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## **Xyce™ Parallel Electronic Simulator Reference Guide, Version 6.2**

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### **Abstract**

This document is a reference guide to the **Xyce** Parallel Electronic Simulator, and is a companion document to the **Xyce** Users' Guide [1] . The focus of this document is (to the extent possible) exhaustively list device parameters, solver options, parser options, and other usage details of **Xyce**. This document is *not* intended to be a tutorial. Users who are new to circuit simulation are better served by the **Xyce** Users' Guide [1] .

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# 1. Introduction

## Welcome to **Xyce**

The **Xyce** Parallel Electronic Simulator has been written to support, in a rigorous manner, the simulation needs of the Sandia National Laboratories electrical designers. It is targeted specifically to run on large-scale parallel computing platforms but also runs well on a variety of architectures including single processor workstations. It also aims to support a variety of devices and models specific to Sandia needs.

## 1.1 Overview

This document is intended to complement the **Xyce** Users' Guide [1] . It contains comprehensive, detailed information about a number of topics pertinent to the usage of **Xyce**. Included in this document is a netlist reference for the input-file commands and elements supported within **Xyce**; a command line reference, which describes the available command line arguments for **Xyce**; and quick-references for users of other circuit codes, such as Orcad's PSpice [2].

## 1.2 How to Use this Guide

This guide is designed so you can quickly find the information you need to use **Xyce**. It assumes that you are familiar with basic Unix-type commands, how Unix manages applications and files to perform routine tasks (e.g., starting applications, opening files and saving your work). Note that while Windows versions of **Xyce** are available, they are command-line programs meant to be run under the *Command Prompt*, and are used almost identically to their Unix counterparts.

### Typographical conventions

Before continuing in this Reference Guide, it is important to understand the terms and typographical conventions used. Procedures for performing an operation are generally indicated with the following typographical conventions.

Notation	Example	Description
Typewriter text	xmpirun -np 4	Commands entered from the keyboard on the command line or text entered in a netlist.
<b>Bold Roman Font</b>	Set nominal temperature using the <b>TNOM</b> option.	SPICE-type parameters used in models, etc.
Gray Shaded Text	DEBUGLEVEL	Feature that is designed primarily for use by <b>Xyce</b> developers.
[text in brackets]	Xyce [options] <netlist>	Optional parameters.
<text in angle brackets>	Xyce [options] <netlist>	Parameters to be inserted by the user.
<object with asterisk>*	K1 <ind. 1> [<ind. n>*]	Parameter that may be multiply specified.
<TEXT1 TEXT2>	.PRINT TRAN + DELIMITER=<TAB COMMA>	Parameters that may only take specified values.

Table 1.1: **Xyce** typographical conventions.

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## 2. Netlist Reference

### Chapter Overview

This chapter contains reference material directed towards working with circuit analyses in **Xyce** using the netlist interface. Included are detailed command descriptions, start-up option definitions and a list of devices supported by the **Xyce** netlist interface.

## 2.1 Netlist Commands

This section outlines the netlist commands that can be used with **Xyce** to setup and control circuit analysis.

### 2.1.1 .AC (AC Analysis)

Calculates the frequency response of a circuit over a range of frequencies.

The .AC command can specify a linear sweep, decade logarithmic sweep, or octave logarithmic sweep.

**General Form**     .AC <sweep type> <points value>  
                         + <start frequency value> <end frequency value>

---

**Examples**            .AC LIN 101 100Hz 200Hz  
                         .AC OCT 10 1kHz 16kHz  
                         .AC DEC 20 1MEG 100MEG

---

#### Arguments and Options

sweep type

Must be LIN, OCT, or DEC, as described below.

LIN Linear sweep

The sweep variable is swept linearly from the starting to the ending value.

OCT Sweep by octaves

The sweep variable is swept logarithmically by octaves.

DEC Sweep by decades

The sweep variable is swept logarithmically by decades.

points value

Specifies the number of points in the sweep, using an integer.

start frequency value

end frequency value

The end frequency value must not be less than the start frequency value, and both must be greater than zero. The whole sweep must include at least one point.

---

**Comments**            AC analysis is a linear analysis. The simulator calculates the frequency response by linearizing the circuit around the bias point.

A .PRINT AC must be used to get the results of the AC sweep analysis. See Section 2.1.21.

## 2.1.2 .DC (DC Sweep Analysis)

Calculates the operating point for the circuit for a range of values. Primarily, this capability is applied to independent voltage sources, but it can also be applied to most device parameters. Note that this may be repeated for multiple sources in the same .DC line.

The .DC command can specify a linear sweep, decade logarithmic sweep, octave logarithmic sweep, or a list of values.

### Linear Sweeps

<b>General Form</b>	<code>.DC [LIN] &lt;sweep variable name&gt; &lt;start&gt; &lt;stop&gt; &lt;step&gt; + [&lt;sweep variable name&gt; &lt;start&gt; &lt;stop&gt; &lt;step&gt;]*</code>
---------------------	---

<b>Examples</b>	<code>.DC LIN V1 5 25 5 .DC VIN -10 15 1 .DC R1 0 3.5 0.05 C1 0 3.5 0.5</code>
-----------------	--

<b>Comments</b>	A .PRINT DC must be used to get the results of the DC sweep analysis. See Section 2.1.21. A .OP comand will result in a linear DC analysis if there is no .DC specified.
-----------------	---

### Decade Sweeps

<b>General Form</b>	<code>.DC DEC &lt;sweep variable name&gt; &lt;start&gt; &lt;stop&gt; &lt;points&gt; + [DEC &lt;sweep variable name&gt; &lt;start&gt; &lt;stop&gt; &lt;points&gt;]*</code>
---------------------	---

<b>Examples</b>	<code>.DC DEC VIN 1 100 2 .DC DEC R1 100 10000 3 DEC VGS 0.001 1.0 2</code>
-----------------	---

### Octave Sweeps

<b>General Form</b>	<code>.DC OCT &lt;sweep variable name&gt; &lt;start&gt; &lt;stop&gt; &lt;points&gt; + [OCT &lt;sweep variable name&gt;&lt;start&gt; &lt;stop&gt; &lt;points&gt;]...</code>
---------------------	--

<b>Examples</b>	<code>.DC OCT VIN 0.125 64 2 .DC OCT R1 0.015625 512 3 OCT C1 512 4096 1</code>
-----------------	---

### List Sweeps

<b>General Form</b>	<code>.DC &lt;sweep variable name&gt; LIST &lt;val&gt; &lt;val&gt; &lt;val&gt;* + [ &lt;sweep variable name&gt; LIST &lt;val&gt; &lt;val&gt;* ]*</code>
---------------------	---

---

**Examples**

```
.DC VIN LIST 1.0 2.0 5.0 6.0 10.0  
.DC VDS LIST 0 3.5 0.05 VGS LIST 0 3.5 0.5  
.DC TEMP LIST 10.0 15.0 18.0 27.0 33.0
```

### 2.1.3 .DCVOLT (Initial Condition, Bias point)

The .DCVOLT sets initial conditions for an operating point calculation. It is identical in function to the .IC command. See section 2.1.10 for detailed guidance.

## 2.1.4 .END (End of Circuit)

Marks the end of netlist file.

## 2.1.5 .ENDS (End of Subcircuit)

Marks the end of a subcircuit definition.

## 2.1.6 .FOUR (Fourier Analysis)

Performs Fourier analysis of transient analysis output.

**General Form**     `.FOUR <freq> <ov> [ov]*`

---

**Examples**            `.FOUR 100K v(5)`  
                      `.FOUR 1MEG v(5,3) v(3)`

---

### Arguments and Options

`freq`

The fundamental frequency used for Fourier analysis. Fourier analysis is performed over the last period ( $1/\text{freq}$ ) of the transient simulation. The DC component and the first nine harmonics are calculated.

`ov`    The desired solution variable, or variables, to be analyzed. Fourier analysis can be performed on several solution variables for each fundamental frequency, `freq`. At least one solution variable must be specified in the `.FOUR` line. The available solution variables are:

- `V(<circuit node>)` the voltage at `<circuit node>`
- `V(<circuit node>,<circuit node>)` to output the voltage difference between the first `<circuit node>` and second `<circuit node>`
- `I(<device>)` the current through a two terminal device
- `I<lead abbreviation>(<device>)` the current into a particular lead of a three or more terminal device (see the Comments, below, for details)
- `N(<device parameter>)` a specific device parameter (see the individual devices in Section 2.3 for syntax)

### Comments

Multiple `.FOUR` lines may be used in a netlist. All results from Fourier analysis will be returned to the user in a file with the same name as the netlist file suffixed with a `.FOUR`.

`<lead abbreviation>` is a single character designator for individual leads on a device with three or more leads. For bipolar transistors these are: c (collector), b (base), e (emitter), and s (substrate). For mosfets, lead abbreviations are: d (drain), g (gate), s (source), and b (bulk). SOI transistors have: d, g, s, e (bulk), and b (body). For PDE devices, the nodes are numbered according to the order they appear, so lead currents are referenced like `I1(<device>)`, `I2(<device>)`, etc.

## 2.1.7 .FUNC (Function)

User defined functions that can be used in expressions appearing later in the same scope as the .FUNC statement.

**General Form**     .FUNC <name>([arg]\*) {<body>}

---

**Examples**

```
.FUNC E(x) {exp(x)}  
.FUNC DECAY(CNST) {E(-CNST*TIME)}  
.FUNC TRIWAV(x) {ACOS(COS(x))/3.14159}  
.FUNC MIN3(A,B,C) {MIN(A,MIN(B,C))}
```

---

### Arguments and Options

name

Function name. Functions cannot be redefined and the function name must not be the same as any of the predefined functions (e.g., SIN and SQRT).

arg The arguments to the function. .FUNC arguments cannot be node names. The number of arguments in the use of a function must agree with the number in the definition. Parameters, TIME, and other functions are allowed in the body of function definitions. Two constants EXP and PI cannot be used as argument names. These constants are equal to  $e$  and  $\pi$ , respectively, and cannot be redefined.

body

May refer to other (previously defined) functions; the second example, DECAY, uses the first example, E.

---

**Comments**     The <body> of a defined function is handled in the same way as any math expression; it must be enclosed in curly braces .

## 2.1.8 .GLOBAL\_PARAM (Global parameter)

User-defined global parameter, which can be time dependent, or can be used in .STEP loops.

**General Form**     .GLOBAL\_PARAM [<name>=<value>]\*

---

**Examples**            .GLOBAL\_PARAM T={27+100\*time}

name

    Name of the global parameter.

value

    Global parameter value. An expression is used for the value when specified within curly braces ({}).

---

**Comments**

You may use parameters defined by .PARAM in expressions used to define global parameters, but you may *not* use global parameters in .PARAM definitions.

Unlike .PARAM parameters, global parameters are evaluated at the time they are needed. They may, therefore, be time dependent, and may depend on other time dependent quantities in the circuit.

Global parameters are accessible, and have the same value, throughout all levels of the netlist hierarchy. It is not legal to redefine global parameters in different levels of the netlist hierarchy.

## 2.1.9 .HB (Harmonic Balance Analysis)

Calculates steady states of nonlinear circuits in the frequency domain.

**General Form**     `.HB <fundamental frequency>`

---

**Examples**         `.HB 1e4`

---

**Arguments and Options**     `fundamental frequency`  
Sets the fundamental frequency for the analysis.

---

**Comments**            Harmonic balance analysis calculates the magnitude and phase of voltages and currents in a nonlinear circuit. Use a `.OPTIONS HBINT` statement to set additional harmonic balance analysis options.

The `.PRINT HB` statement must be used to get the results of the harmonic balance analysis. See section 2.1.21.

## 2.1.10 .IC (Initial Condition, Bias point)

The `.IC/.DCVOLT` command sets initial conditions for operating point calculations. These operating point conditions will be enforced the entire way through the nonlinear solve. Initial conditions can be given for some or all of the circuit nodes.

As the conditions are enforced for the entire solve, only the nodes not specified with `.IC` statements will change over the course of the operating point calculation.

Note that it is possible to specify conditions that are not solvable. Consult the **Xyce** User's Guide for more guidance.

<b>General Form</b>	<code>.IC V(&lt;node&gt;)=&lt;value&gt;</code> <code>.IC &lt;node&gt; &lt;value&gt;</code> <code>.DCVOLT V(&lt;node&gt;)=&lt;value&gt;</code> <code>.DCVOLT &lt;node&gt; &lt;value&gt;</code>
---------------------	--

---

<b>Examples</b>	<code>.IC V(2)=3.1</code> <code>.IC 2 3.1</code> <code>.DCVOLT V(2)=3.1</code> <code>.DCVOLT 2 3.1</code>
-----------------	--

---

<b>Comments</b>	<p>The <code>.IC</code> capability can only set voltage values, not current values.</p> <p><code>.IC</code> lines do not support the use of expressions or parameters. So, only numbers can be used as initial values. However, if a device has an <code>IC</code> instance parameter then expressions and parameters can be used to set that device instance parameter.</p> <p>There are two ways to set initial conditions for the initial voltage drop across a node in a subcircuit. If the devices have <code>IC</code> instance parameters then that may be the best approach. If <code>.IC</code> lines is used instead then the fully-resolved node names must be used. Those names can be found via the <code>-namesfile</code> command line option. The syntax is <code>Xyce -namesfile [netlist]</code>.</p>
-----------------	---

## 2.1.11 .INC or .INCLUDE (Include file)

Include specified file in netlist.

