file parameters

16	<code>readns</code>	File that contains estimate of DC solution (nodeset).
----	---------------------	---

Output parameters

17	<code>stimuli=sources</code>	Stimuli used for xf analysis. Possible values are <code>sources</code> or <code>nodes_and_terminals</code> .
----	------------------------------	---

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Analysis Statements

- 18 `save` Signals to output.
Possible values are `all`, `lvl`, `allpub`, `lvlpub`, `selected`, or `none`.
- 19 `nestlvl` Levels of subcircuits to output.
- 20 `oppoint=no` Should operating point information be computed, and if so, where should it be sent. Operating point information would not be output if operating point is computed in the previous analysis and is unchanged.
Possible values are `no`, `screen`, `logfile`, or `rawfile`.

Convergence parameters

- 21 `restart=yes` Restart the DC solution from scratch if any condition has changed. If not, use the previous solution as initial guess.
Possible values are `no` or `yes`.

Annotation parameters

- 22 `annotate=sweep` Degree of annotation.
Possible values are `no`, `title`, `sweep`, `status`, or `steps`.
- 23 `stats=no` Analysis statistics.
Possible values are `no` or `yes`.
- 24 `title` Analysis title.

You can specify sweep limits by giving the end points or by providing the center value and the span of the sweep. Steps can be linear or logarithmic, and you can specify the number of steps or the size of each step. You can give a step size parameter (`step`, `lin`, `log`, `dec`) to determine whether the sweep is linear or logarithmic. If you do not give a step size parameter, the sweep is linear when the ratio of stop to start values is less than 10, and logarithmic when this ratio is 10 or greater. All frequencies are in Hertz.

The small-signal analysis begins by linearizing the circuit about an operating-point. By default this analysis computes the operating-point if it is not known, or recomputes it if any significant component or circuit parameter has changed. However, if a previous analysis computed an operating point, you can set `prevoppoint=yes` to avoid recomputing it. For example, if you use this option when the previous analysis was a transient analysis, the operating point is the state of the circuit on the final time point.

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Parameter Index

In the following index, the number following each parameter name indicates where to find the description of that parameter.

annotate	22	log	9	probe	15	stats	23
center	4	mod	12	readns	16	step	6
dec	8	nestlvl	19	restart	21	stimuli	17
dev	11	oppoint	20	save	18	stop	3
freq	14	param	13	span	5	title	24
lin	7	prevopoint	1	start	2	values	10

Spectre Syntax

This chapter discusses the following topics:

- [Using analogmodel for Model Passing \(analogmodel\)](#) on page 855
- [Behavioural Source Use Model \(bsource\)](#) on page 856
- [Checkpoint - Restart \(checkpoint\)](#) on page 861
- [Configuring CMI Shared Objects \(cmiconfig\)](#) on page 862
- [Built-in Mathematical and Physical Constants \(constants\)](#) on page 863
- [Convergence Difficulties \(convergence\)](#) on page 864
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- [Sensitivity Analyses \(sens\)](#) on page 887
- [SpectreHDL Usage and Language Summary \(spectrehdl\)](#) on page 888
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- [Verilog-A Usage and Language Summary \(veriloga\)](#) on page 901

Using analogmodel for Model Passing (analogmodel)

Description

`analogmodel` is a reserved word in Spectre that allows you to bind an instance to different masters based on the value of a special instance parameter called `modelname`. An instance of `analogmodel` must have a parameter named `modelname` whose string value will be the name of the master this instance will be bound to. The value of `modelname` can be passed into subcircuits.

The `analogmodel` keyword is used by the Cadence Analog Design Environment to enable model name passing through the schematic hierarchy.

Sample Instance Statement

```
name [(]node1 ... nodeN[)] analogmodel modelname=mastername [[param1=value1]
    ...[paramN=valueN]]
```

<code>name</code>	Name of the statement or instance label.
<code>[(]node1...nodeN[)]</code>	Names of the nodes that connect to the component.
<code>analogmodel</code>	Special device name to indicate that this instance will have its master name specified by the value of the <code>modelname</code> parameter on the instance.
<code>modelname</code>	Parameter to specify the master of this instance indicated by <code>mastername</code> . The <code>mastername</code> must either be a valid string identifier or a netlist parameter that must resolve to a valid master name - a primitive, a model a subckt, or an AHDL module.
<code>param1</code>	Parameter values for the component. Depending on the master type, these can either be device parameters or netlist parameters. This is an optional field.

Example

```
//example spectre netlist to illustrate modelname parameter
simulator lang=spectre
parameters a="low" b="bottom"
ahdl_include "VerilogAStuff.va"
topInst1 (out in) top
```

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Spectre Syntax

```
topInst2 analogmodel modelname="VAMaster" //VAMaster is defined in
"VerilogASTuff.va"
topInst3 (out in) analogmodel modelname="resistor" //topInst3 binds to a
primitive
topInst4 (out in) analogmodel modelname="myOwnRes" //topInst4 binds to modelcard
"myOwnRes" defined below
topinst5 (out in) weiredRes modelname=modelname //modelname is just another
netlist param
model myOwnRes resistor r=100
subckt (out in) top
  parameters a="mid"
  x1 (out in) analogmodel modelname=a //topInst1.x1 binds to "mid"
ends top
subckt (out in) mid
  parameters c="low"
  x1 (out in) analogmodel modelname=b //topInst1.x1.x1 binds to "bottom"
  x2 (out in) analogmodel modelname=c //topInst1.x1.x1.x2 binds to "low"
ends mid
subckt (out in) low
  x1 (out in) analogmodel modelname="bottom" //topInst1.x1.x1.x2.x1 binds to
"bottom"
ends low
subckt (out in) bottom
  x1 (out in) analogmodel modelname="resistor" //x1 binds to primitive
"resistor"
ends bottom
dcl dc
```

Behavioural Source Use Model (bsource)

Description

Behavioral source enables you to model a resistor, inductor, capacitor, voltage or current source as a behavioral component. Using `bsource`, you can express the value of a resistance, capacitance, voltage or current as a combination of node voltages, branch currents, time expression, and built in Spectre expressions.

The syntax for `bsource` is:

```
name (node1 node2) bsource behav_param param_list
```

where `behav_param` can be

Spectre Circuit Simulator Reference

Spectre Syntax

`c=simple_expr` , Capacitance between the nodes

`g=simple_expr`, Conductance between the nodes

`i=generic_expr`, Current through bsource

`l=simple_expr`, Inductance between the nodes

`phi=simple_expr`, Flux in the bsource device

`q=simple_expr`, Charge in bsource device

`r=simple_expr`, Resistance between the nodes

`v=generic_expr`, Voltage across the nodes

`simple_expr` is defined as follows:-

A spectre expression containing,

1. netlist parameters
2. current simulation time, `$time`
3. node voltages, `v(a,b)`, where `a` and `b` are nodes in the spectre netlist or `v(a)`, which is voltage between node `a` and ground
4. branch currents, `i("inst_id:index")`, where `inst_id` is an instance name given in the netlist and `index` is the port index. The default value for `index` is 0.

`generic_expr` is defined as a `simple_expr` or `ddt()` or `idt()` of `simple_expr`

`param_list` is `param_name=value`

`param_name` can be

Multiplicity factor

`m` The value of `m` will be default to 1.

Temperature Parameters

`tc1` Linear temperature co-efficient. Valid for all behavioural elements.

Default value is 0 1/C.

`tc2` Quadratic temperature co-efficient. Valid for all behavioural elements.

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Spectre Syntax

Default value is 0 C^{-2}

tnom Parameters measurement temperature. Valid for all behavioural elements.

Default value is 0.0.

trise Temperature rise for ambient. Valid for all behavioural elements.

Default value is 0.0.

Clipping Parameters

max_val Maximum value of bsource expression. Valid for all behavioural elements, but generally used with i and v elements for clipping the current or voltage between the specified values.

min_val Minimum value of bsource expression. Valid for all behavioural elements, but generally used with i and v elements for clipping the current or voltage between the specified values.

Noise Parameters

af Flicker noise exponent, Valid for r and g elements

Default value is 2.

fexp Flicker noise frequency exponent. Valid for r, g, v, and i elements.

Default value is 1.

isnoisy Specifies whether to generate noise. Valid for r, g, i, and v elements

Valid values are yes and no. Default value is yes.

kf Flicker noise co-efficient. Valid for r and g elements.

white_noise White noise expression. Valid for v and i elements.

flicker_noise Flicker noise expression. Valid for v and i elements.

All the parameters in the param_name table are instance parameters. white_noise and flicker_noise may be assigned behavioural expressions; the other parameters must be assigned constant or parametric expressions.

Instance Parameters Supported

bsource supports the following instance parameters for the Spectre primitives.

Resistor isnoisy, m, r, tc1, tc2, and trise

Capacitor c, m, tc1, tc2, and trise.

Inductor l, m, tc1, tc2 and trise.

Mathematical Definitions

$$i = \text{ddt}(q) = \text{ddt}(\text{simple_expr})$$

$$v = \text{ddt}(\text{phi}) = \text{ddt}(\text{simple_expr})$$

$$v = i * r = i * \text{simple_expr}$$

$$i = g * v = \text{simple_expr} * v$$

$$i = \text{ddt}(c * v)$$

$$v = \text{ddt}(i * l)$$

Temperature effects on bsource:-

The equation for computing temperature factor is given as follows,

$$\text{tempFactor} = [1 + \text{tc1} * (\text{temp} - \text{trise} + \text{tnom}) + \text{tc2} * (\text{temp} - \text{trise} + \text{tnom})^2]$$

Examples

Non-linear resistor/capacitor/inductor modelling

```
res (n1 n2) bsource r=100*(1+(1/2)*v(n1,n2))
res (n1 n2) resistor r=100*(1+(1/2)*v(n1,n2))
cap (n1 n2) bsource c=1.0e-6*(1+(1/2)*v(n1,n2))
cap (n1 n2) capacitor c=1.0e-6*(1+(1/2)*v(n1,n2))
ind (n1 n2) bsource l=0.1*(1+(1/2)*v(n1,n2))
ind (n1 n2) inductor l=0.1*(1+(1/2)*v(n1,n2))
```

Charge model for capacitor

```
cap (n1 n2) bsource q=1.0e-6*v(n1,n2)
```

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Voltage and Current (Sinewave) Source

```
vsrc (n1 n2) bsource v=10.0*sin(2*pi*freq*$time)
isrc (n1 n2) bsource i=1.0e-3*sin(2*pi*freq*$time)
```

Current controlled current source

```
vsrc (n1 n2) vsource v=10
cccs1 (n3 n4) bsource i=gain*i("vsrc:0")
```

Current controlled voltage source

```
vsrc (n1 n2) vsource v=10
ccvs1 (n3 n4) bsource v=100*i("vsrc:0")
```