The file name can be surrounded by double quotes, "filename", but this is not necessary. The directory for the include file is assumed to be the execution directory unless a full or relative path is given as a part of the file name.

<b>General Form</b>	<code>.INC &lt;include file name&gt;</code> <code>.INCLUDE &lt;include file name&gt;</code>
---------------------	--

---

<b>Examples</b>	<code>.INC models.lib</code> <code>.INC "models.lib"</code> <code>.INCLUDE models.lib</code> <code>.INCLUDE "path_to_library/models.lib"</code>
-----------------	--

## 2.1.12 .LIB (Library file)

The .LIB command is similar to .INCLUDE, in that it brings in an external file. However, it is designed to only bring in specific parts of a library file, as designated by an entry name. Note that the **Xyce** version of .LIB has been designed to be compatible with HSPICE [3], not PSpice [4].

There are two forms of the .LIB statement, the call and the definition. The call statement reads in a specified subset of a library file, and the definition statement defines the subsets.

### .LIB call statement

**General Form**     .LIB <file name> <entry name>

---

**Examples**

```
.LIB models.lib nom
.LIB "models.lib" low
.LIB "path/models.lib" high
```

---

#### Arguments and Options

**file name**  
Name of file containing netlist data. Double quotes (") may be used around the file name.

**entry name**  
Entry name, which determines the section of the file to be included. These sections are defined in the included file using the definition form of the .LIB statement.

The library file name can be surrounded by double quotes, as in "path/filename" but this is not necessary. The directory for the library file is assumed to be the execution directory unless a full or relative path is given as a part of the file name. The section name denotes the section or sections of the library file to include.

### .LIB definition statement

The format given above is when the .LIB command is used to reference a library file; however, it is also used as part of the syntax in a library file.

**General Form**

```
.LIB <entry name>
<netlist lines>*
.endl <entry name>
```

---

#### Examples

```
* Library file res.lib
.lib low
```

```
.param rval=2
r3 2 0 9
.endl low

.lib nom
.param rval=3
r3 2 0 8
.endl nom
```

---

## Arguments and Options

entry name

The name to be used to identify this library component. When used on a .LIB call line, these segments of the library file will be included in the calling file.

Note that for each entry name, there is a matched .lib and .endl. Any valid netlist commands can be placed inside the .lib and .endl statements. The following is an example calling netlist, which refers to the library in the examples above:

```
* Netlist file res.cir
V1 1 0 1
R 1 2 {rval}
.lib res.lib nom
.tran 1 ps 1ns
.end
```

In this example, only the netlist commands that are inside of the “nom” library will be parsed, while the commands inside of the “low” library will be discarded. As a result, the value for resistor r3 is 8, and the value for rval is 3.

## 2.1.13 .MEASURE (Measure output)

The .MEASURE statement allows calculation or reporting of simulation metrics to an external file. One can measure when simulated signals reach designated values, or when they are equal to other simulation values. The syntax for a .MEASURE statement is as follows:

**General Form**

```
.MEASURE TRAN <result name> AVG <variable>
+ [FROM=<value>] [TO=<value>] [MIN_THRESH=<value>] [MAX_THRESH=<value>]
+ [FROM=<value>] [TO=<value>]

.MEASURE TRAN <result name> DERIV <variable>
+ [FROM=<value>] [TO=<value>]

.MEASURE TRAN <result name> DUTY <variable>
+ [FROM=<value>] [TO=<value>] [ON=<value>] [OFF=<value>] [MINVAL=<value>]

.MEASURE TRAN <result name> EQN <expression>
+ [FROM=<value>] [TO=<value>]

.MEASURE TRAN <result name> FOUR <variable> AT=freq
+ [NUMFREQ=<value>] [GRIDSIZE=<value>]

.MEASURE TRAN <result name> FREQ <variable>
+ [FROM=<value>] [TO=<value>] [TD=<value>]
+ [ON=<value>] [OFF=<value>] [MINVAL=<value>]

.MEASURE TRAN <result name> INTEG <variable>
+ [FROM=<value>] [TO=<value>]

.MEASURE TRAN <result name> MAX <variable>
+ [TD=<value>]

.MEASURE TRAN <result name> MIN <variable>
+ [TD=<value>]

.MEASURE TRAN <result name> OFF_TIME <variable>
+ [FROM=<value>] [TO=<value>] [OFF=<value>] [MINVAL=<value>]

.MEASURE TRAN <result name> ON_TIME <variable>
+ [FROM=<value>] [TO=<value>] [ON=<value>] [MINVAL=<value>]

.MEASURE TRAN <result name> PP <variable>
+ [TD=<value>]

.MEASURE TRAN <result name> RMS <variable>
+ [FROM=<value>] [TO=<value>]
```

```
.MEASURE TRAN <result name> WHEN <variable>=<variable2>|<value>
+ [TD=<value>] [RISE=r|LAST] [FALL=f|LAST] [CROSS=c|LAST]
+ [MINVAL=<value>] [DEFAULT_VAL=<value>]

.MEASURE TRAN <result name> TRIG <variable>=<variable2>|<value>
+ TARG <variable3>=<variable4>|<value> [TD=<value>]
```

---

## Examples

```
.MEASURE TRAN hit1_75 WHEN V(1)=0.75 MINVAL=0.02
.MEASURE TRAN hit2_75 WHEN V(1)=0.75 MINVAL=0.08 RISE=2
.MEASURE TRAN avgAll AVG V(1)
.MEASURE TRAN dutyAll DUTY V(1) ON=0.75 OFF=0.25
```

---

## Arguments and Options

**result name**

Measured results are reported to the output and log file. Additionally results are stored in a file called `circuitFileName.mt#`, where the suffixed number starts at 0 and increases for multiple iterations of a given simulation. Each line of this file will contain the measurement name, `<result name>`, followed by its value for that run.

AVG, DERIV, DUTY, EQN, FREQ, FOUR, INTEG, MAX, MIN, OFF\_TIME, ON\_TIME, PP, RMS, WHEN, TRIG, TARG

The third argument specifies the type of measurement or calculation to be done. By default, the measurement is performed over the entire simulation. The calculations can be limited to a specific window by using the qualifiers FROM, TO, TD, RISE, FALL, CROSS and MINVAL, which are explained below. The supported types are:

**AVG** Computes the arithmetic mean of `<variable>` for the simulation, or within the extent of the measurement window. The qualifiers FROM and TO can be used to limit the time window.

**DERIV** Computes the derivative of `<variable>`, estimated either by the slope between the first and last value found within the simulation, or within the extent of the measurement window. The qualifiers FROM and TO can be used to limit the time window.

**DUTY** Fraction of time that `<variable>` is greater than ON and does not fall below OFF either for the simulation, or the measurement window specified using the qualifiers FROM TO. The qualifier MINVAL is used as a tolerance on the ON and OFF values as in  $ON \pm MINVAL$  and  $OFF \pm MINVAL$ .

**EQN** Calculates the value of `<expression>` during the simulation. Also supports the qualifiers FROM and TO to limit the time window.

**FOUR** Calculates the fourier transform of the transient waveform for `<variable>`, given the fundamental frequency AT. The DC component and the first NUMFREQ-1 harmonics are determined using an interpolation of GRIDSIZE.

**FREQ** An estimate of the frequency of <variable>, found by cycle counting during the simulation. Cycles are defined through the values of ON and OFF with  $\pm$  MINVAL being used as a tolerance. The time window for counting cycles is either the entire simulation, or is delimited by the qualifier TD for a time delay. Additionally, The qualifiers FROM and TO can be used to limit the time window.

**INTEG** Calculates the integral of outVal through second order numerical integration. The integration window can be limited with the qualifiers FROM and TO.

**MAX** Returns the maximum value of <variable> during the simulation, or limited by the time qualifier TD for a time delay.

**MIN** Returns the minimum value of <variable> during the simulation, or limited by the time qualifier TD for a time delay.

**OFF\_TIME** Returns the time that <variable> is below OFF for the simulation. OFF uses  $\pm$  MINVAL as a tolerance and the measurement window can be limited with the qualifiers FROM and TO.

**ON\_TIME** Returns the time that <variable> is above ON. ON uses  $\pm$  MINVAL as a tolerance and the measurement window can be limited with the qualifiers FROM and TO.

**PP** Returns the difference between the maximum value and the minimum value <variable> during the simulation, or limited by the time qualifier TD for a time delay

**RMS** Computes the root-mean-squared value of <variable> during the simulation, or limited by the time qualifiers FROM and TO.

**TRIG**

**TARG** Measures the time between a trigger event and a target event. The trigger is specified with TRIG <variable>=<variable<sub>2</sub>> or TRIG <variable>=<value>. Likewise, the the target is specified as TARG <variable<sub>3</sub>>=<variable<sub>4</sub>> or TRIG <variable<sub>3</sub>>=<value>. It is also possible to use this measure to find a rise time for variable when the rise time is defined as the time to go from some small fraction of the maxima to some other fraction of the maxima. For example a rise time from 10% to 90% of the maxima. For that case the syntax is TRIG v(node) frac\_max=0.1 TARG v(node) frac\_max=0.9

**WHEN** Returns the time when <variable> reaches <variable<sub>2</sub>> or the constant value, value. The time over which the value is searched can be limited by the qualifiers TD, RISE, FALL and CROSS. The qualifier MINVAL acts as a tolerance. For example when <variable<sub>2</sub>> is specified, the comparison used is when <variable> = <variable<sub>2</sub>>  $\pm$  MINVAL or when a constant, value is given: <variable> = value  $\pm$  MINVAL. If the conditions specified for finding a given value were not found during the simulation then the measure will return the default value of -1. This default return value is settable by the user, for each measure, by adding the option DEFAULT\_VAL=retval on that measure line.

variable  
variable<sub>n</sub>  
value

This represents the test for the stated measurement. <variable> is a simulation quantity, such as a voltage or current. One can compare it to another simulation variable or a fixed quantity. Additionally, the <variable> may be an expression delimited by {, } brackets.

FROM=value

A time *after which* the measurement calculation will start.

T0=value

A time *at which* the measurement calculation will stop.

MIN\_THRESH=value

A minimum, threshold value above which the measurement calculation will be done and below which it will not be done.

MAX\_THRESH=value

A maximum, threshold value above which the measurement calculation will not be done and below which it will be done.

TD=value

A time delay before which this measurement should be taken or checked.

RISE=r | LAST

The number of rises after which the measurement should be checked. If LAST is specified, then the last rise found in the simulation will be used.

FALL=f | LAST

The number of falls after which the measurement should be checked. If LAST is specified, then the last fall found in the simulation will be used.

CROSS=c | LAST

The number of zero crossings after which the measurement should be checked. If LAST is specified, then the last zero crossing found in the simulation will be used.

MINVAL=value

An allowed absolute difference between outVal and the variable to which it is being compared. This has a default value of 1.0e-12. One may need to specify a larger value to avoid missing the test condition in a transient run. AVG DERIV DUTY FREQ INTEG time threshold

ON=value

The value at which a signal is considered to be on for frequency, duty and on time calculations

OFF=value

The value at which a signal is considered to be off for frequency, duty and off time calculations

GOAL=value

This parameter is not implemented in **Xyce**, but is included for compatibility with HSPICE netlists.

WEIGHT=value

This parameter is not implemented in **Xyce**, but is included for compatibility with HSPICE netlists.

## 2.1.14 .MODEL (Model Definition)

The .MODEL command provides a set of device parameters to be referenced by device instances in the circuit.

**General Form**     .MODEL <model name> <model type> (<name>=<value>)\*

---

**Examples**

```
.MODEL RMOD R (RSH=1)
.MODEL MOD1 NPN BF=50 VAF=50 IS=1.E-12 RB=100 CJC=.5PF TF=.6NS
.MODEL NFET NMOS(LEVEL=1 KP=0.5M VTO=2V)
```

---

### Arguments and Options

model name

The model name used to reference the model.

model type

The model type used to define the model. This determines if the model is (for example) a resistor, or a MOSFET, or a diode, etc. For transistors, there will usually be more than one type possible, such as NPN and PNP for BJTs, and NMOS and PMOS for MOSFETs.

name

value

The name of a parameter and its value. Most models will have a list of parameters available for specification. Those which are not set will receive default values. Most will be floating point numbers, but some can be integers and some can be strings, depending on the definition of the model.

## LEVEL Parameter

A common parameter is the **LEVEL** parameter, which is set to an integer value. This parameter will define exactly which model of the given type is to be used. For example, there are many different available MOSFET models. All of them will be specified using the same possible names and types. The way to differentiate (for example) between the BSIM3 model and the PSP model is by setting the appropriate **LEVEL**.

## Model Interpolation

Traditionally, SPICE simulators handle thermal effects by coding temperature dependence of model parameters into each device. These expressions modify the nominal device parameters given in the .MODEL card when the ambient temperature is not equal to **TNOM**, the temperature at which parameters were extracted.

These temperature correction equations may be reasonable at temperatures close to **TNOM**, but Sandia users of **Xyce** have found them inadequate when simulations must be performed over

a wide range of temperatures. To address this inadequacy, **Xyce** implements a model interpolation option that allows the user to specify multiple `.MODEL` cards, each extracted from real device measurements at a different `TNOM`. From these model cards, **Xyce** will interpolate parameters based on the ambient temperature using either piecewise linear or quadratic interpolation.

Interpolation of models is accessed through the model parameter `TEMPMODEL` in the models that support this capability. In the netlist, a base model is specified, and is followed by multiple models at other temperatures.

Interpolation of model cards in this fashion is implemented in the BJT level 1, JFET, MESFET, and MOSFETS levels 1-6, 10, and 18.

The use of model interpolation is best shown by example:

```
Jtest 1a 2a 3 SA2108 TEMP= 40
*
.MODEL SA2108 PJF ( TEMPMODEL=QUADRATIC TNOM = 27
+ LEVEL=2 BETA= 0.003130 VTO = -1.9966 PB = 1.046
+ LAMBDA = 0.00401 DELTA = 0.578; THETA = 0;
+ IS = 1.393E-10          RS = 1e-3)
*
.MODEL SA2108 PJF ( TEMPMODEL=QUADRATIC TNOM = -55
+ LEVEL=2 BETA = 0.00365 VTO = -1.9360 PB = 0.304
+ LAMBDA = 0.00286 DELTA = 0.2540 THETA = 0.0
+ IS = 1.393E-10 RD = 0.0 RS = 1e-3)
*
.MODEL SA2108 PJF ( TEMPMODEL=QUADRATIC TNOM = 90
+ LEVEL=2 BETA = 0.002770 VTO = -2.0350 PB = 1.507
+ LAMBDA = 0.00528 DELTA = 0.630 THETA = 0.0
+ IS = 1.393E-10          RS = 5.66)
```

Note that the model names are all identical for the three `.MODEL` lines, and that they all specify `TEMPMODEL=QUADRATIC`, but with different `TNOM`. For parameters that appear in all three `.MODEL` lines, the value of the parameter will be interpolated using the `TEMP=` value in the device line, which in this example is 40°C, in the first line. For parameters that are not interpolated, such as `RD`, it is not necessary to include these in the second and third `.MODEL` lines.

The only valid arguments for `TEMPMODEL` are **QUADRATIC** and **PWL** (piecewise linear). The quadratic method includes a limiting feature that prevents the parameter value from exceeding the range of values specified in the `.MODEL` lines. For example, the `RS` value in the example would take on negative values for most of the interval between -55 and 27, as the value at 90 is very high. This truncation is necessary as parameters can easily take on values (such as the negative resistance of `RS` in this example) that will cause a **Xyce** failure.

With the BJT parameters `IS` and `ISE`, interpolation is done not on the parameter itself, but on the the log of the parameter, which provides for excellent interpolation of these parameters that vary over many orders of magnitude, and with this type of temperature dependence.

The interpolation scheme used for model interpolation bases the interpolation on the difference between the ambient temperature and the **TNOM** value of the first model card in the netlist, which can sometimes lead to poorly conditioned interpolation. Thus it is often best that the first model card in the netlist be the one that has the “middle” **TNOM**, as in the example above. This ensures that no matter where in the range of temperature values the ambient temperature lies, it is a minimal distance from the base point of the interpolation.

## 2.1.15 .OP (Bias Point Analysis)

The .OP command causes detailed information about the bias point to be printed.

### General Form .OP

---

**Comments** This type of analysis can be specified by itself, in which case **Xyce** will run a nominal operating point. However, if specified with another analysis type, no additional operating point will be calculated, as most analyses require a DC operating point for initialization.

.OP outputs the parameters for all the device models and all the device instances present in the circuit. For large circuits, this can be a very large amount of output, so use with caution.

If no analysis command is provided, .OP will run a DC Operating Point calculation.

# 2.1.16 .OPTIONS Statements

Set various simulation limits, analysis control parameters and output parameters. In general, they use the following format:

**General Form**     .OPTIONS <pkg> [<name>=<value>]\*

---

**Examples**             .OPTIONS TIMEINT ABSTOL=1E-8

---

**Arguments and Options**

pkg	DEVICE	Device Model
	TIMEINT	Time Integration
	NONLIN	Nonlinear Solver
	NONLIN-TRAN	Transient Nonlinear Solver
	NONLIN-HB	HB Nonlinear Solver
	LOCA	Continuation/Bifurcation Tracking
	LINSOL	Linear Solver
	LINSOL-HB	HB Linear Solver
	OUTPUT	Output
	RESTART	Restart
	SENSITIVITY	Direct and Adjoint sensitivities
	HBINT	Harmonic Balance (HB)

name

value

The name of the parameter and the value it will be assigned.

---

**Comments**           Exceptions to this format are the OUTPUT and RESTART options, which use their own format. They are defined under their respective descriptions.

The designator *pkg* refers loosely to a *module* in the code. Thus, the term is used here as identifying a specific module to be controlled via *options* set in the netlist input file.

## .OPTIONS DEVICE (Device Package Options)

The device package parameters listed in Table 2.1 outline the options available for specifying device specific parameters. Some of these (DEFAS, DEFAD, TNOM etc.) have the same meaning as they do for the .OPTION line from Berkeley SPICE (3f5). Parameters which apply globally to all device models will be specified here. Parameters specific to a particular device instance or model are specified in section 2.3.

Table 2.1: Options for Device Package

Option	Description	Default
DEFAD	MOS Drain Diffusion Area	0.0
DEFAS	MOS Source Diffusion Area	0.0
DEFL	MOS Default Channel Length	1.0E-4
DEFW	MOS Default Channel Width	1.0E-4
GMIN	Minimum Conductance	1.0E-12
MINRES	This is a minimum resistance to be used in place of the default zero value of semiconductor device internal resistances. It is only used when model specifications (.MODEL cards) leave the parameter at its default value of zero, and is not used if the model explicitly sets the resistance to zero.	0.0
MINCAP	This is a minimum capacitance to be used in place of the default zero value of semiconductor device internal capacitances. It is only used when model specifications (.MODEL cards) leave the parameter at its default value of zero, and is not used if the model explicitly sets the capacitance to zero.	0.0
TEMP	Temperature	27.0 °C (300.15K)
TNOM	Nominal Temperature	27.0 °C (300.15K)
NUMJAC	Numerical Jacobian flag (only use for small problems)	0 (FALSE)
VOLTLIM	Voltage limiting	1 (TRUE)
icFac	This is a multiplicative factor which is applied to right-hand side vector loads of .IC initial conditions during the DCOP phase.	10000.0
LAMBERTW	This flag determines if the Lambert-W function should be applied in place of exponentials in hard-to-solve devices. This capability is implemented in the diode and BJT. Try this for BJT circuits that have convergence problems. For best effect, this option should be tried with voltlim turned off. A detailed explanation of the Lambert-W function, and its application to device modeling can be found in reference [5].	0 (FALSE)
MAXTimestep	Maximum time step size	1.0E+99
<b><i>MOSFET Homtopy parameters</i></b>		
VDSSCALEMIN	Scaling factor for Vds	0.3
VGSTCONST	Initial value for Vgst	4.5 Volt
LENGTHO	Initial value for length	5.0e-6
WIDTHO	Initial value for width	200.0e-6

Table 2.1: Options for Device Package

Option	Description	Default
TOX0	Initial value for oxide thickness	6.0e-8
<b><i>Debug output parameters</i></b>		
DEBUGLEVEL	The higher this number, the more info is output	1
DEBUGMINTIMESTEP	First time-step debug information is output	0
DEBUGMAXTIMESTEP	Last time-step of debug output	65536
DEBUGMINTIME	Same as DEBUGMINTIMESTEP except controlled by time (sec.) instead of step number	0.0
DEBUGMAXTIME	Same as DEBUGMAXTIMESTEP except controlled by time (sec.) instead of step number	100.0

### **.OPTIONS TIMEINT (Time Integration Options)**

The time integration parameters listed in Table 2.2 give the available options for helping control the time integration algorithms for transient analysis.

Time integration options are set using the `.OPTIONS TIMEINT` command.

Table 2.2: Options for Time Integration Package.

Option	Description	Default
METHOD	Time integration method. This parameter is only relevant when running <b>Xyce</b> in transient mode. Supported methods: <ul style="list-style-type: none"> <li>■ bdf or 6 (Backward Difference Formula orders 1-5)</li> <li>■ trap or 7 (variable order Trapezoid)</li> <li>■ gear or 8 (Gear method)</li> </ul>	trap or 7 (variable order Trapezoid)
RELTOL	Relative error tolerance	1.0E-03
ABSTOL	Absolute error tolerance	1.0E-06
RESTARTSTEPSCALE	This parameter is a scalar which determines how small the initial time step out of a breakpoint should be. In the current version of the time integrator, the first step after a breakpoint isn't subjected to much error analysis, so for very stiff circuits, this step can be problematic.	0.005

Table 2.2: Options for Time Integration Package.

Option	Description	Default
NLNEARCONV	This flag sets if “soft” failures of the nonlinear solver, when the convergence criteria are almost, but not quite, met, should result in a “success” code being returned from the nonlinear solver to the time integrator. If this is enabled, it is expected that the error analysis performed by the time integrator will be the sole determination of whether or not the time step is considered a “pass” or a “fail”. This is on by default, but occasionally circuits need tighter convergence criteria.	0 (FALSE)
NLSMALLUPDATE	This flag is another “soft” nonlinear solver failure flag. In this case, if the flag is set, time steps in which the nonlinear solver stalls, and is using updates that are numerically tiny, can be considered to have converged by the nonlinear solver. If this flag is set, the time integrator is responsible for determining if a step should be accepted or not.	1 (TRUE)
RESETTRANLS	The nonlinear solver resets its settings for the transient part of the run to something more efficient (basically a simpler set of options with smaller numbers for things like max Newton step). If this is set to false, this resetting is turned off. Normally should be left as default.	1 (TRUE)
MAXORD	This parameter determines the maximum order of integration that time integrators will attempt. For BDF 1-5, this can be reduced down to 1 to use Backward Euler. Setting this option does not guarantee that the integrator will integrate at this order, it just sets the maximum order the integrator will attempt. In order to guarantee a particular order is used, see the option MINORD below.	5 for BDF 1-5, 2 for variable order Trapezoid and Gear
MINORD	This parameter determines the minimum order of integration that time integrators will attempt to maintain. The integrator will start at Backward Euler and move up in order as quickly as possible to achieve MINORD and then it will keep the order above this. If MINORD is set at 2 and MAXORD is set at 2, then the integrator will move to second order as quickly as possible and stay there.	1
NEWLTE	This flag sets a new and more aggressive local truncation error estimation strategy to speedup the simulation. Note the default reltol is 1e-3 with newlte.	1 (TRUE)

Table 2.2: Options for Time Integration Package.

Option	Description	Default
NEWBPSTEPPING	<p>This flag sets a new time stepping method after a break point. Previously, <b>Xyce</b> treats each breakpoint identically to the DCOP point, in which the initial time step out of the DCOP is made to be very very small, because the LTE calculation is unreliable. As a result, <b>Xyce</b> takes an incredibly small step out of each breakpoint and then tries to grow the stepsize from there. When NEWBPSTEPPING is set, <b>Xyce</b> can take a reasonable large step out of every non-DCOP breakpoint, and then just relies on the step control to ensure that the step is small enough.</p> <p>Note that the new time stepping method after a break point does not work well with the old LTE calculation since the old LTE calculation is conservative and it tends to reject the first time step out of a break point. We recommend to use newlte if you choose to use the new time stepping method out of a break point. When using BDF15 method, newbpstepping should be disabled since no new time stepping strategy is implemented for BDF15 method.</p>	1 (TRUE)
ERROPTION	<p>This parameter determines if Local Truncation Error (LTE) control is turned on or not. If ERROPTION is on, then step-size selection is based on the number of Newton iterations nonlinear solve. For BDF15, if the nonlinear solve succeeds, then the step is doubled, otherwise it is cut by one eighth. For Trapezoid and Gear, if the number of nonlinear iterations is below NLMIN then the step is doubled. If the number of nonlinear iterations is above NLMAX then the step is cut by one eighth. In between, the step-size is left alone. Because this option can lead to very large time-steps, it is very important to specify an appropriate DELMAX option. If the circuit has breakpoints, then the option MINTIMESTEPSBP can also help to adjust the maximum time-step by specifying the minimum number of time points between breakpoints.</p>	0 (Local Truncation Error is used)
NLMIN	<p>This parameter determines the lower bound for the desired number of nonlinear iterations during a Trapezoid time or Gear integration solve with ERROPTION=1.</p>	3
NLMAX	<p>This parameter determines the upper bound for the desired number of nonlinear iterations during a Trapezoid time or Gear integration solve with ERROPTION=1.</p>	8
DELMAX	<p>This parameter determines the maximum time step-size used with ERROPTION=1. If a maximum time-step is also specified on the .TRAN line, then the minimum of that value and DELMAX is used.</p>	1e99

Table 2.2: Options for Time Integration Package.