Voltage controlled voltage source

```
vsrc (n1 n2) resistor r=100k
vcvs1 (n3 n4) bsource v=gain*v(n1,n2)
```

Voltage controlled current source

```
vsrc (n1 n2) resistor r=100k
vccs1 (n3 n4) bsource i=v(n1,n2)/2000.0
```

Giving voltage clipping limit

```
res (n1 n2) bsource r=100*(1+(1/2)*v(n1,n2)) max_val=105 min_val=95
res (n1 n2) bsource r=100 tc1=0.01 tc2=0.003 trise=10 tnom=30
```

Doing Altergroup with bsource

```
vsrcl (n1 n2) bsource v=10*sin(2*pi*freq1*$time)
vsrc2 (n3 n4) bsource v=10*cos(2*pi*freq2*$time)
cccs1 (n5 n6) bsource i=gain*i("vsrcl:0")
res (n5 n6) bsource r=100*(1+(1/2)*v(n5,n6))
tran1 tran stop=1u
altAnal altergroup {
cccs1 (n5 n6) bsource i=gain*i("vsrc2:0")
res (n5 n6) bsource r=100*(1+(1/3)*pow(v(n5,n6),2))
}
tran2 tran stop=1u
```

Note: For resistor/capacitor/inductor, spectre netlist could directly be given with non-linear equation. However, they will internally be treated as behavioural source.

bsource Compilation

In this release the performance of bsource devices has been improved by performing a one time compilation step. The performance improvement obtained is proportional to the complexity of the bsource expression. Following the initial compilation, recompilation will only be performed if the bsource expression is changed.

Bsource compilation is enabled by default. If you are making frequent changes to bsource expressions

used in your design, the overhead of the compilation step may become an issue. To turn off compilation

set the CDS_AHDLCMI_ENABLE shell environment variable to NO e.g:

```
setenv CDS_AHDLCMI_ENABLE NO
```

To re-enable bsource compilation set the CDS_AHDLCMI_ENABLE to YES e.g:

```
setenv CDS_AHDLCMI_ENABLE YES
```

or undefine the CDS_AHDLCMI_ENABLE environment variable e.g:

```
unsetenv CDS_AHDLCMI_ENABLE
```

Checkpoint - Restart (checkpoint)

Description

Spectre has the ability to save checkpoint files while the analyses are running, and to restart an analysis from its checkpoint file. Checkpoint files can be generated in several ways:

- 1) Periodically based on real time (wall clock time).
- 2) Asynchronous UNIX signals.
- 3) By other methods unique to the analyses.

To generate checkpoint files periodically based on real time, set the Spectre option `ckptclock` to the time interval in seconds that you want checkpoints. This option is turned on by default with a value of 1800 seconds (30 minutes). Spectre will delete the checkpoint file if the simulation completes normally. If the simulation terminates abnormally, the checkpoint file will not be deleted.

If Spectre receives the UNIX signal `USR2`, then Spectre will immediately write a checkpoint file. If Spectre receives interrupt signals like `QUIT`, `TERM`, `INT`, or `HUP`, Spectre will attempt

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to write a checkpoint file and then exit. After other fatal signals, it may not be possible for Spectre to write a checkpoint file.

The name of the checkpoint file is a combination of the circuit name, the analysis name, and the extension `.ckpt`. For example, if the circuit is named `test1` and the transient analysis is named `timeSweep`, then the checkpoint file will be named `test1.timeSweep.tran.ckpt`.

Spectre keeps only the latest checkpoint file. When a new checkpoint is created, it creates the file under a temporary name. After the file has been successfully written, it deletes the previous checkpoint file and renames the new file.

Currently only the transient analyses supports checkpoint and restart.

Checkpoint

The transient analysis can generate checkpoint files by using the above methods, or by generating a checkpoint file periodically based on the transient simulation time. This is accessed by a transient analysis parameter called `ckptperiod`, which is turned off by default.

Restart

To restart an analysis from a checkpoint file, use the `+recover` option on the Spectre command line. Spectre will look through the requested analyses to see if a checkpoint file exists for any of them. If a checkpoint file for a given analysis does exist, Spectre will skip over any analyses previous to that one, and start the analysis using the information from the file.

Configuring CMI Shared Objects (cmiconfig)

Description

Spectre supports the ability to install devices dynamically from shared objects at run time. CMI Configuration files are used to determine and locate the set of shared objects to be installed. Spectre first reads the default CMI configuration file which specifies the default shared objects provided by Cadence. The configuration file specified by the value of the `CMI_CONFIG` environment variable is then read. The third configuration file that Spectre reads is `~/cmiconfig`. Finally, the configuration file specified in Spectres `-cmiconfig` command line argument is read. Each CMI configuration file modifies the existing configuration established by the configuration files read before.

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The following commands can be used in a CMI configuration file.

<code>setpath</code>	Specifies and resets the search path. <code>setpath path</code> or <code>setpath (path1 path2 ... pathN)</code>
<code>prepend</code>	Adds a path before the current search path. <code>prepend path</code> or <code>prepend (path1 path2 ... pathN)</code>
<code>append</code>	Adds a path after the current search path. <code>append path</code> or <code>append (path1 path2 ... pathN)</code>
<code>load</code>	Add a shared object to the list of shared objects to load. <code>loads path/ shared_object_name</code>
<code>unload</code>	Removes a shared object to the list of shared objects to load. <code>unload shared_object_name</code>

For example, given the following CMI configuration file

```
append /hm/spectre_dev/tools.sun4v/spectrecmi/lib/cmi/1.0
load libbjtx+tfet.so
load libmosx.so
```

The shared objects `libbjtx+tfet.so` and `libmosx.so` are loaded from `/hm/spectre_dev/tools.sun4v/spectrecmi/lib/cmi/1.0` in addition to the default shared objects provided by Cadence.

Built-in Mathematical and Physical Constants (constants)

Description

Spectre supports the following list of built-in mathematical and physical constants:

<code>M_</code>	is a mathematical constant	
<code>M_E</code>	2.7182818284590452354	$\exp(1) = e$
<code>M_LOG2E</code>	1.4426950408889634074	$\log_2(e)$
<code>M_LOG10E</code>	0.43429448190325182765	$\log_{10}(e)$
<code>M_LN2</code>	0.69314718055994530942	$\ln(2)$

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M_LN10	2.30258509299404568402	$\ln(10)$
M_PI	3.14159265358979323846	pi
M_TWO_PI	6.28318530717958647652	$2 * \pi$
M_PI_2	1.57079632679489661923	$\pi/2$
M_PI_4	0.78539816339744830962	$\pi/4$
M_1_PI	0.31830988618379067154	$1/\pi$
M_2_PI	0.63661977236758134308	$2/\pi$
M_2_SQRTPI	1.12837916709551257390	$2/\sqrt{\pi}$
M_SQRT2	1.41421356237309504880	$\sqrt{2}$
M_SQRT1_2	0.70710678118654752440	$\sqrt{1/2}$
M_DEGPERRAD	57.2957795130823208772	number of degrees per radian
P_	is a physical constant	
P_Q	1.6021918e-19	charge of electron in coulombs
P_C	2.997924562e8	speed of light in vacuum in meters/sec
P_K	1.3806226e-23	Boltzmanns constant in joules/kelvin
P_H	6.6260755e-34	Plancks constant in joules*sec
P_EPS0	8.85418792394420013968e-12	Permittivity of vacuum in farads/meter
P_U0	$(4.0e-7 * M_PI)$	Permeability of vacuum in henrys/meter
P_CELSIUS0	273.15	Zero celsius in kelvin

These constants can be used in expressions, or anywhere that a numeric value of expression is expected.

Convergence Difficulties (convergence)

Description

If you are having convergence difficulties, try the following suggestions:

1. Carefully evaluate and resolve any notice, warning or error messages.

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2. Assure topology checker is used (set `topcheck=full` on options statement) and heed any warnings it gives.
3. Perform sanity checking on the parameter values using the parameter range checker (use `+param param-limits-file` as a command line argument) and heed any warnings. Print the minimum and maximum parameter value using the `info` analysis. Assure that the bounds given for instance, model, output, temperature-dependent, and operating-point (if possible) parameters are reasonable.
4. Small floating resistors connected to high impedance nodes can cause convergence difficulties. Avoid very small floating resistors, particularly small parasitic resistors in semiconductors. Use voltage sources or iprobes to measure current rather than small resistors.
5. Use realistic device models. Check all component parameters, particularly nonlinear device model parameters, to assure that they are reasonable.
6. Increase the value of `gmin` (on options statement).
7. Loosen tolerances, particularly absolute tolerances like `iabstol` (on options statement). If tolerances are set too tight, they might preclude convergence.
8. Try to simplify the nonlinear component models in order to avoid regions in the model that may contribute to convergence problems.

DC Convergence Suggestions

Once you have a solution, write it to a nodeset file using the `write` parameter, and read it back in on subsequent simulations using the `readns` parameter.

1. If you have an estimate of what the solution should be, use nodeset statements or a nodeset file and set as many nodes as possible.
2. If convergence difficulties occur when using nodesets or initial conditions, try increasing `rforce` (on options statement).
3. If this is not the first analysis, perhaps the solution from the previous analysis is far from the solution for this analysis. If so, set `restart=yes`.
4. If simulating a bipolar analog circuit, assure the region parameter on all transistors and diodes is set correctly.

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5. If analysis fails at an extreme temperature, but succeeds at room temperature, then try adding a DC analysis that sweeps temperature. Start at room temperature, sweep to the extreme temperature, and write the final solution to a nodeset file.
6. Use numeric pivoting in the sparse matrix factorization by setting `pivotdc=yes` (on options statement). Sometimes it is also necessary to increase the pivot threshold to somewhere in the range of 0.1 to 0.5 using `pivrel` (on options statement).
7. Divide the circuit into smaller pieces and simulate them individually, but be careful to assure that the results will be close to what they would be if the rest of the circuit was present. Use the results to generate nodesets for the whole circuit.
8. If all else fails, replace the DC analysis with a transient analysis and modify all the independent sources to start at zero and ramp to their DC values. Run the transient analysis well beyond the time when all the sources have reached their final value (remember that transient analysis is very cheap when all of the signals in the circuit are not changing) and write the final point to a nodeset file. To make the transient analysis more efficient set the integration method to backward Euler (`method=euler`) and loosen the local truncation error criteria by increasing `lteratio`, say to 50. Occasionally, this approach will fail or be very slow because the circuit contains an oscillator. Often times the oscillation can be eliminated for the sake of finding the dc solution by setting the minimum capacitance from each node to ground (`cmin`) to a large value.