Option	Description	Default
MINTIMESTEPSBP	This parameter determines the minimum number of time-steps to use between breakpoints. This enforces a maximum time-step between breakpoints equal to the distance between the last breakpoint and the next breakpoint divided by MINTIMESTEPSBP.	10
TIMESTEPSREVERSAL	This parameter determines whether time-steps are rejected based upon the step-size selection strategy in ERROPTION=1. If it is set to 0, then a step will be accepted with successful nonlinear solves independent of whether the number of nonlinear iterations is between NLMIN and NLMAX. If it is set to 1, then when the number of nonlinear iterations is above NLMAX, the step will be rejected and the step-size cut by one eighth and retried. If ERROPTION=0 (use LTE) then TIMESTEPSREVERSAL=1 (reject steps) is set. This has the consequence that for the BDF15 integrator, TIMESTEPREVERSAL=1.	0 (do not reject steps)
DOUBLED COPSTEP	This option should only be set to TRUE for a PDE device run. PDE devices often have to solve an extra "setup" problem to get the initial condition. This extra setup problem solves a nonlinear Poisson equation (see the device appendix for more details), while the normal step solves a full drift-diffusion(DD) problem. The name of this flag refers to the fact that the code is essentially taking two DC operating point steps instead of one. If you set this to TRUE, but have no PDE devices in the circuit, the code will repeat the same identical DCOP step twice. Generally there is no point in doing this.	0 (FALSE), if no PDE devices are present. 1 (TRUE), if at least one PDE device is in the circuit.
FIRSTDCOPSTEP	This is the index of the first DCOP step taken in a simulation for which DOUBLED COPSTEP is set to TRUE. The special initialization (nonlinear Poisson) step is referred to as step 0, while the normal (drift-diffusion) step is indexed with a 1. These two options(FIRSTDCOPSTEP and LASTDCOPSTEP) allow you to set the 1st or second DCOP step to be either kind of step. If FIRSTDCOPSTEP and LASTDCOPSTEP are both set to 0, then only the initial setup step happens. If FIRSTDCOPSTEP and LASTDCOPSTEP are both set to 1, then the initialization step doesn't happen, and only the real DD problem is attempted, with a crude initial guess. You should <i>never</i> set FIRSTDCOPSTEP to 1 and SECONDDCOPSTEP to 0. Normally, they should always be left as the defaults.	0
LASTDCOPSTEP	This is the second step taken in a simulation for which DOUBLED COPSTEP is set to TRUE.	1
BPENABLE	Flag for turning on/off breakpoints (1 = ON, 0 = OFF). It is unlikely anyone would ever set this to FALSE, except to help debug the breakpoint capability.	1 (TRUE)

Table 2.2: Options for Time Integration Package.

Option	Description	Default
EXITTIME	If this is set to nonzero, the code will check the simulation time at the end of each step. If the total time exceeds the exittime, the code will ungracefully exit. This is a debugging option, the point of which is to have the code stop at a certain time during a run without affecting the step size control. If not set by the user, it isn't activated.	-
EXITSTEP	Same as EXITTIME, only applied to step number. The code will exit at the specified step. If not set by the user, it isn't activated.	-

### **.OPTIONS HBINT (Harmonic Balance Options)**

The Harmonic Balance parameters listed in Table 2.3 give the available options for helping control the harmonic balance algorithms for harmonic balance analysis.

Harmonic Balance options are set using the `.OPTIONS HBINT` command.

Table 2.3: Options for HB.

Option	Description	Default
NUMFREQ	Number of harmonic frequencies to be calculated. Must be an odd number.	21
STARTUPPERIODS	Number of periods to integrate through before calculating the initial conditions. This option is only used when TAHB=1.	0
SAVEICDATA	Write out the initial conditions to a file.	0
TAHB	This flag sets transient assisted HB. When TAHB=0, transient analysis is not performed to get an initial guess. When TAHB=1, it uses transient analysis to get an initial guess.	1
VOLTLM	This flag sets voltage limiting for HB. During the initial guess calculation which normally uses transient simulation, the voltage limiting flag is determined by <code>.options device voltlim</code> . During the HB phase, the voltage limiting flag is determined by <code>.options hbint voltlim</code> .	1

### **.OPTIONS NONLIN (Nonlinear Solver Options)**

The nonlinear solver parameters listed in Table 2.4 provide methods for controlling the nonlinear solver for DC, Transient and harmonic balance. Note that the nonlinear solver options for DCOP, transient and harmonic balance are specified in separate options statements, using `.OPTIONS NONLIN`, `.OPTIONS NONLIN-TRAN` and `.OPTIONS NONLIN-HB`, respectively. The defaults for each

are specified in the third and fourth columns of Table 2.4.

Table 2.4: Options for Nonlinear Solver Package.

Option	Description	NONLIN Default	NONLIN- TRAN Default
NOX	Use NOX nonlinear solver.	1 (TRUE)	1 (TRUE)
NLSTRATEGY	Nonlinear solution strategy. Supported Strategies: <ul style="list-style-type: none"> <li>■ 0 (Newton)</li> <li>■ 1 (Gradient)</li> <li>■ 2 (Trust Region)</li> </ul>	0 (Newton)	0 (Newton)
SEARCHMETHOD	Line-search method used by the nonlinear solver. Supported line-search methods: <ul style="list-style-type: none"> <li>■ 0 (Full Newton - no line search)</li> <li>■ 1 (Interval Halving)</li> <li>■ 2 (Quadratic Interpolation)</li> <li>■ 3 (Cubic Interpolation)</li> <li>■ 4 (More'-Thuente)</li> </ul>	0 (Full New- ton) (NOTE: for itera- tive linear solves, the default is Quadratic Linesearch - 2)	0 (Full Newton)
CONTINUATION	Enables the use of Homotopy/Continuation algorithms for the nonlinear solve. Options are: <ul style="list-style-type: none"> <li>■ 0 (Standard nonlinear solve)</li> <li>■ 1 (Natural parameter homotopy. See LOCA options list)</li> <li>■ 2/mos (Specialized dual parameter homotopy for MOSFET circuits)</li> <li>■ 3/gmin (GMIN stepping, similar to that of SPICE)</li> </ul>	0 (Standard nonlinear solve)	0 (Standard nonlinear solve)
ABSTOL	Absolute residual vector tolerance	1.0E-12	1.0E-06
RELTOL	Relative residual vector tolerance	1.0E-03	1.0E-02
DELTAXTOL	Weighted nonlinear-solution update norm convergence tolerance	1.0	0.33
RHSTOL	Residual convergence tolerance (unweighted 2-norm)	1.0E-06	1.0E-02
SMALLUPDATETOL	Minimum acceptable norm for weighted nonlinear-solution update	1.0E-06	1.0E-06
MAXSTEP	Maximum number of Newton steps	200	20
MAXSEARCHSTEP	Maximum number of line-search steps	2	2
NORMLVL	Norm level used by the nonlinear solver algorithms (NOTE: not used for convergence tests)	2	2
IN_FORCING	Inexact Newton-Krylov forcing flag	0 (FALSE)	0 (FALSE)

Table 2.4: Options for Nonlinear Solver Package.

Option	Description	NONLIN Default	NONLIN- TRAN Default
AZ_TOL	Sets the minimum allowed linear solver tolerance. Valid only if IN_FORCING=1.	1.0E-12	1.0E-12
RECOVERYSTEPTYPE	<p>If using a line search, this option determines the type of step to take if the line search fails. Supported strategies:</p> <p>■ 0 (Take the last computed step size in the line search algorithm)</p> <p>■ 1 (Take a constant step size set by RECOVERYSTEP)</p>	0	0
RECOVERYSTEP	Value of the recovery step if a constant step length is selected	1.0	1.0
DLSDEBUG	Debug output for direct linear solver	0 (FALSE)	0 (FALSE)
DEBUGLEVEL	The higher this number, the more info is output	1	1
DEBUGMINTIMESTEP	First time-step debug information is output	0	0
DEBUGMAXTIMESTEP	Last time-step of debug output	99999999	99999999
DEBUGMINTIME	Same as DEBUGMINTIMESTEP except controlled by time (sec.) instead of step number	0.0	0.0
DEBUGMAXTIME	Same as DEBUGMAXTIMESTEP except controlled by time (sec.) instead of step number	1.0E+99	1.0E+99
<b>Parameters not supported by NOX</b>			
LINOPT	Linear optimization flag	0 (FALSE)	0 (FALSE)
CONSTRAINTBT	Constraint backtracking flag	0 (FALSE)	0 (FALSE)
CONSTRAINTMAX	Global maximum setting for constraint backtracking	DBL_MAX (Machine Dependent Constant)	DBL_MAX
CONSTRAINTMIN	Global minimum setting for constraint backtracking	-DBL_MAX (Machine Dependent Constant)	-DBL_MAX
CONSTRAINTCHANGE	Global percentage-change setting for constraint backtracking	sqrt(DBL_MAX) (Machine Dependent Constant)	sqrt(DBL_MAX)

### .OPTIONS LOCA (Continuation and Bifurcation Tracking Package Options)

The continuation selections listed in Table 2.5 provide methods for controlling continuation and bifurcation analysis. These override the defaults and any that were set simply in the continuation package. This option block is only used if the nonlinear solver or transient nonlinear solver enable continuation through the CONTINUATION flag.

There are two specialized homotopy methods that can be set in the nonlinear solver options line. One is MOSFET-based homotopy, which is specific to MOSFET circuits. This is specified using `continuation=2` or `continuation=mos`. The other is GMIN stepping, which is specified using `continuation=3` or `continuation=gmin`. For either of these methods, while it is possible to modify their default LOCA options, it is generally not necessary to do so. Note that **Xyce** automatically attempts GMIN stepping if the initial attempt to find the DC operating point fails. If GMIN stepping is specified in the netlist, **Xyce** will not attempt to find a DC operating point without GMIN stepping.

LOCA options are set using the `.OPTIONS LOCA` command.

Table 2.5: Options for Continuation and Bifurcation Tracking Package.

Option	Description	Default
STEPPER	Stepping algorithm to use:	
	■ 0 (Natural or Zero order continuation)	0 (Natural)
	■ 1 (Arc-length continuation)	
PREDICTOR	Predictor algorithm to use:	
	■ 0 (Tangent)	0 (Tangent)
	■ 1 (Secant)	
	■ 2 (Random)	
	■ 3 (Constant)	
STEPCONTROL	Algorithm used to adjust the step size between continuation steps:	
	■ 0 (Constant)	0 (Constant)
	■ 1 (Adaptive)	
CONPARAM	Parameter in which to step during a continuation run	VA:V0
INITIALVALUE	Starting value of the continuation parameter	0.0
MINVALUE	Minimum value of the continuation parameter	-1.0E20
MAXVALUE	Maximum value of the continuation parameter	1.0E20
BIFPARAM	Parameter to compute during bifurcation tracking runs	VA:V0
MAXSTEPS	Maximum number of continuation steps (includes failed steps)	20
MAXNLITERS	Maximum number of nonlinear iterations allowed (set this parameter equal to the MAXSTEP parameter in the NONLIN option block)	20
INITIALSTEPSIZE	Starting value of the step size	1.0
MINSTEPSIZE	Minimum value of the step size	1.0E20
MAXSTEPSIZE	Maximum value of the step size	1.0E-4

Table 2.5: Options for Continuation and Bifurcation Tracking Package.

Option	Description	Default
AGGRESSIVENESS	Value between 0.0 and 1.0 that determines how aggressive the step size control algorithm should be when increasing the step size. 0.0 is a constant step size while 1.0 is the most aggressive.	0.0
RESIDUALCONDUCTANCE	If set to a nonzero (small) number, this parameter will force the GMIN stepping algorithm to stop and declare victory once the artificial resistors have a conductance that is smaller than this number. This should only be used in transient simulations, and <i>ONLY</i> if it is absolutely necessary to get past the DC operating point calculation. It is almost always better to fix the circuit so that residual conductance is not necessary.	0.0

### .OPTIONS LINSOL (Linear Solver Options)

**Xyce** uses both sparse direct solvers as well as Krylov iterative methods for the solution of the linear equations generated by Newton's method. For the advanced users, there are a variety of options that can be set to help improve these solvers. Transformations of the linear system have a "TR\_" prefix on the flag. Many of the options for the Krylov solvers are simply passed through to the underlying Trilinos/AztecOO solution settings and thus have an "AZ\_" prefix on the flag.

Linear solver options are set using the .OPTIONS LINSOL command.

Table 2.6: Options for Linear Solver Package.

Option	Description	Default
TYPE	Determines which linear solver will be used.	
	■ KLU	
	■ SuperLU (optional)	
	■ AztecOO	KLU
	■ Belos	(Serial)
	■ ShyLU (optional)	AztecOO
	Note that while KLU and SuperLU (optional) are available for parallel execution they will solve the linear system in serial. Therefore they will be useful for moderate problem sizes but will not scale in memory or performance for large problems	(Parallel)
TR_partition	Perform load-balance partitioning on the linear system	0 (NONE, Serial) 1 (Isorropia, Parallel)
TR_partition_type	Type of load-balance partitioning on the linear system	"GRAPH"

Table 2.6: Options for Linear Solver Package.

Option	Description	Default
TR_singleton_filter	Triggers use of singleton filter for linear system	0 (FALSE, Serial) 1 (TRUE, Parallel)
TR_amd	Triggers use of approximate minimum-degree (AMD) ordering for linear system	0 (FALSE, Serial) 1 (TRUE, Parallel)
TR_global_btf	Triggers use of block triangular form (BTF) ordering for linear system, requires TR_amd=0 and TR_partition=0	0 (FALSE)
TR_reindex	Reindexes linear system parallel global indices in lexicographical order, recommended with singleton filter	1 (TRUE)
TR_solvermap	Triggers remapping of column indices for parallel runs, recommended with singleton filter	1 (TRUE)
<b><i>Iterative linear solver parameters</i></b>		
adaptive_solve	Triggers use of AztecOO adaptive solve algorithm for preconditioning of iterative linear solves	0 (FALSE)
use_aztec_precond	Triggers use of native AztecOO preconditioners for the iterative linear solves	0 (FALSE)
AZ_max_iter	Maximum number of iterative solver iterations	500
AZ_precond	AztecOO iterative solver preconditioner flag (used only when use_aztec_precond=1)	AZ_dom_decomp (14)
AZ_solver	Iterative solver type	AZ_gmres (1)
AZ_conv	Convergence type	AZ_r0 (0)
AZ_pre_calc	Type of precalculation	AZ_recalc (1)
AZ_keep_info	Retain calculation info	AZ_true (1)
AZ_orthog	Type of orthogonalization	AZ_modified (1)
AZ_subdomain_solve	Subdomain solution for domain decomposition preconditioners	AZ_ilut (9)
AZ_ilut_fill	Approximate allowed fill-in factor for the ILUT preconditioner	2.0
AZ_drop	Specifies drop tolerance used in conjunction with LU or ILUT preconditioners	1.0E-03
AZ_reorder	Reordering type	AZ_none (0)
AZ_scaling	Type of scaling	AZ_none (0)
AZ_kspace	Maximum size of Krylov subspace	500

Table 2.6: Options for Linear Solver Package.

Option	Description	Default
AZ_tol	Convergence tolerance	1.0E-12
AZ_output	Output level	AZ_none (0) 50 (if verbose build)
AZ_diagnostics	Diagnostic information level	AZ_none (0)
AZ_overlap	Schwarz overlap level for ILU preconditioners	0
AZ_rthresh	Diagonal shifting relative threshold for ILU preconditioners	1.0001
AZ_athresh	Diagonal shifting absolute threshold for ILU preconditioners	1.0E-04
ShyLU_rthresh	Relative dropping threshold for Schur complement preconditioner (ShyLU only)	1.0E-03

### **.OPTIONS LINSOL-HB (Linear Solver Options)**

For harmonic balance (HB) analysis, only Krylov iterative methods are available for the solution of the steady state. Furthermore, only matrix-free techniques are available for preconditioning the HB Jacobian, so many of the standard linear solver options are not available. The linear solver options for HB are set using the `.OPTIONS LINSOL-HB` command.

Table 2.7: Options for Linear Solver Package for HB.

Option	Description	Default
TYPE	Determines which linear solver will be used.	AztecOO
	■ AztecOO	
	■ Belos	
AZ_kspace	Maximum size of Krylov subspace	500
AZ_tol	Convergence tolerance	1.0E-12
prec_type	Preconditioning type	"none"

### **.OPTIONS OUTPUT (Output Options)**

The main purpose of the `.OPTIONS OUTPUT` command is to allow control of the output frequency of data to files specified by `.PRINT TRAN` commands. The format is:

```
.OPTIONS OUTPUT INITIAL_INTERVAL=<interval> [<t0> <i0> [<t1> <i1>]* ]
```

where INITIAL\_INTERVAL=<interval> specifies the starting interval time for output and <tx> <ix> specifies later simulation times <tx> where the output interval will change to <ix>. The solution is output at the exact intervals requested; this is done by interpolating the solution to the requested time points.

## **.OPTIONS RESTART (Checkpointing Options)**

The .OPTIONS RESTART command is used to control all checkpoint output and restarting.

The checkpointing form of the .OPTIONS RESTART command takes the following format:

### **General Format:**

```
.OPTIONS RESTART [PACK=<0|1>] JOB=<job prefix>
+ [INITIAL_INTERVAL=<initial interval time> [<t0> <i0> [<t1> <i1>]* ]]
```

PACK=<0|1> indicates whether the restart data will be byte packed or not. Parallel restarts must always be packed while Windows/MingW runs are always not packed. Otherwise, data will be packed by default unless explicitly specified. JOB=<job prefix> identifies the prefix for restart files. The actual restart files will be the job name with the current simulation time appended (e.g. name1e-05 for JOB=name and simulation time 1e-05 seconds). Furthermore, INITIAL\_INTERVAL=<initial interval time> identifies the initial interval time used for restart output. The <tx> <ix> intervals identify times <tx> at which the output interval (<ix>) should change. This functionality is identical to that described for the .OPTIONS OUTPUT command.

## **Examples**

To generate checkpoints at every time step (default):

**Example:**            .OPTIONS RESTART JOB=checkpt

To generate checkpoints every 0.1  $\mu s$ :

**Example:**            .OPTIONS RESTART JOB=checkpt INITIAL\_INTERVAL=0.1us

To generate unpacked checkpoints every 0.1  $\mu s$ :

**Example:**            .OPTIONS RESTART PACK=0 JOB=checkpt INITIAL\_INTERVAL=0.1us

To specify an initial interval of 0.1  $\mu s$ , at 1  $\mu s$  change to interval of 0.5  $\mu s$ , and at 10  $\mu s$  change to interval of 0.1  $\mu s$ :

### **Example:**

```
.OPTIONS RESTART JOB=checkpt INITIAL_INTERVAL=0.1us 1.0us
+ 0.5us 10us 0.1us
```

## **.OPTIONS RESTART (Restarting Options)**

To restart from an existing restart file, specify the file by either `FILE=<restart file name>` to explicitly use a restart file or by `JOB=<job name> START_TIME=<specified name>` to specify a file prefix and a specified time. The time must exactly match an output file time for the simulator to correctly identify the correct file. To continue generating restart output files, `INITIAL_INTERVAL=<interval>` and following intervals can be appended to the command in the same format as described above. New restart files will be packed according to the previous restart file read in.

The restarting form of the `.OPTIONS RESTART` command takes the following format:

### **General Format:**

```
.OPTIONS RESTART FILE=<restart file name>|JOB=<job name> START_TIME=<time>
+ [ INITIAL_INTERVAL=<interval> [<t0> <i0> [<t1> <i1>]* ]]
```

### **Examples**

Example restarting from checkpoint file at  $0.133\ \mu s$ :

**Example:** `.OPTIONS RESTART JOB=checkpoint START_TIME=0.133us`

To restart from checkpoint file at  $0.133\ \mu s$ :

**Example:** `.OPTIONS RESTART FILE=checkpoint0.000000133`

Restarting from  $0.133\ \mu s$  and continue checkpointing at  $0.1\ \mu s$  intervals:

### **Example:**

```
.OPTIONS RESTART FILE=checkpoint0.000000133 JOB=checkpoint_again
+ INITIAL_INTERVAL=0.1us
```

## **.OPTIONS RESTART: special notes for use with two-level-Newton**

Large parallel problems which involve power supply parasitics often require a two-level solve, in which different parts of the problem are handled separately. In most respects, restarting a two-level simulation is similar to restarting a conventional simulation. However, there are a few differences:

- When running with a two-level algorithm, **Xyce** requires (at least) two different input files. In order to do a restart of a two-level **Xyce** simulation, it is necessary to have an `.OPTIONS RESTART` statement in each file.
- It is necessary for the statements to be consistent. For example, the output times must be exactly the same, meaning the initial intervals must be exactly the same.

- **Xyce** will *not* check to make sure that the restart options used in different files match, so it is up to the user to ensure matching options.
- Finally, as each netlist that is part of a two-level solve will have its own `.OPTIONS RESTART` statement, that means that each netlist will generate and/or use its own set of restart files. As a result, the restart file name used by each netlist must be unique.

### `.OPTIONS SENSITIVITY` (Direct and Adjoint Sensitivity Options)

The sensitivity selections listed in Table 2.8 provide methods for controlling direct and adjoint sensitivity analysis.

SENSITIVITY options are set using the `.OPTIONS SENSITIVITY` command. They are only used if the netlist also includes a `.SENS` statement.

Table 2.8: Options for Sensitivity Package.

Option	Description	Default
ADJOINT	Flag to enable adjoint sensitivity calculation	false
DIRECT	Flag to enable direct sensitivity calculation	false
OUTPUTSCALED	Flag to enable output of scaled sensitivities	false
OUTPUTUNSCALED	Flag to enable output of unscaled sensitivities	true
STDOUTPUT	Flag to enable output of sensitivities to std output	true

## 2.1.17 .PARAM (Parameter)

User defined parameter that can be used in expressions throughout the netlist.

**General Form**     `.PARAM [<name>=<value>]*`

---

**Examples**            `.PARAM A_Param=1K`  
                         `.PARAM B_Param={A_Param*3.1415926535}`

---

**Arguments and Options**     `name`  
                                 `value`  
                                 The name of a parameter and its value.

---

**Comments**            Parameters defined using `.PARAM` are evaluated when the netlist is read in, and therefore must evaluate to constants at the time the netlist is parsed. It is therefore illegal to use any time- or solution-dependent terms in parameter definitions, including the `TIME` variable or any nodal voltages. Since they must be constants, these parameters may also not be used in `.STEP` loops.

## 2.1.18 .PREPROCESS REPLACEGOUND (Ground Synonym)

The purpose of ground synonym replacement is to treat nodes with the names GND, GND!, GROUND or any capital/lowercase variant thereof as synonyms for node 0. The general invocation is

**General Form**     .PREPROCESS REPLACEGOUND <bool>

---

### Arguments and Options

bool

If TRUE, **Xyce** will treat all instances of GND, GND!, GROUND, etc. as synonyms for node 0 but, if FALSE, **Xyce** will treat these nodes as separate. Only one .PREPROCESS REPLACEGOUND statement is permissible in a given netlist file.

## 2.1.19 .PREPROCESS REMOVEUNUSED (Removal of Unused Components)

If a given netlist file contains devices whose terminals are all connected to the same node (*e.g.*, R2 1 1 1M), it may be desirable to remove such components from the netlist before simulation begins. This is the purpose of the command

**General Form**     .PREPROCESS REMOVEUNUSED [<value>]

---

### Arguments and Options

value  
is a list of components separated by commas. The allowed values are

C	Capacitor
D	Diode
I	Independent Current Source
L	Inductor
M	MOSFET
Q	BJT
R	Resistor
V	Independent Voltage Source

---

### Examples

.PREPROCESS REMOVEUNUSED R,C

.PREPROCESS will attempt to search for all resistors and capacitors in a given netlist file whose individual device terminals are connected to the same node and remove these components from the netlist before simulation ensues. A list of components that are supported for removal is given above. Note that for MOSFETS and BJTs, three terminals on each device (the gate, source, and drain in the case of a MOSFET and the collector, base, and emitter in the case of a BJT) must be the same for the device to be removed from the netlist. As before, only one .PREPROCESS REMOVEUNUSED line is allowed in a given netlist file.

## 2.1.20 .PREPROCESS ADDRESISTORS (Adding Resistors to Dangling Nodes)

We refer to a *dangling node* as a circuit node in one of the following two scenarios: either the node is connected to only one device terminal, and/or the node has no DC path to ground. If several such nodes exist in a given netlist file, it may be desirable to automatically append a resistor of a specified value between the dangling node and ground. To add resistors to nodes which are connected to only one device terminal, one may use the command

**General Form**     `.PREPROCESS ADDRESISTORS ONETERMINAL <value>`

---

### Arguments and Options

`value`

is the value of the resistor to be placed between nodes with only one device terminal connection and ground. For instance, the command

---

### Examples

`.PREPROCESS ADDRESISTORS ONETERMINAL 1G`

will add resistors of value 1G between ground and nodes with only one device terminal connection and ground. The command

---

### Examples

`.PREPROCESS ADDRESISTORS NODCPATH <value>`

acts similarly, adding resistors of value <VALUE> between ground and all nodes which have no DC path to ground.