Transient Convergence Suggestions

1. Assure that a complete set of parasitic capacitors is used on nonlinear devices to avoid jumps in the solution waveforms. On MOS models, specify nonzero source and drain areas.
2. Use the `cmin` parameter to install a small capacitor from every node in the circuit to ground. This usually eliminates any jumps in the solution.

Export (export)

The export feature is not supported. It is designated for internal use only.

Expressions (expressions)

Description

An expression is a construct that combines operands with operators to produce a result that is a function of the values of the operands and the semantic meaning of the operators. Any legal operand is also an expression in itself. Legal operands include numeric constants and references to top-level netlist parameters or subcircuit parameters. Calls to algebraic and trigonometric functions are also supported. The complete lists of operators, algebraic, and trigonometric functions are given after some examples.

Examples

```
simulator lang=spectre
parameters p1=1 p2=2           // declare some top-level parameters
r1 (1 0) resistor r=p1         // the simplest type of expression
r2 (1 0) resistor r=p1+p2      // a binary (+) expression
r3 (1 0) resistor r=5+6/2      // expression of constants, = 8
x1 s1 p4=8                     // instantiate a subcircuit, defined in the following lines
subckt s1
parameters p1=4 p3=5 p4=6     // subcircuit parameters
    r1 (1 0) resistor r=p1     // another simple expression
    r2 (1 0) resistor r=p2*p2  // a binary multiply expression
    r3 (1 0) resistor r=(p1+p2)/p3 // a more complex expression
    r4 (1 0) resistor r=sqrt(p1+p2) // an algebraic function call
    r5 (1 0) resistor r=3+atan(p1/p2) // a trigonometric function call
    r6 (1 0) RESMOD r=(p1 ? p4+1 : p3) // the ternary operator
ends
// a model card, containing expressions
model RESMOD resistor tc1=p1+p2 tc2=sqrt(p1*p2)
// some expressions used with analysis parameters
time_sweep tran start=0 stop=(p1+p2)*50e-6 // use 5*50e-6 = 150 us
// a vector of expressions (see notes on vectors below)
dc_sweep dc param=p1 values=[0.5 1 +p2 (sqrt(p2*p2)) ] // sweep p1
```

Where Expressions Can Be Used:

The Spectre native netlist language allows expressions to be used where numeric values are expected on the right-hand side of an "=" sign, or within a vector, where the vector itself is on the right-hand side of an "=" sign. Expressions can be used when specifying device or

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analysis instance parameter values (for example specifying the resistance of a resistor or the stop time of a transient analysis, as outlined in the preceding example), when specifying model parameter values in model cards (for example specifying "bf=p1*0.8" for a bipolar model parameter, bf), or when specifying initial conditions and nodesets for individual circuit nodes.

Operators

The following operators are supported, listed in order of decreasing precedence. Parentheses can be used to change the order of evaluation. For a binary expression like "a+b", "a" is the first operand and "b" is the second operand. All operators are left associative, with the exceptions of the "to the power of" operator (**) and the ternary operator (? :), which are right associative. For logical operands, any nonzero value is considered true. The relational and equality operators return a value of 1 to indicate true or 0 to indicate false. There is no short circuiting of logical expressions involving && and ||.

Operator	Symbol(s)	Value
Unary +, Unary -	+, -	Value of operand, negative of operand.
To the power of	**	First operand to raised to power of second operand.
Multiply, Divide	*, /	Product, Quotient of operands.
Binary Plus/Minus	+, -	Sum, Difference of operands
Shift	<<, >>	First operand shifted left (, right) by number of bits specified by second operand.
Relational	<, <=, >, >=	Less than, less than or equal, greater than, greater than or equal.
Equality	==, !=	True if operands are equal, not equal.
Bitwise AND	&	Bitwise AND (of integer operands).
Bitwise Exclusive NOR	~^ (or ^~)	Bitwise Exclusive NOR (of integer operands).
Bitwise OR		Bitwise OR (of integer operands).
Logical AND	&&	True only if both operands true.
Logical OR		True if either operand is true.
Ternary Operator	(cond) ? x : y	Returns x if cond is true, y if not where x and y are expressions.

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Algebraic and Trigonometric Functions

The trigonometric and hyperbolic functions expect their operands to be specified in radians. The atan2() and hypot() functions are useful for converting from Cartesian to polar form.

Function	Description	Domain
log(x)	Natural logarithm	$x > 0$
log10(x)	Decimal logarithm	$x > 0$
exp(x)	Exponential	$x < 80$
sqrt(x)	Square Root	$x > 0$
min(x,y)	Minimum value	All x, all y
max(x,y)	Maximum value	All x, all y
abs(x)	Absolute value	All x
pow(x,y)	x to the power of y	All x, all y
int(x)	Integer value of x	All x
floor(x)	Argest integer $\leq x$	All x
ceil(x)	Smallest integer $\geq x$	All x
fmod(x,y)	Floating point modulus	All x, all y, except $y=0$
int(x)	Integer value of x	All x
floor(x)	Largest integer $\leq x$	All x
ceil(x)	Smallest integer $\geq x$	All x
fmod(x,y)	Floating point modulus	All x, all y, except $y=0$
sin(x)	Sine	All x
cos(x)	Cosine	All x
tan(x)	Tangent	All x, except $x = n*(\pi/2)$, where n odd
asin(x)	Arc-sine	$-1 \leq x \leq 1$
acos(x)	Arc-cosine	$-1 \leq x \leq 1$
atan(x)	Arc-tangent	All x
atan2(x,y)	Arc-tangent of x/y	All x, all y

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hypot(x,y)	$\sqrt{x^2 + y^2}$	All x, all y
sinh(x)	Hyperbolic sine	All x
cosh(x)	Hyperbolic cosine	All x
tanh(x)	Hyperbolic tangent	All x
asinh(x)	Arc-hyperbolic sine	All x
acosh(x)	Arc-hyperbolic cosine	$x \geq 1$
atanh(x)	Arc-hyperbolic tangent	$-1 \leq x \leq 1$

User-defined functions are also supported. See `spectre -h functions` for a description of user-defined functions.

A large number of built-in mathematical and physical constants are available for use in expressions. See `spectre -h constants` for the list of these constants

Using Expressions in Vectors

Expressions can be used as vector elements, as in the following example:

```
dc_sweep dc param=p1 values=[0.5 1 +p2 (sqrt(p2*p2)) ] // sweep p1
```

Note that when expressions are used within vectors, anything other than constants, parameters, or unary expressions (unary +, unary -) must be surrounded by parentheses. Vector elements should be space separated for clarity, though this is not mandatory. The preceding "dc_sweep" example shows a vector of four elements, namely 0.5, 1, +p2, and $\sqrt{p2 \cdot p2}$. Note that the square root expression is surrounded by parentheses.

User Defined Functions (functions)

Description

Spectre's user-defined function capability allows you to build upon the provided set of built-in mathematical and trigonometric functions. You can write your own functions, and call these functions from within any expression. The syntax for calling a user-defined function is the same as the syntax for calling a built-in algebraic or trigonometric function. Note that user-defined functions must be defined before they are referenced (called). Arguments to user-defined functions will be taken as real values, and the functions will return real values. A user-defined function may contain only a single statement in braces and this statement must return an expression (which will typically be an expression involving the function arguments). The

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return expression may reference the built in parameters `temp` and `tnom`. User-defined functions must be declared at the top level only, and must not be declared within subcircuits. User-defined functions may be called from anywhere that an expressions can be currently used in Spectre. User-defined functions may call other functions (both user-defined and built-in), however any user-defined function will need to be declared before it can be called. User-defined functions can override built-in mathematical and trigonometric functions.

NOTE: Only real values for arguments and return values are supported in this release.

See `spectre -h expressions` for a list of built-in algebraic and trigonometric functions.

Definition

```
real myfunc( [real arg1, ...real argn] ) {
```

Examples

```
real myfunc( real a, real b ) {
    return a+b*2+sqrt(a*sin(b));
}
```

An example of a function calling a previously defined function follows

```
real yourfunc( real a, real b ) {
    return a+b*myfunc(a,b);    // call "myfunc"
}
```

The final example shows how a user-defined function may be called from an expression in the Spectre netlist:

```
r1 (1 0) resistor r=myfunc(2.0, 4.5)
```

Global Nodes (global)

Description

The global statement allows a set of nodes to be designated as common to the main circuit and all subcircuits. Thus, components inside subcircuits can be attached to global nodes even though the subcircuits terminals are not attached to these nodes.

Any number of global nodes may be specified using the global statement. To do this, follow the keyword `global` with a list of the node names that you wish to declare as global. The first node name that appears in this list is taken to be the name of the ground node. Ground is also

known as the datum or reference node. If a global statement is not used, 0 is taken to be the name of the ground node.

At most one global statement is allowed and, if present, it must be the first statement in the file (however, you can have `simulator lang=spectre` statement, before the global statement so that you can use mixed case names for the node names). Ground is always treated as global even if a global statement is not used.

Definition

```
global <ground
```

Initial Conditions (ic)

Description

The `ic` statement is used to provide initial conditions for nodes in the transient analysis. It can occur multiple times in the input and the information provided in all the occurrences is collected. Initial conditions will only be accepted for inductor currents and node voltages where the nodes have a path of capacitors to ground. For more information, read the description of transient analysis. It should be noted that specifying `cmin` for a transient analysis, will not satisfy the condition that a node has a capacitive path to ground.

Definition

```
ic <node=value
```

This statement takes a list of signals as an argument. The concept of nodes has been generalized to signals where a signal is a value associated with a topological node of the circuit or some other unknown that is solved by the simulator, such as the current through an inductor or the voltage of the internal node in a diode. Topological nodes can be either at the top-level or in a subcircuit.

For example,

```
ic 7=0 out=1 OpAmp1.comp=5 L1:1=1.0u
```

where `7=0` implies that node 7 should start at 0V, node `out` should start at 1V, node `comp` in subcircuit `OpAmp1` should start at 5V, and the current through the first terminal of `L1` should start at 1uA.

The Structural if-statement (if)

Description

The structural if-statement can be used to conditionally instantiate other instance statements.

Definition

```
if <condition
```

The condition is a boolean expression based on the comparisons of various arithmetic expressions which are evaluated during circuit hierarchy flattening. The `statement1` and `statement2` fields can be ordinary instance statements, if-statements, or a list of these within braces (`{}`). (Note that ordinary instance statements need a newline to terminate them.) The `else` part is optional. When if-statements are nested without braces, an `else` matches the closest previous unmatched `if` at the same level.

- It is possible to have duplicate instance names within the if statement under strict topological conditions. These are:
 - References to instance with duplicate names is only possible within a structural if statement which has both an "if" part and an "else" part.
 - both the "if" part and the "else" part must be either a simple one-statement block, or another structural if statement to which these same rules apply.
 - The duplicate instances must have the same number of terminals and be bound to the same list of nodes.
 - The duplicate instances must refer to the same primitive or model.
 - Where duplicate instances refer to a model, the underlying primitive must be the same.