The `.PREPROCESS ADDRESISTORS` command is functionally different from either of the prior `.PREPROCESS` commands in the following way: while the other commands augment the netlist file for the current simulation, a `.PREPROCESS ADDRESISTORS` statement creates an auxiliary netlist file which explicitly contains a set of resistors that connect dangling nodes to ground. If the original netlist file containing a `.PREPROCESS ADDRESISTORS` statement is called `filename`, invoking **Xyce** on this file will produce a file `filename_xyce.cir` which contains the resistors that connect dangling nodes to ground. One can then run **Xyce** on this file to run a simulation in which the dangling nodes are tied to ground. Note that, in the original run on the file `filename`, **Xyce** will continue to run a simulation as usual after producing the file `filename_xyce.cir`, but this simulation will *not* include the effects of adding resistors between the dangling nodes and ground. Refer to the **Xyce** User's Guide for more detailed examples on the use of `.PREPROCESS ADDRESISTOR` statements.

Note that it is possible for a node to have one device terminal connection and, simultaneously, have no DC path to ground. In this case, if both a `ONETERMINAL` and `NODCPATH` command are invoked, only the resistor for the `ONETERMINAL` connection is added to the netlist; the `NODCPATH` connection is omitted.

As before, each netlist file is allowed to contain only one `.PREPROCESS ADDRESISTORS ONETERMINAL` and one `.PREPROCESS ADDRESISTORS NODCPATH` line each, or else **Xyce** will exit in error.

## 2.1.21 .PRINT (Print output)

Send analysis results to an output file.

**Xyce** allows multiple output files to be created during the run and supports several options for each.

**General Form**     `.PRINT <print type> [FORMAT=<STD|NOINDEX|PROBE|TECPLOT|RAW|CSV>]  
+ [FILE=<output filename>] [WIDTH=<print field width>]  
+ [PRECISION=<floating point output precision>]  
+ [FILTER=<absolute value below which a number outputs as 0.0>]  
+ [DELIMITER=<TAB|COMMA>] [TIMESCALEFACTOR=<real scale factor>]  
+ <output variable> [output variable]*`

---

**Examples**

```
.print tran format=tecplot V(1) I(Vsrc) {V(1)*(I(Vsrc)**2.0)}  
  
.PRINT TRAN FORMAT=PROBE FILE=foobar.csd V(1) {abs(V(1))-5.0}  
  
.PRINT DC FILE=foobar.txt WIDTH=19 PRECISION=15 FILTER=1.0e-10  
+ I(VSOURCE5) I(VSOURCE6)  
  
.print tran FORMAT=RAW V(1) I(Vsrc)  
  
R1 1 0 100  
X1 1 2 3 MySubcircuit  
V1 3 0 1V  
.SUBCKT MYSUBCIRCUIT 1 2 3  
R1 1 2 100K  
R2 2 4 50K  
R3 4 3 1K  
.ENDS  
  
.PRINT DC V(X1:4) V(2) I(V1)
```

---

### Arguments and Options

print type

A print type is the name of an analysis, one of the analysis specific print subtypes, or a specialized output command.

Analysis	Print Type	Description
.AC	AC	Sets default variable list and formats for print subtypes
.AC	AC_IC	Overrides variable list and format for AC initial conditions
.DC	DC	
.HB	HB	
.HB	HB_FD	Overrides variable list and format for HB frequency domain
.HB	HB_IC	Overrides variable list and format for HB initial conditions
.HB	HB_STARTUP	Overrides variable list and format for HB start up
.HB	HB_TD	Overrides variable list and format for HB time domain
.TRAN	TRAN	
<b>Specialized Output Commands</b>		
<i>Homotopy</i>	HOMOTOPY	Sets variable list and format for homotopy
.SENS	SENS	Sets variable list and format for sensitivity

A netlist may contain many .PRINT commands, but only commands with analysis types which are appropriate for the analysis being performed are processed. This feature allows you to generate multiple formats and variable sets in a single analysis run.

For analysis types that generate multiple output files, the print subtype allows you to specify variables and output parameters for each of those output files. If there is no .PRINT <subtype> provided in the netlist, the variables and parameters from the analysis type will be used.

FORMAT=<STD|NOINDEX|PROBE|TECLOT|RAW|CSV>

The output format may be specified using the FORMAT option. The STD format outputs the data divided up into data columns. The NOINDEX format is the same as the STD format except that the index column is omitted. The PROBE format specifies that the output should be formatted to be compatible with the PSpice Probe plotting utility. The TECLOT format specifies that the output should be formatted to be compatible with the Tecplot plotting program. The RAW format specifies that the output should comply with the SPICE binary rawfile format. Use with the **-a** command line option to output an ascii rawfile. The CSV format specifies that the output file should be a comma-separated value file with a header indicating the variables printed in the file. It is similar to, but not identical to using DELIMITER=COMMA; the latter will also print

a footer that is not compatible with most software that requires CSV format.

FILE=<output filename>

Specifies the name of the file to which the output will be written.

WIDTH=<print field width>

Controls the output width used in formatting the output.

PRECISION=<floating point precision>

Number of floating point digits past the decimal for output data.

FILTER=<filter floor value>

Used to specify the absolute value below which output variables will be printed as 0.0.

DELIMITER=<TAB|COMMA>

Used to specify an alternate delimiter in the STD or NOINDEX format output.

TIMESCALEFACTOR=<real scale factor>

Specify a constant scaling factor for time. Time is normally printed in units of seconds, but if one would like the units to be milliseconds, then set TIMESCALEFACTOR=1000.

<output variable>

Following the analysis type and other options is a list of output variables. There is no upper bound on the number of output variables. The output is divided up into data columns and output according to any specified options (see options given above). Output variables can be specified in four ways:

- V(<circuit node>) to output the voltage at <circuit node>
- V(<circuit node>,<circuit node>) to output the voltage difference between the first <circuit node> and second <circuit node>
- I(<device>) to output current through a two terminal device
- I<lead abbreviation>(<device>) to output current into a particular lead of a three or more terminal device (see the Comments, below, for details)
- N(<device internal variable>) to output a specific device internal variable (see the individual devices in Section 2.3 for syntax)
- {expression} to output an expression
- <device>:<parameter> to output a device parameter
- <model>:<parameter> to output a device parameter

When the analysis type is AC or HB, additional output variable formats are available:

- VR(<circuit node>) to output the real component of voltage response at a point in the circuit

- `VI(<circuit node>)` to output the imaginary component of voltage response at a point in the circuit
- `VM(<circuit node>)` to output the magnitude of voltage response
- `VP(<circuit node>)` to output the phase of voltage response
- `VDB(<circuit node>)` to output the magnitude of voltage response in decibels.
- `VR(<circuit node>,<circuit node>)` to output the real component of voltage response at a point in the circuit
- `VI(<circuit node>,<circuit node>)` to output the imaginary component of voltage response at a point in the circuit
- `VM(<circuit node>,<circuit node>)` to output the magnitude of voltage response
- `VP(<circuit node>,<circuit node>)` to output the phase of voltage response
- `VDB(<circuit node>,<circuit node>)` to output the magnitude of voltage response in decibels.
- `IR(<device>)` to output the real component of the current through a two terminal device
- `II(<device>)` to output the imaginary component of the current through a two terminal device
- `IM(<device>)` to output the magnitude of the current through a two terminal device
- `IP(<device>,<circuit node>)` to output the phase of the current through a two terminal device
- `IDB(<device>)` to output the magnitude of the current through a two terminal device in decibels.

In AC analysis, outputting a voltage node without any of these optional designators results in output of the real and imaginary parts of the signal. Note that under AC analysis, printing of device lead currents, e.g. `I(<device>)`, is not supported.

Note that when using the variable list for time domain output, usage of frequency domain functions like `VDB` can result in `-Inf` output being written to the output file. This is easily solved by defining the time domain equivalent command to specify the correct output for time domain data.

Further explanation of current specification is in comments section below.

---

## Comments

- Currents are positive flowing from node 1 to node 2 for two node devices, and currents are positive flowing into a particular lead for multi-terminal devices.
- `<circuit node>` is simply the name of any node in your top-level circuit, or `<subcircuit name>:<node>` to reference nodes that are internal to a subcircuit.

- `<device>` is the name of any device in your top-level circuit, or `<subcircuit name>:<device>` to reference devices that are internal to a subcircuit.
- `<lead abbreviation>` is a single character designator for individual leads on a device with three or more leads. For bipolar transistors these are: c (collector), b (base), e (emitter), and s (substrate). For mosfets, lead abbreviations are: d (drain), g (gate), s (source), and b (bulk). SOI transistors have: d, g, s, e (bulk), and b (body). For PDE devices, the nodes are numbered according to the order they appear, so lead currents are referenced like I1(`<device>`), I2(`<device>`), etc.
- The "lead current" method of printing from devices in Xyce is done at a low level with special code added to each device; the method is therefore only supported in specific devices that have this extra code. So, if `.PRINT I(Y)` does not work, for a device called Y, then you will need to attach an ammeter (zero-volt voltage source) in series with that device and print the ammeter's current instead.
- Lead currents of subcircuit ports are not supported. However, access is provided via specific node names (e.g., `X1:internalNodeName`) or specific devices (e.g., `X1:V3`) inside the subcircuit.
- Wildcards are partially supported on `.PRINT` lines. `V(*)` will print all of the node voltages in the circuit. `I(*)` will print all of the currents that are implemented in voltage sources. If, for example, an ammeter (zero-volt voltage source) is placed in series with a resistor then the current through that resistor would be included in the `I(*)` output.
- For STD formatted output, the values of the output variables are output as a series of columns (one for each output variable).
- When the command line option `-r <raw-file-name>` is used, all of the output is diverted to the *raw-file-name* file as a concatenation of the plots, and each plot includes all of the variables of the circuit. Using the `-a` options in conjunction with the `-r` option results in a raw file that is output all in ascii characters.
- Any output going to the same file from one simulation of **Xyce** results in the concatenation of output. However, if a simulation is re-run then the original output will be over-written.
- During analysis a number of output files may be generated. The selection of which files are created depends on a variety of factors, most obvious of which is the `.PRINT` command. See section 2.1 for more details.
- Frequency domain values are output as complex values for Raw, TecPlot and Probe formats when a complex variable is printed. For STD and CSV formats, the output appears in two columns, the real part followed by the imaginary part. The print variables VR, VI, VM, VDB and VP print the scalar values for the real part, imaginary part, magnitude, magnitude in decibels, and phase, respectively.
- When outputting a device or model parameter, it is usually necessary to specify both the device name and the parameter name, separated by a colon. For example, the saturation current of a diode model DMOD would be requested as `DMOD:IS`.

## Print AC Analysis

AC Analysis generates two output files, the primary output is in the frequency domain and the initial conditions output is in the time domain.

Note that when using the `.PRINT AC` to create the variable list for DC type output, usage of frequency domain functions like `VDB` can result in `-Inf` output being written to the output file. This is easily solved by defining a `.PRINT AC_IC` command to specify the correct output for initial condition data.

Homotopy and sensitivity output can also be generated.

Table 2.9: Print AC Analysis Type

Trigger	Files	Columns/Description
<code>.PRINT AC</code>	<i>circuit-file.FD.prn</i>	INDEX FREQ
<code>.PRINT AC FORMAT=NOINDEX</code>	<i>circuit-file.FD.prn</i>	FREQ
<code>.PRINT AC FORMAT=CSV</code>	<i>circuit-file.FD.csv</i>	FREQ
<code>.PRINT AC FORMAT=RAW</code>	<i>circuit-file.raw</i>	FREQ
<code>runxyce -a</code> <code>.PRINT AC FORMAT=RAW</code>	<i>circuit-file.raw</i>	FREQ
<code>.PRINT AC FORMAT=TECPLOT</code>	<i>circuit-file.FD.dat</i>	FREQ
<code>.PRINT AC FORMAT=PROBE</code>	<i>circuit-file.FD.csd</i>	–
<b>Add .OP To Netlist To Enable AC_IC Output</b>		
<code>.PRINT AC_IC</code>	<i>circuit-file.TD.prn</i>	INDEX TIME
<code>.PRINT AC_IC FORMAT=CSV</code>	<i>circuit-file.TD.csv</i>	TIME
<code>.PRINT AC_IC FORMAT=RAW</code>	<i>circuit-file.raw</i>	TIME
<code>runxyce -a</code> <code>.PRINT AC_IC FORMAT=RAW</code>	<i>circuit-file.raw</i>	TIME
<code>.PRINT AC_IC FORMAT=TECPLOT</code>	<i>circuit-file.TD.dat</i>	TIME
<code>.PRINT AC_IC FORMAT=PROBE</code>	<i>circuit-file.TD.csd</i>	–
<b>Command Line Raw Override Output</b>		
<code>runxyce -r</code>	<i>circuit-file.raw</i>	All circuit variables printed
<code>runxyce -r -a</code>	<i>circuit-file.raw</i>	All circuit variables printed
<b>Additional Output Available</b>		
<code>.OP</code>	<i>log file</i>	Operating point data
<code>.SENS</code> <code>.PRINT SENS</code>		see Print Sensitivity
<code>.OPTIONS NONLIN CONTINUATION=&lt;method&gt;</code> <code>.PRINT HOMOTOPY</code>		see Print Homotopy

## Print DC Analysis

DC Analysis generates output based on the format specified by the `.PRINT` command.

Homotopy and sensitivity output can also be generated.

Table 2.10: Print DC Analysis Type

Trigger	Files	Columns/Description
<code>.PRINT DC</code>	<i>circuit-file.prn</i>	INDEX
<code>.PRINT DC FORMAT=NOINDEX</code>	<i>circuit-file.prn</i>	—
<code>.PRINT DC FORMAT=CSV</code>	<i>circuit-file.csv</i>	—
<code>.PRINT DC FORMAT=RAW</code>	<i>circuit-file.raw</i>	—
<code>runxyce -a</code> <code>.PRINT DC FORMAT=RAW</code>	<i>circuit-file.raw</i>	—
<code>.PRINT DC FORMAT=TECPLOT</code>	<i>circuit-file.dat</i>	—
<code>.PRINT DC FORMAT=PROBE</code>	<i>circuit-file.csd</i>	—
<b>Command Line Raw Override Output</b>		
<code>runxyce -r</code>	<i>circuit-file.raw</i>	All circuit variables
<code>runxyce -r -a</code>	<i>circuit-file.raw</i>	All circuit variables
<b>Additional Output Available</b>		
<code>.OP</code>	<i>log file</i>	Operating point data
<code>.SENS</code> <code>.PRINT SENS</code>		see Print Sensitivity
<code>.OPTIONS NONLIN CONTINUATION=&lt;method&gt;</code> <code>.PRINT HOMOTOPY</code>		see Print Homotopy

## Print Harmonic Balance Analysis

HB Analysis generates two output files in the frequency domain and two in the time domain based on the format specified by the `.PRINT` command. Additional startup and initial conditions output can be generated based on `.OPTIONS` commands.

Note that when using the `.PRINT HB` to create the variable list for time domain output, usage of frequency domain functions like `VDB` can result in `-Inf` output being written to the output file. This is easily solved by defining a `.PRINT HB_TD`, `.PRINT HB_IC` and `.PRINT HB_STARTUP` commands to specify the correct output for initial condition time domain data.

Homotopy output can also be generated.

Table 2.11: Print HB Analysis Type

Trigger	Files	Columns/Description
.PRINT HB	circuit-file.HB.TD.prn circuit-file.HB.FD.prn circuit-file.hb_ic.prn	INDEX TIME INDEX FREQ INDEX TIME
.PRINT HB FORMAT=NOINDEX	circuit-file.HB.TD.prn circuit-file.HB.FD.prn circuit-file.hb_ic.prn	TIME FREQ TIME
.PRINT HB FORMAT=CSV	circuit-file.HB.TD.csv circuit-file.HB.FD.csv circuit-file.hb_ic.csv	TIME FREQ TIME
.PRINT HB FORMAT=TECPLOT	circuit-file.HB.TD.dat circuit-file.HB.FD.dat circuit-file.hb_ic.dat	TIME FREQ TIME
.PRINT HB_FD	circuit-file.HB.TD.prn	INDEX FREQ
.PRINT HB_FD FORMAT=NOINDEX	circuit-file.HB.FD.prn	FREQ
.PRINT HB_FD FORMAT=CSV	circuit-file.HB.FD.csv	FREQ
.PRINT HB_FD FORMAT=TECPLOT	circuit-file.HB.FD.dat	FREQ
.PRINT HB_TD	circuit-file.HB.TD.prn	INDEX TIME
.PRINT HB_TD FORMAT=NOINDEX	circuit-file.HB.TD.prn	TIME
.PRINT HB_TD FORMAT=CSV	circuit-file.HB.TD.csv	TIME
.PRINT HB_TD FORMAT=TECPLOT	circuit-file.HB.TD.dat	TIME
<b>Command Line Raw Override Output</b>		
runxyce -r	circuit-file.raw	All circuit variables printed
runxyce -r -a	circuit-file.raw	All circuit variables printed
<b>Startup Period</b>		
.OPTIONS HBINT STARTUPPERIODS=<n> .PRINT HB_STARTUP	circuit-file.startup.prn	INDEX TIME
.OPTIONS HBINT STARTUPPERIODS=<n> .PRINT HB_STARTUP FORMAT=NOINDEX	circuit-file.startup.prn	TIME
.OPTIONS HBINT STARTUPPERIODS=<n> .PRINT HB_STARTUP FORMAT=CSV .OPTIONS HBINT STARTUPPERIODS=<n>	circuit-file.startup.csv	TIME
.OPTIONS HBINT STARTUPPERIODS=<n> .PRINT HB_STARTUP FORMAT=TECPLOT .OPTIONS HBINT STARTUPPERIODS=<n>	circuit-file.startup.dat	TIME
<b>Initial Conditions</b>		
.OPTIONS HBINT SAVEICDATA=1 .PRINT HB_IC	circuit-file.hb_ic.prn	INDEX TIME
.OPTIONS HBINT SAVEICDATA=1 .PRINT HB_IC FORMAT=NOINDEX	circuit-file.hb_ic.prn	TIME

Table 2.11: Print HB Analysis Type

Trigger	Files	Columns/Description
.OPTIONS HBINT SAVEICDATA=1 .PRINT HB_IC FORMAT=CSV	<i>circuit-file.hb_ic.csv</i>	TIME
.OPTIONS HBINT SAVEICDATA=1 .PRINT HB_IC FORMAT=TECPLOT	<i>circuit-file.hb_ic.dat</i>	TIME
<b>Additional Output Available</b>		
.OP	<i>log file</i>	Operating point data
.SENS .PRINT SENS		see Print Sensitivity
.OPTIONS NONLIN CONTINUATION=<method> .PRINT HOMOTOPY		see Print Homotopy

## Print Transient Analysis

Transient Analysis generates time domain output based on the format specified by the .PRINT command.

Homotopy and sensitivity output can also be generated.

Table 2.12: Print Transient Analysis Type

Trigger	Files	Columns/Description
.PRINT TRAN	<i>circuit-file.prn</i>	INDEX TIME
.PRINT TRAN FORMAT=NOINDEX	<i>circuit-file.prn</i>	TIME
.PRINT TRAN FORMAT=CSV	<i>circuit-file.csv</i>	TIME
.PRINT TRAN FORMAT=RAW	<i>circuit-file.raw</i>	TIME
runxyce -a .PRINT TRAN FORMAT=RAW	<i>circuit-file.raw</i>	TIME
.PRINT TRAN FORMAT=TECPLOT	<i>circuit-file.dat</i>	TIME
.PRINT TRAN FORMAT=PROBE	<i>circuit-file.csd</i>	
<b>Command Line Raw Override Output</b>		
runxyce -r	<i>circuit-file.raw</i>	All circuit variables printed
runxyce -r -a	<i>circuit-file.raw</i>	All circuit variables printed
<b>Additional Output Available</b>		
.OP	<i>log file</i>	Operating point data
.SENS .PRINT SENS		see Print Sensitivity
.OPTIONS NONLIN CONTINUATION=<method> .PRINT HOMOTOPY		see Print Homotopy

## Print Homotopy

Homotopy output is generated by the inclusion of the  
.OPTIONS NONLIN CONTINUATION=<method> command.

Table 2.13: Print Homotopy

Trigger	Files	Columns/Description
.OPTIONS NONLIN CONTINUATION=<method> .PRINT <analysis-type>	circuit-file.HOMOTOPY.prn	INDEX TIME

## Print Sensitivity

Sensitivity is enabled by inclusion of the  
.SENS command.

Table 2.14: Print Sensitivities

Trigger	Files	Columns/Description
.SENS objfunc=<obj> p=[ $p_1$ ] [, $p_n$ ]* .PRINT SENS	circuit-file.SENS.prn	obj dobj/d( $p_1$ ) dobj/d( $p_n$ )
.SENS objfunc=<obj> p=[ $p_1$ ] [, $p_n$ ]* .PRINT SENS FORMAT=TECPLOT	circuit-file.SENS.prn	obj dobj/d( $p_1$ ) dobj/d( $p_n$ )

## Parameter Stepping

During parameter stepping, enabled with the .STEP command, the output generated by each of analysis types varies. Generally the FORMAT indicates this variation, however some combinations of analysis and format can result in additional variation.

The following table lists how the output differs for each analysis type and format.

Print Type	Format	Description
AC	STD	1, 3, 4, 11, 12, 13
AC	CSV	4, 11
AC	PROBE	16
AC	TECPLOT	4, 12, 13, 18
AC	RAW	19
AC	RAW (runxyce -a)	19
AC_IC	STD	1, 4, 11, 12, 13
AC_IC	CSV	4, 11
AC_IC	PROBE	16
AC_IC	TECPLOT	12, 13, 18

Print Type	Format	Description
AC_IC	RAW	19
AC_IC	RAW (runxyce -a)	19
DC	STD	1, 11, 12
DC	CSV	11
DC	PROBE	17
DC	TECPLOT	4, 12, 13, 18
DC	RAW	19
DC	RAW (runxyce -a)	19
HB_TD	STD	1, 2, 4, 11, 12, 13
HB_TD	CSV	11
HB_TD	TECPLOT	12, 13, 18
HB_FD	STD	1, 3, 4, 11, 12, 13
HB_FD	CSV	4, 11
HB_FD	TECPLOT	4, 12, 13, 18
HB_IC	STD	1, 2, 4, 11, 12, 13
HB_IC	CSV	11
HB_IC	TECPLOT	12, 13, 18
HB_STARTUP	STD	1, 2, 4, 11, 12, 13
HB_STARTUP	CSV	11
HB_STARTUP	TECPLOT	12, 13, 18
TRAN	STD	1, 2, 11, 12
TRAN	CSV	2, 11
TRAN	PROBE	17
TRAN	TECPLOT	2, 4, 12, 13, 18
TRAN	RAW	2, 19
TRAN	RAW (runxyce -a)	2, 19
<b><i>Specialized Output Commands</i></b>		
HOMOTOPY	STD	1, 2, 4, 11, 12, 13
HOMOTOPY	PROBE	17
HOMOTOPY	TECPLOT	2, 4, 12, 13, 18
SENSITIVITY	STD	1, 2, 11, 12, 14
SENSITIVITY	PROBE	17

### Description

- |   |  |
|---|--|
| 1 | INDEX column added to output variable list |
| 2 | TIME column added to output variable list  |
| 3 | FREQ column added to output variable list  |

Description	
4	Frequency domain data written as $\text{Re}(var)$ and $\text{Im}(var)$
11	INDEX resets to zero at start of each .STEP
12	Prints 'End of Xyce(TM) Parameter Sweep' at end of .STEP simulation
13	Prints 'End of Xyce(TM) Simulation' at end of non-.STEP simulation
14	Prints 'End of Xyce(TM) Sensitivity Simulation' at end of non-.STEP simulation
16	Two '#' at the end of each .STEP (BUG)
17	One '#' at end of each .STEP
18	New ZONE for each .STEP, and AUXDATA for each .STEP parameter
19	Prints 'Plotname: Step Analysis: Step $s$ of $n$ params' at the start of each .STEP

## 2.1.22 .SAVE (Save operating point conditions)

Stores the operating point of a circuit in the specified file for use in subsequent simulations. The data may be saved as .IC or .NODESET lines.

**General Form**     .SAVE [TYPE=<IC|NODESET>] [FILE=<filename>] [LEVEL=<all|none>]  
                      + [TIME=<save\_time>]

---

**Examples**        .SAVE TYPE=IC FILE=mycircuit.ic  
                      .SAVE TYPE=NODESET FILE=myothercircuit.ic  
  
                      .include mycircuit.ic

---

**Comments**        The file created by .SAVE will contain .IC or .NODESET lines containing all the voltage node values at the DC operating point of the circuit. The default **TYPE** is NODESET. The default filename is *netlist.cir.ic*.

                      The resulting file may be used in subsequent simulations to obtain quick DC convergence simply by including it in the netlist, as in the third example line above. **Xyce** has no corresponding .LOAD statement.

                      The **LEVEL** parameter is included for compatibility with HSPICE netlists. If none is specified, then no save file is created. The default **LEVEL** is all.

**TIME** is also an HSPICE compatibility parameter. This is unsupported in **Xyce**. **Xyce** outputs the save file only at time=0.0.

## 2.1.23 .SENS (Compute DC or transient sensitivities)

Computes sensitivities for a user-specified objective function with respect to a user-specified list of circuit parameters.