This feature allows automatic model selection based on any netlist or subcircuit parameter. As an example, consider using Spectres inline subcircuits and structural if statement to implement automatic model selection based on bipolar device area. Here, the duplicate instances are the inline components.

```
model npn_default bjt is=3.2e-16 va=59.8
model npn10x10 bjt is=3.5e-16 va=61.5
model npn20x20 bjt is=3.77e-16 va=60.5
// npn_mod chooses scaled models binned on area!
// if ( area < 100e-12 ) use model npn10x10
// else if ( area < 400e-12 ) use model npn20x20
// else use model npn_default
```

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```
inline subckt npn_mod (c b e s)
  parameters area=5e-12
  if ( area < 100e-12 ) {
    npn_mod (c b e s) npn10x10 // 10u * 10u, inline device
  } else if ( area < 400e-12 ) {
    npn_mod (c b e s) npn20x20 // 20u * 20u, inline device
  } else {
    npn_mod (c b e s) npn_default // 5u * 5u, inline device
  }
ends npn_mod
q1 (1 2 0 0) npn_mod area=350e-12 // gets 20x20 model
q2 (1 3 0 0) npn_mod area=25e-12 // gets 10x10 model
q3 (1 3 0 0) npn_mod area=1000e-12 // gets default model
```

Include File (include)

Description

File inclusion allows the circuit description to be spread over several files. The include statement itself is replaced by the contents of the file named. An included file may also contain include statements. If the name given is not an absolute path specification, then it is taken relative to the directory of the file currently being read.

In order to read existing SPICE library and model files, Spectre automatically switches to SPICE input mode when it opens an include file. Thus, all files that use the Spectre native language must begin with a `simulator lang=spectre` statement. The one exception is files that end with a ".scs" file extension which are treated specially and are read in Spectre input mode. This language mode treatment applies to files included by both Spectres include statement, and CPPs #include statement.

After reading the include file, Spectre restores the language processing mode to what it was before the file was included, and continues reading the original file starting at the line after the include statement. Lines cannot be continued across file boundaries.

The CPP #include statement differs from Spectres include statement in that CPP macro processing will not be performed on files included by Spectre, but will be performed on files included by CPP. If your netlist contains a #include statement, you must run CPP to perform this inclusion, otherwise an error will occur.

Spectre Circuit Simulator Reference

Spectre Syntax

If the file to be included cannot be found in the same directory as the including file, Both Spectres include and CPPs #include will search for the file to be included along the search path specified by the -I command line arguments.

Spectres include statement allows you to embed special characters in the name of the file to be included. Spectres include statement will automatically expand the "~" character to the users home directory, and will expand environment variables and % codes, such as

```
include "~/models/${SIMULATOR}_pd/npn.scs"
```

which will look in the directory given by the environment variable SIMULATOR followed by "_pd", which is under the "models" directory, in the users home directory. Note: These special character features are not available using CPPs #include statement.

Definition

```
include "filename"
```

Spectre Netlist Keywords (keywords)

Description

The following lists the reserved Spectre keywords, including netlist keywords, built-in algebraic and trigonometric functions, and built-in mathematical and physical constants. When creating a netlist, you should avoid using any of the keywords from this list in any context other than that in which it was intended. Creating nodenames, parameter names, instance names, or model names from any of these keywords will result in an error.

Keyword	Keyword Type
M_1_PI	Mathematical Constant
M_2_PI	Mathematical Constant
M_2_SQRTPI	Mathematical Constant
M_DEGPER RAD	Mathematical Constant
M_E	Mathematical Constant
M_LN10	Mathematical Constant
M_LN2	Mathematical Constant

Spectre Circuit Simulator Reference

Spectre Syntax

M_LOG10E	Mathematical Constant
M_LOG2E	Mathematical Constant
M_PI	Mathematical Constant
M_PI_2	Mathematical Constant
M_PI_4	Mathematical Constant
M_SQRT1_2	Mathematical Constant
M_SQRT2	Mathematical Constant
M_TWO_PI	Mathematical Constant
P_C	Mathematical Constant
P_CELSIUS0	Mathematical Constant
P_EPS0	Mathematical Constant
P_H	Mathematical Constant
P_K	Mathematical Constant
P_Q	Mathematical Constant
P_U0	Mathematical Constant
abs	Algebraic Function
acos	Trigonometric Function
acosh	Trigonometric Function
altergroup	Netlist Keyword
asin	Trigonometric Function
asinh	Trigonometric Function
atan	Trigonometric Function
atan2	Trigonometric Function
atanh	Trigonometric Function
ceil	Algebraic Function
correlate	Netlist Keyword
cos	Trigonometric Function
cosh	Trigonometric Function
else	Netlist Keyword

Spectre Circuit Simulator Reference

Spectre Syntax

end	Netlist Keyword
ends	Netlist Keyword
exp	Algebraic Function
export	Netlist Keyword
floor	Algebraic Function
fmod	Algebraic Function
for	Netlist Keyword
function	Netlist Keyword
global	Netlist Keyword
hypot	Algebraic Function
ic	Netlist Keyword
if	Netlist Keyword
inline	Netlist Keyword
int	Algebraic Function
library	Netlist Keyword
local	Netlist Keyword
log	Algebraic Function
log10	Algebraic Function
march	Netlist Keyword
max	Algebraic Function
min	Algebraic Function
model	Netlist Keyword
nodeset	Netlist Keyword
parameters	Netlist Keyword
paramset	Netlist Keyword
plot	Netlist Keyword
pow	Algebraic Function
print	Netlist Keyword
pwr	Netlist Keyword

Spectre Circuit Simulator Reference

Spectre Syntax

real	Netlist Keyword
return	Netlist Keyword
save	Netlist Keyword
sens	Netlist Keyword
sin	Trigonometric Function
sinh	Trigonometric Function
sqrt	Algebraic Function
statistics	Netlist Keyword
subckt	Netlist Keyword
tan	Trigonometric Function
tanh	Trigonometric Function
to	Netlist Keyword
truncate	Netlist Keyword
vary	Netlist Keyword

Library - Sectional Include (library)

Description

Library inclusion allows the circuit description to be spread over several files. The library statement itself is replaced by the contents of the section of the library file. A library section may also contain library reference statements. If the file name given is not an absolute path specification, then it is taken relative to the directory of the file currently being read.

There are two kinds of library statements. One that references a library section, and one that defines a library section. The definition of a library section is prohibited in the netlist.

In order to read existing SPICE library and model files, Spectre automatically switches to SPICE input mode when it opens a library file. Thus, all files that use the Spectre native language must contain a `simulator lang=spectre` statement within each section of the library or the file can have a `scs` filename extension. After reading the library section, Spectre restores the language processing mode and continues reading the original file starting at the line after the library statement. Lines cannot be continued across file boundaries.

Spectre Circuit Simulator Reference

Spectre Syntax

Spectre allows only one library per file, but a library may contain multiple sections (typically one section per process corner for example.)

Definition

Inside netlist (reference library section)

Sample Library

```
library corner_lib
```

```
section tt
```

```
model nch bsim3v3 type=n mobmod=1 capmod=2 version=3.1
+ xj=1.7e-7 vsat=7.99e4 at=3.6e4 a0=0.799 ags=0.4
+ a1=0 a2=1 keta=-0.05 nch=2.8e17 ngate=1.31e20 k1=0.74
model pch bsim3v3 type=p mobmod=1 capmod=2 version=3.1
+ xj=1.7e-7 vsat=1.38e5 at=1e5 a0=1.3 ags=0.3
+ a1=1.1e-4 a2=1 keta=0 nch=4.1e17 ngate=7.6e19 k1=0.88
model knpn bjt is=10e-13 bf=170 va=58.7 ik=5.63e-3 rb=rbn rbm=86
+ re=3.2 cje=0.25e-12 pe=0.76 me=0.34 tf=249e-12 cjc=0.34e-12 pc=0.55
+ mc=0.35 ccs=2.4e-12 ms=0.35 ps=0.53 rc=169
model kpnp bjt type=PNP is=10e-13 bf=60 va=43.1 ik=0.206e-3 rb=rpb rbm=64.3
+ re=33.8 cje=0.16e-12 pe=0.5 me=0.26 tf=36e-9 cjc=0.72e-12 pc=0.58
+ mc=0.34 ccs=2.5e-12 ps=0.53 ms=0.35 rc=276
```

```
endsection
```

```
section ss
```

```
model nch bsim3v3 type=n mobmod=1 capmod=2 version=3.1
+ xj=1.7e-7 vsat=7.99e4 at=3.6e4 a0=0.799 ags=0.4
+ a1=0 a2=1 keta=-0.05 nch=2.8e17 ngate=1.31e20 k1=0.74
model pch bsim3v3 type=p mobmod=1 capmod=2 version=3.1
+ xj=1.7e-7 vsat=1.38e5 at=1e5 a0=1.3 ags=0.3
+ a1=1.1e-4 a2=1 keta=0 nch=4.1e17 ngate=7.6e19 k1=0.88
model knpn bjt is=10e-13 bf=70 va=58.7 ik=5.63e-3 rb=rbn rbm=86
+ re=3.2 cje=0.25e-12 pe=0.76 me=0.34 tf=249e-12 cjc=0.34e-12 pc=0.55
+ mc=0.35 ccs=2.4e-12 ms=0.35 ps=0.53 rc=169
model kpnp bjt type=PNP is=10e-13 bf=30 va=43.1 ik=0.206e-3 rb=rpb rbm=64.3
+ re=33.8 cje=0.16e-12 pe=0.5 me=0.26 tf=36e-9 cjc=0.72e-12 pc=0.58
+ mc=0.34 ccs=2.5e-12 ps=0.53 ms=0.35 rc=276
```

```
endsection
```

```
section ff
```

```
model nch bsim3v3 type=n mobmod=1 capmod=2 version=3.1
+ xj=1.7e-7 vsat=7.99e4 at=3.6e4 a0=0.799 ags=0.4
```

Spectre Circuit Simulator Reference

Spectre Syntax

```
+ a1=0 a2=1 keta=-0.05 nch=2.8e17 ngate=1.31e20 k1=0.74
model pch bsim3v3 type=p mobmod=1 capmod=2 version=3.1
+ xj=1.7e-7 vsat=1.38e5 at=1e5 a0=1.3 ags=0.3
+ a1=1.1e-4 a2=1 keta=0 nch=4.1e17 ngate=7.6e19 k1=0.88
model knpn bjt is=10e-13 bf=220 va=58.7 ik=5.63e-3 rb=rbn rbm=86
+ re=3.2 cje=0.25e-12 pe=0.76 me=0.34 tf=249e-12 cjc=0.34e-12 pc=0.55
+ mc=0.35 ccs=2.4e-12 ms=0.35 ps=0.53 rc=169
model kpnp bjt type=PNP is=10e-13 bf=90 va=43.1 ik=0.206e-3 rb=rbp rbm=64.3
+ re=33.8 cje=0.16e-12 pe=0.5 me=0.26 tf=36e-9 cjc=0.72e-12 pc=0.58
+ mc=0.34 ccs=2.5e-12 ps=0.53 ms=0.35 rc=276
endsection
endlibrary
```

Node Sets (nodeset)

Description

The `nodeset` statement is used to provide an initial guess for nodes in any DC analysis or the initial condition calculation for the transient analysis. It can occur multiple times in the input, the information provided in all the occurrences is collected. For more information, read the description of DC analysis.

Definition

```
nodeset <node=value
```

This statement takes a list of signals as an argument. The concept of nodes has been generalized to signals where a signal is a value associated with a topological node of the circuit or some other unknown that is solved by the simulator, such as the current through an inductor or the voltage of the internal node in a diode. Topological nodes can be either at the top-level or in a subcircuit.

For example,

```
nodeset 7=0 out=1 OpAmp1.comp=5 L1:1=1.0u
```

where `7=0` implies that node 7 should be about 0V, node `out` should be about 1V, node `comp` in subcircuit `OpAmp1` should be about 5V, and the current through the first terminal of `L1` should be about 1uA.

Parameter Soft Limits (param_limits)

Description

The parameter values passed to Spectre components and analysis are subject to both hard and soft limits. If you set a parameter to a value that violates a hard limit, such as giving $z0=0$ to a transmission line, Spectre issues an error message and quits. If the given parameter value violates a soft limit, only a warning is issued, but Spectre uses the value of the component as given. Hard limits are used to prevent you from using values that would cause Spectre to fail or put a model in an invalid region. Soft limits are used to call attention to unusual parameter values that might have been given mistakenly. If a parameter value violates a soft limit, a message similar to one of the following sample messages is printed:

Parameter `rb` has the unusually small value of 1uOhms.

or

Parameter `rb` has the unusually large value of 1MOhms.

Spectre has built-in soft limits on a few parameter values. However, it is possible for you to override these limits, or provide limits on parameters that do not have built-in limits. To do so, create a parameter range limits file, and invoke Spectre giving the name of the file after the `+param` command line option. For example,

```
spectre +param limits-file input-file
```

Limits are given using the following syntax:

```
[PrimitiveName] [model] [LowerLimit <[=]] [|]Param[|] [<[=] UpperLimit]
```

The limits can be given as strict (using `<=`) or nonstrict (using `<`). If the limits are strict, there can be no space between `<` and `=`. The limits for one parameter are given on one line. There is no way of continuing the specification of the limits for a parameter over more than one line. If a parameter is given more than once, the limits given the last time override earlier limits. The primitive name must be a Spectre primitive name, not a name used for SPICE compatibility. So, for example, `mos3` must be used rather than `mos`. Parameter limits can be written using Spectre native mode metric scale factors. Thus a limit of `f <= 1.0e6` could also be written as `f <= 1M`.

Examples

```
mos3      0.5u <= l <= 100u
          0.5u <= w
          0 < as <= 1e-8
          0 < ad <= 1e-8
```

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Spectre Syntax

```
model |vto| <= 3
```

Notice that it is not necessary to give the primitive name each time. If not given, it is assumed to be the same as the previous parameter. Upper and lower limits may be given, and if not given there will be no limit on the parameter value. Thus, in the example, if w is less than 0.5 μm , a warning will be issued, but there is no limit on how large w can be. If a parameter is mentioned, but no limits given, then all limits are disabled for that parameter. Limits are placed on model parameters by giving the model keyword. If the model keyword is not given, the limits are applied to instance parameters. Notice that you can also place upper or lower limits on the absolute value of a parameter. For example,

```
resistor 0.1 < |r| < 1M
```

indicates that the absolute value of r should be greater than 0.1 Ohm and less than 1 MOhm. There can be no spaces between the absolute value symbols and the parameter name.

Examples

```
1 <= x < 0.5
```

```
1 <= y <= 1
```

```
1 < z < 1
```

In the first case the lower bound is larger than the upper bound, which indicates that the range of x is all real numbers except those from 0.5 to 1 and 0.5 itself. The limits are applied separately, thus x must be both greater than or equal to 1 ($1 \leq x$) and less than 0.5 ($x < 0.5$). The second case specifies that y should be 1 and the third case specifies that z should not be 1.

It is possible to specify limits for any scalar parameter that takes either a real number, an integer, or an enumeration. To specify the limits of a parameter that takes enumerations use the indices associated with the enumerations. For example, consider the region parameter of the bjt. There are four possible regions: off, fwd, rev, or sat (see `spectre -help bjt`). Each enumeration is assigned a number starting at 0 and counting up. Thus, off=0, fwd=1, rev=2, and sat=3. The specification

```
bjt          3 <= region <= 1
```

indicates that a warning should be printed if region=rev because the conditions ($3 \leq \text{region}$) and ($\text{region} \leq 1$) exclude only (region=2) and region 2 is rev.

It is possible to read a parameter limits file from within another file. To do so, use an include statement. For example,

```
include "filename"
```

will temporarily suspend the reading of the current file until the contents of `filename` have been read. Include statements may be nested arbitrarily deep with the condition that the operating system may limit the number of files that Spectre may have open at once. Paths in

Spectre Circuit Simulator Reference

Spectre Syntax

file names are taken to be relative to the directory that contains the current file, not from the directory in which Spectre was invoked.

Spectre can be instructed to always read a parameter limits file by using the SPECTRE_DEFAULTS environment variable. For example, if you put the following in your shell initialization file (.profile for sh, .cshrc for csh)

```
setenv SPECTRE_DEFAULTS "+param /cds/etc/spectre/param.lmts"
```

Spectre would always read the specified limits file.

Netlist Parameters (parameters)

Description

The Spectre native netlist language allows parameters to be specified and referenced in the netlist, both at the top-level scope and within subcircuit declarations (run `spectre -h subckt` for more details on parameters within subcircuits).

Definition

```
parameters <param=value
```

Examples:

```
simulator lang=spectre
parameters p1=1 p2=2           // declare some parameters
r1 (1 0) resistor r=p1         // use a parameter, value=1
r2 (1 0) resistor r=p1+p2     // use parameters in an expression, value=3
x1 s1 p4=8                    // "s1" is defined below, pass in value 8 for "p4"
subckt s1
parameters p1=4 p3=5 p4=6     // note: no "p2" here, p1 "redefined"
r1 (1 0) resistor r=p1         // local definition used: value=4
r2 (1 0) resistor r=p2         // inherit from parent(top-level) value=2
r3 (1 0) resistor r=p3         // use local definition, value=5
r4 (1 0) resistor r=p4         // use passed-in value, value=8
r5 (1 0) resistor r=p1+p2/p3  // use local+inherited/local = (4+2/5) = 4.4
ends
time_sweep tran start=0 stop=(p1+p2)*50e-6 // use 5*50e-6 = 150 us
dc_sweep dc param=p1 values=[0.5 1 +p2 (sqrt(p2*p2)) ] // sweep p1
```

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Spectre Syntax

Parameter Declaration

Parameters can be declared anywhere in the top-level circuit description or on the first line of a subcircuit definition. Parameters must be declared before they are used (referenced). Multiple parameters can be declared on a single line. When parameters are declared in the top-level, their values must be specified. When parameters are declared within subcircuits, their default values are optionally specified.

Parameter Inheritance

Subcircuit definitions inherit parameters from their parent (enclosing subcircuit definition, or top-level definition). This inheritance continues across all levels of nesting of subcircuit definitions, that is, if a subcircuit `s1` is defined, which itself contains a nested subcircuit definition `s2`, then any parameters accessible within the scope of `s1` are also accessible from within `s2`. Also, any parameters declared within the top-level circuit description are also accessible within both `s1` and `s2`. However, any subcircuit definition can redefine a parameter that it has inherited. In this case, if no value is specified for the redefined parameter when the subcircuit is instantiated, then the redefined parameter uses the locally defined default value, rather than inheriting the actual parameter value from the parent.

Parameter Namespace

Parameter names must not conflict with device or analysis instance names, that is, it is not possible to reference a parameter called `r1` if there is an instance of a resistor (or other device or analysis) called `r1`. Parameter names must also not be used where a node name is expected.

Parameter Referencing

Spectre netlist parameters can be referenced anywhere that a numeric value is normally specified on the right-hand side of an "=" sign or within a vector, where the vector itself is on the right-hand side of an "=" sign. This includes referencing of parameters in expressions (run `spectre -h expressions` for more details on netlist expression handling), as indicated in the preceding examples. You can use expressions containing parameter references when specifying device or analysis instance parameter values (for example specifying the resistance of a resistor or the stop time of a transient analysis, as outlined in the preceding example), when specifying model parameter values in model cards (for example specifying "`bf=p1*0.8`" for a bipolar model parameter, `bf`), or when specifying initial conditions and nodesets for individual circuit nodes.

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Spectre Syntax

Altering/Sweeping Parameters

Just as certain Spectre analyses (for example `sweep`, `alter`, `ac`, `dc`, `noise`, `sp`, `xf`) can sweep device instance or model parameters, they can also sweep netlist parameters. Run `spectre -h <analysis>` to see the particular details for any of these analyses, where `<analysis>` is the analysis of interest.

Temperature as a parameter

You can use the reserved parameters `temp` and `tnom` anywhere that an expression can be used, including within expressions and user-defined functions. The `temp` parameter always represents the simulator (circuit) temperature, and `tnom` always represents the measurement temperature. All expressions involving `temp` or `tnom` are re-evaluated any time the circuit temperature or measurement temperature changes.

You can also alter or sweep the `temp` and `tnom` parameters using any of the techniques available for altering or sweeping netlist or subcircuit parameters (with the exception of `altergroups`).

This capability allows you to write temperature dependent models for example, by using `temp` in an equation for a model or instance parameter. For example,

```
r1 1 0 res r=(temp-tnom)*15+10k // temp is temperature
o1 options temp=55           // causes a change in above resistor r1
```

Reserved Parameters

The following parameters are reserved, and may not be declared as either top-level parameters or subcircuit parameters: `temp`, `tnom`, `scale`, `scalem`, `freq`, `time`.

Parameter Set - Block of Data (paramset)

Description

A parameter set is a block of data, which can be referenced by a sweep analysis. Within a `paramset` the first row contains an array of top-level netlist parameters. All other rows contain numbers which are used to alter the value of the parameters during the sweep. Each row represents an iteration of the sweep. This data should be bound within braces. The opening brace is required at the end of the line defining the `paramset`. The `paramset` cannot be defined within subcircuits or cannot be nested.

Spectre Circuit Simulator Reference

Spectre Syntax

Definition

<Name

Example

```
data paramset {
  p1 p2 p3
  1.1 2.2 3.3
  4.4 5.5 6.6
}
```

Output Selections (save)

Description

The `save` statement indicates that the values of specific nodes or signals should be saved in the output file. It works in conjunction with the `save` parameter on most analyses. The output file is written in Cadence Waveform Storage Format (WSF), Cadence Parameter Storage Format (PSF) or in Nutmeg/SPICE3 format as controlled by a a command line argument or a global option (see the options statement). The proper postprocessor should be used to view the output, generate plots, or do any further processing.

Definition

```
save <node|component|terminal
```

This statement takes a list of signals as an argument. The concept of nodes has been generalized to signals where a signal is a value associated with a topological node of the circuit or some other unknown that is solved by the simulator, such as the current through an inductor or the voltage of the internal node in a diode. Topological nodes can be either at the top-level or in a subcircuit.

For example,

```
save 7 out OpAmp1.comp M1:currents D3:oppoint L1:1 R4:pwr
```

which tells that node 7, node `out`, node `comp` in subcircuit `OpAmp1`, the currents through the terminals of `M1`, the `oppoint` information for diode `D3`, the current through the first terminal of `L1`, and the instantaneous power dissipated by `R4` should be saved. These outputs are saved in addition to any outputs specified with the `save` parameter on the analysis.

To specify a component terminal current, give the name of the component and the name or the index of the terminal separated by a colon. If `currents` is specified after the component and the colon, then all the terminal currents for the component are saved unless the

component has only two terminals, in which case only the current through the first terminal is saved. Current is positive if it enters the terminal flowing into the component.

If a component name is followed by a colon and `oppoint`, then the operating point information associated with the component is computed and saved. If the colon is followed by an operating point parameter name (see each component for list of operating point parameters), then the value of that parameter is output.

If only a component name is given, all available information about the component, including the terminal currents and the operating point parameter values, is saved.

Sensitivity Analyses (sens)

Description

Use the `sens` control statement to find partial or normalized sensitivities of the output variables with respect to component and instance parameters for the list of the analyses performed. Currently DC and AC sensitivity analyses are supported. The results of the sensitivity analyses are stored in the output files written in Cadence Parameter Storage Format (PSF). The global option parameter `senstype` (see the options statement) is used to control the type of sensitivity being calculated. In addition, you can use `+sensdata filename` command line argument or a global option (see the options statement) to direct sensitivity analyses results into a specified ASCII file.

Definition

```
sens (output_variables_list) to (design_parameters_list) for (analyses_list)
```

where

```
output_variables_list = ovar1 ovar2 ...
```

```
design_parameters_list = dpar1 dpar2 ...
```

```
analyses_list = anal1 anal2 ...
```

The list of the design parameters may include valid instance and model parameters. You can also specify device instances or device models without a modifier. In this case Spectre will attempt to compute sensitivities with respect to all corresponding instance or model parameters. Caution should be exercised in using this option as warnings or errors may be generated since many instance and model parameters cannot be modified. If no design parameters are specified then all the instance and model parameters are added. The list of

Spectre Circuit Simulator Reference

Spectre Syntax

the output variables for both AC and DC analyses may include node voltages and branch currents. For DC analyses, it also may include device instance operating point parameters.

Examples

```
sens (q1:betadc 2 Out) to (vcc:dc nbjt1:rb) for (analDC)
```

For this statement DC sensitivities of betadc operating point parameter of transistor `q1` and of nodes `2` and `Out` will be computed with respect to the `dc` voltage level of voltage source `vcc` and the model parameter `rb` for the DC analysis `analDC`. The results will be stored in the output file `analDC.sens.dc`.

```
sens (1 n2 7) to (q1:area nbjt1:rb) for (analAC)
```

For this statement AC sensitivities of nodes `1`, `n2`, `7` will be computed with respect to the area parameter of transistor `q1` and the model parameter `rb` for each frequency of the AC analysis `analAC`. The results will be stored in the output file `analAC.sens.ac`.

```
sens (1 n2 7) for (analAC)
```

For this statement AC sensitivities of nodes `1`, `n2`, `7` will be computed with respect to all instance and model parameters of all devices in the design for each frequency of the AC analysis `analAC`. The results will be stored in the output file `analAC.sens.ac`.

```
sens (vbb:p q1:int_c q1:gm 7) to (q1:area nbjt1:rb) for (analDC1)
```

For this statement DC sensitivities of the branch current `vbb:p`, the operating point parameter `gm` of the transistor `q1`, the internal collector voltage `q1:int_c` and the node `7` voltage will be computed with respect to the instance parameter `area` for instance `q1` and the model parameter `rb` for model `nbt1`.

SpectreHDL Usage and Language Summary (spectrehdl)

Description

SpectreHDL is a proprietary analog hardware description language. It allows analog circuit behavior to be described at a high level of abstraction, using a language which is similar to Verilog-A (run `spectre -h veriloga` for some details on the Verilog-A modeling language supported by Spectre). Behavioral descriptions of modules/components may be instantiated in a Spectre netlist along with regular Spectre primitives.

SpectreHDL descriptions are written in file(s) separate from the Spectre netlist file. These descriptions are written in modules (see the module `alpha` below). To include a module in the Spectre netlist, first add the line

```
ahdl_include "Ahdlfile.def"
```

Spectre Circuit Simulator Reference

Spectre Syntax

to the Spectre netlist file (where Ahdfile.def is the name of the file in which the required module is defined). The module is instantiated in the Spectre netlist in the same manner as Spectre primitives. For example,

```
name (node1 node2) alpha arg1=4.0 arg2=2 arg3="parameterized resistor"
```

This instantiates an element `alpha`, having two nodes and three parameters.

SpectreHDL modules can be debugged using `hdldebug`. `hdldebug` has a GUI and a command line mode. Please refer to the Verilog-A Debugging Tool User Guide for more information.

Module Template

The following is a SpectreHDL module template

```
module alpha( n1, n2 ) ( arg1, arg2, arg3 )
node [V,I] n1, n2;
parameter real arg1 = 2.0;
parameter integer arg2;
parameter string arg3;
{
    real local;
    // this is a comment
    initial {
        // initializations performed before the
        // start of an analysis.
    }
    analog {
        // module behavioral description
        V(n1, n2) <- I(n1, n2) * arg1;
    }
    final {
        // tasks performed at the end of an analysis
    }
}
```

Language Summary

The following provides a summary of the SpectreHDL analog hardware description language. For more information refer to the SpectreHDL Reference Manual.

Spectre Circuit Simulator Reference

Spectre Syntax

Derivative and Integral Operators

dot(x)	Differentiate x wrt time.
integ(x <, ic <, assert>>)	Integrate x wrt time. Output = ic during DC analysis. assert causes the integration to be reset.
idtmod(x, <ic <, modulus <, offset> > >)	Circular Integration of x wrt time. Output = ic during DC analysis. Integration is performed with given offset and modulus if specified.

Built-In Mathematical Functions

abs(x)	Absolute value
exp(x)	Exponential if $x < 80$
ceil(x)	Smallest integer $> x$
floor(x)	Largest integer $< x$
ln(x)	Natural logarithm
log(x)	Log base 10
sqrt(x)	Square root
min(x,y)	Minimum
max(x,y)	Maximum
pow(x,y)	x to the power of y

Simulator Time-Step Control Functions

\$threshold(x, direction <, abstol <, reltol_factor>>)	Set breakpoint when x crosses zero.
\$bound_step(max_step)	Limit timestep, (timestep \leq max_step).
\$break_point(target <, period>)	Set breakpoints at time = target and at times = $N \cdot \text{period} + \text{target}$ if period is specified.

Spectre Circuit Simulator Reference

Spectre Syntax

`$last_crossing(x, direction)`

Return time when expression last crossed zero in a given direction.

Waveform Filter Functions

`$transition(x <, delay <, trise <, tfall>>>)`

Specify details of signal transitions. For efficient simulation, it is recommended that x not be a continuous signal, i.e. a function of a probe. See the SpectreHDL manual for further explanation of this issue.

`$slew(x <, SRpos <, SRneg>>)`

Model slew rate behavior.

`$tdelay(x, time_delay, max_delay)`

Response(t) = x(t - time_delay).

`$zdelay(x <, period <, ttransition <, sample offset time <, ic>>>>)`

Fixed period sample and hold function.

`$zi_nd(x, numer, denom, period <, ttransition <,sample offset time>>)`

z-domain filter function, numerator-denominator form.

`$zi_zd(x, zeros, denom, period <, ttransition <,sample offset time>>)`

z-domain filter function, zero-denominator form.

`$zi_np(x, numer, poles, period <, ttransition <,sample offset time>>)`

z-domain filter function, numerator-pole form.

`$zi_zp(x, zeros, poles, period <, ttransition <,sample offset time>>)`

z-domain filter function, zero-pole form.

`$laplace_nd(x, numer, denom, <, abstol >)`

s-domain filter function, numerator-denominator form.

`$laplace_zd(x, zeros, denom, <, abstol >)`

s-domain filter function, zero-denominator form.

`$laplace_np(x, numer, poles, <, abstol >)`

s-domain filter function, numerator-pole form.

Spectre Circuit Simulator Reference

Spectre Syntax

`$laplace_zp(x, zeros, poles, <, abstol >)`
s-domain filter function, zero-pole form.

Noise Functions

`$white_noise(power <, tag >)`
Generates white noise with given power. Noise contributions with the same tag are combined for a module.

`$flicker_noise(power, exp <, tag >)`
Generates pink noise with given power at 1 Hz that varies in proportion to $1/f^{\text{exp}}$. Noise contributions with the same tag are combined for a module.

`$noise_table(vector <, tag >)`
Generates noise where power is determined by linear interpolation from the given vector of frequency-power pairs. Noise contributions with the same tag are combined for a module.

AC Analysis Stimuli

`$ac_stim(<analysis_name <, mag > >)`
Small signal source of specified magnitude, active for given analysis.

Interpolation Functions

`$build_table(type, response, inVec1, sizeVec1 <, inVec2, sizeVec2 ...>)`
Build a table for B-Spline interpolation.

`$interpolate(interp_table, v1<, v2 <, v3 <, v4 >>>)`
Perform interpolation at given point.

Simulator IO Functions

`$strobe("format string" <, arguments>)`
Print data to stdout every time step.

Spectre Circuit Simulator Reference

Spectre Syntax

<code>\$debug("format string" <, arguments>)</code>	Print data to stdout every iteration.
<code>\$fstrobe(fptr, "format string" <, arguments>)</code>	Print data to a file every time step.
<code>\$fdebug(fptr, "format string" <, arguments>)</code>	Print data to a file every iteration.
<code>\$fread(fptr, "format string" <, arguments>)</code>	Read data from a file.
<code>\$warning("format string" <, arguments>)</code>	Warning message.
<code>\$error("format string" <, arguments>)</code>	Error message. Abort analysis.
<code>\$fatal("format string" <, arguments>)</code>	Fatal message. Abort simulation.
<code>\$fopen("filename", mode)</code>	Open a file.
<code>\$fflush(fptr)</code>	Flush a file to disk.
<code>\$fclose(fptr)</code>	Close a file.
<code>\$popen("command", "mode")</code>	Open a pipe with given command in given mode.
<code>\$pclose(fptr)</code>	Close a pipe.
<code>\$read_table("filename", table_id <, comment_str<, delim_str> >)</code>	Read from a file into a 2-D real array.
<code>\$write_table("filename", table_id <,delim_str>)</code>	Write from a 2-D real array to a file.
<code>\$halt("string")</code>	Halt the simulation, printing given string.
<code>\$system("system command")</code>	Sends a command to the operating system.

Spectre Circuit Simulator Reference

Spectre Syntax

\$str("format_string" < ,arg1 < ,arg2 < ..etc> > >)	Create a string from arguments in given format.
\$strcmp(str1, str2)	Compares two strings lexicographically.
\$strtoint(int_as_str)	Converts a string, int_as_str, to an integer.
\$strtoreal(real_as_str)	Converts a string, real_as_str, to a real.
\$strcpy(des_str, src_str)	Copies src_str to des_src.
\$strcat(des_str, src_str)	Appends src_str to des_src.
\$strlen(str)	Returns the number of characters in str.
\$substr(input_str, start_pos, end_pos)	Returns the substring of input_str between start_pos and end_pos.
\$strstr(input_str, sub_str)	Returns the first position where sub_str is found in input_str.
\$strchr(input_str, character)	Returns the first position where character is found in input_str.
\$strrchr(input_str, character)	Returns the last position where character is found in input_str.
\$strspn(input_str, span_set)	Returns the number of continuous characters from the start of input_str that are in span_set.
\$strcspn(input_str, span_set)	Returns the number of continuous characters from the start of input_str that are not in span_set.
\$ascii(character)	Returns the ascii code of character.

Simulator Environment Functions

\$time()	Returns current simulation time.
\$temp()	Returns ambient simulation temperature.

Spectre Circuit Simulator Reference

Spectre Syntax

`$vt(<temp>)` Returns thermal voltage. If `temp` is defined, returns the thermalvoltage at `temp`.

`$analysis(analysis_string1<, analysis_string2 <, ...>>)`
Returns true(1) if the current analysis phase matches one of the given analyses strings. Valid analyses strings are; "dc", "tran", "ac", "pss", "noise", "pdisto", "qpss", "pac", "pnoise", "pxf", "sp", "tdr", "xf", "envlp", "psp", "qpssp", "qpac", "qpnoise", "qpxf", "static", or "ic".

Simulator Tolerance Functions

`$reltol()` Returns relative tolerance.

`$abstol(name)` Returns absolute tolerance of quantity `name`.

Parameter Functions

`$param_given(param)` Returns 1 if `param` was set. `param` can be a model parameter or an instance parameter.

`$pwr(x)` Assignment of model power consumption. Adds the expression `x` to the `pwr` parameter of a module.

Data Types

`integer` Discrete numerical type.

`real` Continuous numerical type.

`string` Text string type.

`stream` File pointer and text stream type.

`enum { name1 <, name2 <, name3 <, ... >>> }`
Discrete name type.

`void` Null or empty type.

`table` Interpolation table type.

Spectre Circuit Simulator Reference

Spectre Syntax

node [PotentialName, FlowName
Interconnection point type.

Data Qualifiers

parameter Indicates that a variable is a parameter and so may be given a different value when the module is instantiated and may have a range specifier.

const Indicates that a variable must be given a constant value when declared that can never be changed.

global Used only for internal nodes. Means that the internal node is an alias for a global node of the same name in the netlist.

Structural Statements

Structural statements are used inside the module block but outside the analog, final and initial blocks.

model module_or_primitive new_model (<param1 = expr1 <,...>>);
Used to create a model called new_model from module_or_primitive.

module_or_primitive inst_name (<node1 <, ..>>)(<param1 = expr1 <,...>>);
Creates a new instance of module_or_primitive called inst_name.

SpectreRF Summary (spectrerf)

Description

SpectreRF is a optional collection of analyses that are useful for circuits that are driven with a large periodic signal. Examples include mixers, oscillators, switched-capacitor filters, sample-and-holds, chopper stabilized amplifiers, frequency multipliers, frequency dividers, and samplers. They efficiently and directly compute the periodic and quasiperiodic steady-state solution of such circuits. They are capable of computing the large-signal and small-signal behavior, including noise behavior. Thus, SpectreRF is capable of computing the noise figure or intermodulation distortion of a mixer, the phase noise and harmonic distortion of an oscillator, and the frequency-response and noise behavior of a switched-capacitor filter. For

more information on the SpectreRF analyses, run `spectre -help analysisName` where `analysisName` is `pss`, `pac`, `pxf`, `pnoise`, `psp`, `qpss`, `qpac`, `qpxf`, `qpnoise`, `qpssp`, or `envlp`.

Subcircuit Definitions (subckt)

Description

Hierarchical circuit description

The `subckt` statement is used to define a subcircuit. Subcircuit definitions are simply circuit macros that can be expanded anywhere in the circuit any number of times. When an instance in your input file refers to a subcircuit definition, the instances specified within the subcircuit are inserted into the circuit. Subcircuits may be nested. Thus a subcircuit definition may contain instances of other subcircuits. Subcircuits may also contain component, analysis or model statements. Subcircuit definitions may also be nested, in which case the innermost subcircuit definition can only be referenced from within the subcircuit in which it is defined, and cannot be referenced from elsewhere.

Instances that instantiate a subcircuit definition are referred to as subcircuit calls. The node names (or numbers) specified in the subcircuit call are substituted, in order, for the node names given in the subcircuit definition. All instances that refer to a subcircuit definition must have the same number of nodes as are specified in the subcircuit definition and in the same order. Node names inside the subcircuit definition are strictly local unless declared otherwise in the input file with a global statement.

Subcircuit Parameters:

Parameter specification in subcircuit definitions is optional. In the case of nested subcircuit definitions, any parameters which have been declared for the outer subcircuit definition are also available within the inner subcircuit definition. Any parameters that are specified are referred to by name optionally followed by an equals sign and a default value. If, when making a subcircuit call, you do not specify a particular parameter, this default value is used in the macro expansion. Subcircuit parameters can be used in expressions within the subcircuit consisting of subcircuit parameters,

constants, and various mathematical operators. Run `spectre -h expressions` for more details on Spectre expression handling capability. Run `spectre -h parameters` for more details on how Spectre handles netlist parameters, including subcircuit parameters, and how they inherit within nested subcircuit definitions.

Spectre Circuit Simulator Reference

Spectre Syntax

Subcircuits always have an implicitly defined parameter m . This parameter is passed to all components in the subcircuit and each component is expected to multiply it by its own multiplicity factor. In this way, it is possible to efficiently model several copies of the subcircuit in parallel. It is an error to attempt to explicitly define m on a `parameters` line. Also, because m is only implicitly defined, it is not available for use in expressions in the subcircuit.

Inline Subcircuits

An inline subckt is a special case of a subckt where one of the devices or models instantiated within this subckt does not get its full hierarchical name, but rather inherits the subckt call name itself. An inline subckt is syntactically denoted by the presence of the keyword "inline" before the "subckt". It is called in the same manner as a regular subcircuit. The body of the inline subcircuit can typically contain one of the following, corresponding to different use models:

- multiple device instances (one of which is the "inline" component)
- multiple device instances, (one of which is "inline") and one or more parameterized models
- a single "inline" device instance and a parameterized model to which the device instance refers

The "inline" component is denoted by giving it the same name as the inline subcircuit itself. When the subcircuit is flattened, the "inline" component does not take on a hierarchical name such as X1.M1, but rather takes on the name of the subckt call itself, such as X1. Any non-inline components in the subckt take on the regular hierarchical name, just as if the subcircuit were a regular subckt (i.e. not an inline subckt).

Probing the inline device

Spectre allows the following list of items to be saved or probed for primitive devices. These would also apply to devices modeled as the inline components of inline subcircuits:

1. all terminal currents e.g. "save q1:currents"
2. specific (index) terminal current e.g. "save q1:1" (#1=collector)
3. specific (named) terminal current e.g. "save q1:b" ("b"=base)
4. save all operating point info e.g. "save q1:oppoint"
5. save specific operating point info e.g. "save q1:vbe"

6. save all currents and oppoint info e.g. "save q1"

Parameterized Models and Inline Subckts

Inline subckts can be used in the same way as regular subcircuits to implement parameterized models, however inline subckts provide some powerful new options. When an inline subcircuit contains both a parameterized model and an inline device which references that model, then the user can create instances of the device, and each device will automatically get an appropriately scaled model assigned to it. For example, the instance parameters to an inline subckt could represent something like emitter width and length of a BJT device and within the subckt a model card could be created which is parameterized for emitter width and length and scales accordingly. When the designer instantiates the macro, he/she supplies the values for the emitter width and length, and a device is instantiated with an appropriate geometrically scaled model. Again, the inline device does not get a hierarchical name, and can be probed in the same manner as the inline device in the previous section on modeling parasitics, that is, it can be probed just as if it was a simple device, and not actually embedded in a subckt

Automatic Model Selection using Inline Subckts:

See `spectre -h if` for a description on how to combine Spectres `structural if` statement with inline subckts to perform automatic model selection based on any netlist/subckt parameter.

Definition

```
[inline] subckt <Name
```

Example 1: subckt

```
subckt coax (i1 o1 i2 o2)
  parameters zin=50 zout=50 vin=1 vout=1 len=0
  inner i1 o1 i2 o2  tline z0=zin vel=vin len=len
  outer o1 0  o2 0  tline z0=zout vel=vout len=len
ends coax
```

defines a parameterized coaxial transmission line macro from two ideal transmission lines. To instantiate this subcircuit, one could use an instance statement such as:

```
Coax1 pin nin out gnd coax zin=75 zout=150 len=35m
```

Example 2: inline subckt - parasitics

Consider the following example of an inline subcircuit, which contains a mosfet instance, and two parasitic capacitances:

Spectre Circuit Simulator Reference

Spectre Syntax

```
inline subckt s1 (a b)                // "s1" is name of subckt
  parameters p1=1u p2=2u
  s1 (a b 0 0) mos_mod l=p1 w=p2      // "s1" is "inline" component
  cap1 (a 0) capacitor c=1n
  cap2 (b 0) capacitor c=1n
ends s1
```

The following circuit creates a simple mos device instance M1, and calls the inline subcircuit "s1" twice (M2 & M3)

```
M1 (2 1 0 0) mos_mod
M2 (5 6) s1 p1=6u p2=7u
M3 (6 7) s1
```

This expands/flattens to:

```
M1 (2 1 0 0) mos_mod
M2 (5 6 0 0) mos_mod l=6u w=7u // the "inline" component, inherits call name
M2.cap1 (5 0) capacitor c=1n // a regular hierarchical name
M2.cap2 (6 0) capacitor c=1n
M3 (6 7 0 0) mos_mod l=1u w=2u // the "inline" component, inherits call name
M3.cap1 (6 0) capacitor c=1n
M3.cap2 (7 0) capacitor c=1n
```

Here the final flattened names of the three mosfets (one for each instance) are M1, M2 and M3, rather than M1, M2.s1 and M3.s1 as they would be if s1 was a regular subcircuit. The parasitic capacitors (which the user is not really interested in, or perhaps even aware of, if the inline subckt definition was written by a separate modeling engineer) have full hierarchical names however.

Example 3: inline subckt - scaled models

Consider the following example, in which a parameterized model is declared within an inline subcircuit for a bipolar transistor. The model parameters are the emitter width, length, and area, and also the temperature delta (trise) of the device above nominal. Ninety nine instances of a 4*4 transistor are then placed, and one instance of a transistor with area=50 is placed. Each transistor gets an appropriately scaled model.

* declare a subckt, which instantiates a transistor with a parameterized model. The parameters are emitter width and length.

```
inline subckt bjtmod (c b e s)
  parameters le=1u we=2u area=le*we trise=0
  model mod1 bjt type=npn bf=100+(le+we)/2*(area/le-12)
  +      is=1e-12*(le/we)*(area/le-12)
  bjtmod (c b e s) mod1 trise=trise // "inline" component
ends bjtmod
```

Spectre Circuit Simulator Reference

Spectre Syntax

* some instances of this subckt

```
q1 (2 3 1 0) bjtmod le=4u we=4u          // trise defaults to zero
q2 (2 3 2 0) bjtmod le=4u we=4u trise=2
q3 (2 3 3 0) bjtmod le=4u we=4u
.
.
q99 (2 3 99 0) bjtmod le=4u we=4u
q100 (2 3 100 0) bjtmod le=1u area=50e-12
```

Verilog-A Usage and Language Summary (veriloga)

Description

Verilog-A is an analog hardware description language standard from Open Verilog International. It allows analog circuit behavior to be described at a high level of abstraction, using a language which is similar to SpectreHDL (run `spectre -h spectrehdl` for some details on the SpectreHDL modeling language). Behavioral descriptions of modules/components may be instantiated in a Spectre netlist along with regular Spectre primitives.

Verilog-A descriptions are written in file(s) separate from the Spectre netlist file. These descriptions are written in modules (see the module `alpha` below). To include a module in the Spectre netlist, first add the line

```
ahdl_include "VerilogAfile.va"
```

to the Spectre netlist file (where `VerilogAfile.va` is the name of the file in which the required module is defined). The module is instantiated in the Spectre netlist in the same manner as Spectre primitives for example,

```
name (node1 node2) alpha arg1=4.0 arg2=2
```

This instantiates an element `alpha`, having two nodes and two parameters.

Verilog-A modules can be debugged using `hdldebug`. `hdldebug` has a GUI and a command line mode. Please refer to the Verilog-A Debugging Tool User Guide for more information.

Module Template

The following is a Verilog-A module template

```
include "discipline.h"
```

Spectre Circuit Simulator Reference

Spectre Syntax

```
include "constants.h"
module alpha( n1, n2 );
electrical n1, n2;
parameter real arg1 = 2.0;
parameter integer arg2 = 0;
    real local;
    // this is a comment
    analog begin
        @ ( initial_step ) begin
            // performed at the first timestep of an analysis
        end
        // module behavioral description
        @ ( final_step ) begin
            // performed at the last time step of an analysis
        end
    end
endmodule
```

Language Summary

The following provides a summary of the Verilog-A analog hardware description language. For more information, refer to the *Verilog-A Reference Manual*.

Analog Operators/Waveform Filters

`ddt(x <,abstol>)` Differentiate x wrt time.

`idt(x, ic <, assert <, abstol> >)`
Integrate x wrt time. Output = ic during dc analysis and when assert is 1.

`idtmod(x, <ic <, modulus <, offset> > >)`
Circular Integration of x wrt time. Output = ic during DC analysis. Integration is performed with given offset and modulus if specified.

`transition(x <, delay <, trise <, tfall>>>)`
Specify details of signal transitions. For efficient simulation, it is recommended that x not be a continuous signal, i.e. a function of a probe. See the Verilog-A manual for further explanation of this issue.

Spectre Circuit Simulator Reference

Spectre Syntax

slew(x <, SRpos <, SRneg>>) Model slew rate behavior.

delay(x, time_delay, max_delay)
Response(t) = x(t - time_delay).

zi_nd(x, numer, denom, period, < ttransition <,sample offset time >)
z-domain filter function, numerator-denominator form.

zi_zd(x, zeros, denom, period, < ttransition <,sample offset time >)
z-domain filter function, zero-denominator form.

zi_np(x, numer, poles, period, < ttransition <,sample offset time >)
z-domain filter function, numerator-pole form.

zi_zp(x, zeros, poles, period, < ttransition <,sample offset time >)
z-domain filter function, zero-pole form.

laplace_nd(x, numer, denom, <, abstol >)
s-domain filter function, numerator-denominator form.

laplace_zd(x, zeros, denom, <, abstol >)
s-domain filter function, zero-denominator form.

laplace_np(x, numer, poles, <, abstol >)
s-domain filter function, numerator-pole form.

laplace_zp(x, zeros, poles, <, abstol >)
s-domain filter function, zero-pole form.

Built-In Mathematical Functions

abs(x)	Absolute value
exp(x)	Exponential if $x < 80$
ln(x)	Natural logarithm
log(x)	Log base 10
sqrt(x)	Square root
min(x,y)	Minimum

Spectre Circuit Simulator Reference

Spectre Syntax

<code>max(x,y)</code>	Maximum
<code>pow(x,y)</code>	x to the power of y

Noise Functions

`white_noise(power <, tag >)`
Generates white noise with given power. Noise contributions with the same tag are combined for a module.

`flicker_noise(power, exp <, tag >)`
Generates pink noise with given power at 1 Hz that varies in proportion to $1/f^{\text{exp}}$. Noise contributions with the same tag are combined for a module.

`noise_table(vector <, tag >)`
Generates noise where power is determined by linear interpolation from the given vector of frequency-power pairs. Noise contributions with the same tag are combined for a module.

AC Analysis Stimuli

`ac_stim(<analysis_name <, mag > >)`
Small signal source of specified magnitude, active for given analysis.

Analog Events

Analog events must be contained in an analog event detection statement; `@(analog_event)` statement.

`cross(x, direction <, timetol <, abstol >>)`
Generates an event when x crosses zero.

`above(x, <, timetol <, abstol >>)`
Generates an event when x becomes greater than or equal to zero. An above event can be generated and detected during initialization. By contrast, a cross event can be generated and detected only after at least one transient time step is complete.

Spectre Circuit Simulator Reference

Spectre Syntax

timer(start_time <, period>)

Set (optionally periodic) breakpoint event at time = start_time.

initial_step< (arg1 <, arg2 <, etc... > >)

Generate an event at the initial step of an analysis. arg1, arg2, etc. may be any of: "dc", "tran", "ac", "pss", "noise", "pdisto", "qpss", "pac", "pnoise", "pxf", "sp", "tdr", "xf", "envlp", "psp", "qpssp", "qpac", "qpnoise", "qpxf", "static", or "ic".

final_step< (arg1 <, arg2 <, etc... > >)

Generate an event at the final step of an analysis. arg1, arg2, etc. may be any of: "dc", "tran", "ac", "pss", "noise", "pdisto", "qpss", "pac", "pnoise", "pxf", "sp", "tdr", "xf", "envlp", "psp", "qpssp", "qpac", "qpnoise", "qpxf", "static", or "ic".

Timestep Control

bound_step(max_step)

Limit timestep, (timestep <= max_step).

last_crossing(x, direction)

Return time when expression last crossed zero in a given direction.

discontinuity(n)

Hint to simulator that discontinuity is present in nth derivative.

Simulator IO Functions

\$display(argument_list)

Print data to stdout. Formatting strings may be interspersed between arguments/data.

\$fdisplay(fptr, argument_list)

Print data to a file. Formatting strings may be interspersed between arguments/data.

\$strobe(argument_list)

Print data to stdout. Formatting strings may be interspersed between arguments/data.

\$fstrobe(fptr, argument_list)

Print data to a file. Formatting strings may be interspersed between arguments/data.

\$fopen("filename")

Open a file for writing.

Spectre Circuit Simulator Reference

Spectre Syntax

\$fclose(fpnr)	Close a file.
\$finish<(n)>	Finish the simulation.
\$stop<(n)>	Stop the simulation.

Simulator Environment Functions

\$realtime	Returns current simulation time.
\$temperature	Returns ambient simulation temperature (K).
\$vt	Returns thermal voltage.
\$vt(temp)	Returns thermal voltage at given temp.
\$analysis(analysis_string1<, analysis_string2 <, ...>>)	Returns true(1) if the current analysis phase matches one of the given analyses strings. Valid analyses strings are; "dc", "tran", "ac", "pss", "noise", "pdisto", "qpss", "pac", "pnoise", "pxf", "sp", "tdr", "xf", "envlp", "psp", "qpssp", "qpac", "qpnoise", "qpxf", "static", or "ic".

Parameter Functions

\$pwr(x)	Assignment of model power consumption. Adds the expression x to the pwr parameter of a module.
------------	--

Data Types

integer	Discrete numerical type.
real	Continuous numerical type.

Data Qualifiers

parameter	Indicates that a variable is a parameter and so may be given a different value when the module is instantiated, and that it may not be assigned a different value inside the module.
-----------	--

Spectre Circuit Simulator Reference

Spectre Syntax

Structural Statements

Structural statements are used inside the module block but outside the analog block.

```
module_or_primitive #(<.param1(expr1)<,...>>) inst_name (<node1 <, ..>> );
```

Creates a new instance of module_or_primitive called inst_name.

Spectre Circuit Simulator Reference
Spectre Syntax

References

This section gives additional details about the source documents referred to in the text.

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