**General Form**     .SENS objfunc=<output expression> param=<circuit parameter(s)>

---

**Examples**     .SENS objfunc={0.5\*(V(B)-3.0)\*\*2.0} param=R1:R,R2:R  
                 .options SENSITIVITY direct=1 adjoint=1

---

**Comments**     This capability can be applied to either DC or transient analysis. For DC, both direct and adjoint sensitivities are supported, and the user can optionally request either direct or adjoint sensitivities, or both.

*For transient analysis, only direct sensitivities are supported.* If adjoint sensitivities are requested for a transient simulation, the code will attempt to compute them, but the results are not reliable.

For sensitivity analysis, it is necessary to specify circuit parameters on the .SENS line. Unlike the SPICE version, this capability will not automatically use every parameter in the circuit.

## 2.1.24 .STEP (Step Parametric Analysis)

Calculates a full analysis (.DC, .TRAN, .AC, etc.) over a range of parameter values. This type of analysis is very similar to .DC analysis. Similar to .DC analysis, .STEP supports sweeps which are linear, decade logarithmic, octave logarithmic, or a list of values.

LIN Linear sweep

The sweep variable is swept linearly from the starting to the ending value.

OCT Sweep by octaves

The sweep variable is swept logarithmically by octaves.

DEC Sweep by decades

The sweep variable is swept logarithmically by decades.

### Linear Sweeps

<b>General Form</b>	<code>.STEP &lt;parameter name&gt; &lt;initial&gt; &lt;final&gt; &lt;step&gt;</code>
---------------------	--

<b>Examples</b>	<code>.STEP TEMP -45 -55 -10</code> <code>.STEP R1 45 50 5</code> <code>.STEP C101:C 45 50 5</code> <code>.STEP DLEAK:IS 1.0e-12 1.0e-11 1.0e-12</code> <code>.STEP V1 20 10 -1</code>
-----------------	--

<b>Arguments and Options</b>	
------------------------------	--

<code>initial</code>	Initial value for the parameter.
<code>final</code>	Final value for the parameter.
<code>step</code>	Value that the parameter is incremented at each step.

<b>Comments</b>	<p>STEP parameter analysis will sweep a parameter from its initial value to its final value, at increments of the step size. At each step of this sweep, it will conduct a full analysis (.DC, .TRAN, .AC, etc.) of the circuit.</p> <p>The specification is similar to that of a .DC sweep, except that each parameter gets its own .STEP line in the input file, rather than specifying all of them on a single line.</p> <p>Output, as designated by a .PRINT statement, is slightly more complicated in the case of a .STEP simulation. If the user has specified a .PRINT line in the input file, <b>Xyce</b> will output two files. All steps of the sweep to a single output file as usual, but with the results of each step appearing one after another with</p>
-----------------	---

the “Index” column starting over at zero. Additionally, a file with a “.res” suffix will be produced indicating what parameters were used for each iteration of the step loops.

## Decade Sweeps

<b>General Form</b>	<code>.STEP DEC &lt;sweep variable name&gt; &lt;start&gt; &lt;stop&gt; &lt;points&gt;</code>
---------------------	--

---

<b>Examples</b>	<code>.STEP DEC VIN 1 100 2</code> <code>.STEP DEC R1 100 10000 3</code> <code>.STEP DEC TEMP 1.0 10.0 3</code>
-----------------	---

## Octave Sweeps

<b>General Form</b>	<code>.STEP OCT &lt;sweep variable name&gt; &lt;start&gt; &lt;stop&gt; &lt;points&gt;</code>
---------------------	--

---

<b>Examples</b>	<code>.STEP OCT VIN 0.125 64 2</code> <code>.STEP OCT TEMP 0.125 16.0 2</code> <code>.STEP OCT R1 0.015625 512 3</code>
-----------------	---

## List Sweeps

<b>General Form</b>	<code>.STEP &lt;sweep variable name&gt; LIST &lt;val&gt; &lt;val&gt; &lt;val&gt;...</code> <code>+ [&lt;sweep variable name&gt; LIST &lt;val&gt; &lt;val&gt; ...]...</code>
---------------------	--

---

<b>Examples</b>	<code>.STEP VIN LIST 1.0 2.0 10. 12.0</code> <code>.STEP TEMP LIST 8.0 21.0</code>
-----------------	---

## 2.1.25 .SUBCKT (Subcircuit)

The .SUBCKT statement begins a subcircuit definition by giving its name, the number and order of its nodes and the names and default parameters that direct its behavior. The .ENDS statement signifies the end of the subcircuit definition. See Section 2.3.26 for more information on using subcircuits with the X device.

**General Form**

```
.SUBCKT <name> [node]*  
+ [PARAMS: [<name>=<value>]* ]  
...  
.ENDS
```

---

**Examples**

```
.SUBCKT OPAMP 10 12 111 112 13  
...  
.ENDS  
  
.SUBCKT FILTER1 INPUT OUTPUT PARAMS: CENTER=200kHz,  
+ BANDWIDTH=20kHz  
...  
.ENDS  
  
.SUBCKT PLRD IN1 IN2 IN3 OUT1  
+ PARAMS: MNTYMXDELY=0 IO_LEVEL=1  
...  
.ENDS  
  
.SUBCKT 74LS01 A B Y  
+ PARAMS: MNTYMXDELY=0 IO_LEVEL=1  
...  
.ENDS
```

---

### Arguments and Options

name

The name used to reference a subcircuit.

node

An optional list of nodes. This is not mandatory since it is feasible to define a subcircuit without any interface nodes.

PARAMS:

Keyword that provides values to subcircuits as arguments for use as expressions in the subcircuit. Parameters defined in the PARAMS: section may be used in expressions within the body of the subcircuit and will take the default values specified in the subcircuit definition unless overridden by a PARAMS: section when the subcircuit is instantiated.

---

**Comments**

A subcircuit designation ends with a `.ENDS` command. The entire netlist between `.SUBCKT` and `.ENDS` is part of the definition. Each time the subcircuit is called via an `X` device, the entire netlist in the subcircuit definition replaces the `X` device.

There must be an equal number of nodes in the subcircuit call and in its definition. As soon as the subcircuit is called, the actual nodes (those in the calling statement) substitute for the argument nodes (those in the defining statement).

Node zero cannot be used in this node list, as it is the global ground node.

Subcircuit references may be nested to any level. Subcircuits definitions may also be nested; a `.SUBCKT` statement and its closing `.ENDS` may appear between another `.SUBCKT/.ENDS` pair. A subcircuit defined inside another subcircuit definition is local to the outer subcircuit and may not be used at higher levels of the circuit netlist.

Subcircuits should include only device instantiations and possibly these statements:

- `.MODEL` (model definition)
- `.PARAM` (parameter)
- `.FUNC` (function)

Models, parameters, and functions defined within a subcircuit are scoped to that definition. That is they are only accessible within the subcircuit definition in which they are included. Further, if a `.MODEL`, `.PARAM` or a `.FUNC` statement is included in the main circuit netlist, it is accessible from the main circuit as well as all subcircuits.

Node, device, and model names are scoped to the subcircuit in which they are defined. It is allowable to use a name in a subcircuit that has been previously used in the main circuit netlist. When the subcircuit is flattened (expanded into the main netlist), all of its names are given a prefix via the subcircuit instance name. For example, `Q17` becomes `X3:Q17` after expansion. After expansion, all names are unique. The single exception occurs in the use of global node names, which are not expanded.

## 2.1.26 .TRAN (Transient Analysis)

Calculates the time-domain response of a circuit for a specified duration.

**General Form**     `.TRAN <initial step value> <final time value>`  
                      `+ [<start time value> [<step ceiling value>]] [NOOP] [UIC]`  
                      `+ [{schedule( <time>, <maximum time step>, ... )}]`

---

**Examples**            `.TRAN 1us 100ms`  
                          `.TRAN 1ms 100ms 0ms .1ms`  
                          `.TRAN 0 2.0e-3 {schedule( 0.5e-3, 0, 1.0e-3, 1.0e-6, 2.0e-3, 0 )}`

---

### Arguments and Options

`initial step value`

Used to calculate the initial time step (see below).

`final time value`

Sets the end time (duration) for the analysis.

`start time value`

Sets the time at which output of the simulation results is to begin.  
Defaults to zero.

`step ceiling value`

Sets a maximum time step. Defaults to  $((\text{final time value}) - (\text{start time value})) / 10$ , unless there are breakpoints (see below).

`NOOP`

Specifies that no operating point calculation is to be performed.

`UIC` Specifies that no operating point calculation is to be performed, and that the specified initial condition (from `.IC` lines) should be used in its place. Unspecified values are set to zero. Finally, the `.IC` capability can only set voltage values, not current values.

`schedule(<time>, <maximum time step>, ...)`

Specifies a schedule for maximum allowed time steps. The list of arguments,  $t_0, \Delta t_0, t_1, \Delta t_1$ , *etc.* implies that a maximum time step of  $\Delta t_0$  will be used while the simulation time is greater than or equal to  $t_0$  and less than  $t_1$ . A maximum time step of  $\Delta t_1$  will be used when the simulation time is greater or equal to  $t_1$  and less than  $t_2$ . This sequence will continue for all pairs of  $t_i, \Delta t_i$  that are given in the `{schedule()}`. If  $\Delta t$  is zero or negative, then no maximum time step is enforced (other than hardware limits of the host computer).

---

**Comments**

The transient analysis calculates the circuit's response over an interval of time beginning with `TIME=0` and finishing at `<final time value>`. Use a `.PRINT` (print) statement to get the results of the transient analysis.

Before calculating the transient response **Xyce** computes a bias point for the circuit that is different from the regular bias point. This is necessary because at the start of a transient analysis, the independent sources can have different values than their DC values. Specifying `NOOP` on the `.TRAN` line causes **Xyce** to begin the transient analysis without performing the usual bias point calculation.

The time integration algorithms within **Xyce** use adaptive time-stepping methods that adjust the time-step size according to the activity in the analysis. The default ceiling for the internal time step is  $(\text{<final time value>}-\text{<start time value>})/10$ . This default ceiling value is automatically adjusted if breakpoints are present, to ensure that there are always at least 10 time steps between breakpoints. If the user specifies a ceiling value, however, it overrides any internally generated ceiling values.

**Xyce** is not strictly compatible with SPICE in its use of the values on the `.TRAN` line. In SPICE, the first number on the `.TRAN` line specifies the printing interval. In **Xyce**, the first number is the `<initial step value>`, which is used in determining the initial step size. The actual initial step size is chosen to be the smallest of three quantities: the `<initial step value>`, the `<step ceiling value>`, or 1/200th of the time until the next breakpoint.

The third argument to `.TRAN` simply determines the earliest time for which results are to be output. Simulation of the circuit always begins at `TIME=0` irrespective of the setting of `<start time value>`.

## 2.1.27 Miscellaneous Commands

### \* (Comment)

A netlist comment line. Whitespace at the beginning of a line is also interpreted as a comment.

### ; (In-line Comment)

Add a netlist in-line comment.

### + (Line Continuation)

Continue the text of the previous line.

## 2.2 Expressions

**Xyce** supports use of mathematical expressions in several contexts:

- for the values of device instance and model parameters.
- in definition of parameters in `.PARAM` and `.GLOBAL_PARAM` statements.
- for output on `.PRINT` lines.

In all contexts where expressions are allowed, they should be enclosed in braces (`{}`). For netlist compatibility with other simulators they may be enclosed in single quotation marks instead (`'`), but these are simply replaced with braces at a very early stage of netlist parsing. It is recommended that the braces be used in netlists written specifically for **Xyce**.

The expression package in **Xyce** supports all standard arithmetic operators, trigonometric functions, a collection of arithmetic functions, and some functions to mimic the pulse, sine, exp, and sffm time-dependent functions in the independent current and voltage sources. These functions are listed in tables 2.18 and 2.19.

### Operators

Table 2.17: Operators

Class of Operator	Operator	Meaning
arithmetic	+	addition or string concatenation
	-	subtraction
	*	multiplication
	/	division
	**	exponentiation
logical <sup>1</sup>	~	unary NOT
		boolean OR
	^	boolean XOR
	&	boolean AND
relational	==	equality
	!=	non-equality
	>	greater-than
	>=	greater-than or equal
	<	less-than
	<=	less-than or equal

<sup>1</sup>Logical and relational operators are used only with the `IF()` function.

## Arithmetic Functions

Table 2.18: Arithmetic Functions

Function	Meaning	Explanation
<i>Arithmetic functions</i>		
ABS(x)	$ x $	absolute value of $x$
AGAUSS( $\mu, \alpha, n$ )	$\mu - \alpha < result < \mu + \alpha$	<p>Random number sampled from normal distribution with mean <math>\mu</math> and standard deviation <math>\alpha/n</math></p> <p>The number returned will differ from the mean by at most <math>\alpha</math></p> <p>A deviation <math>\alpha</math> will be <math>n</math> standard deviations from the mean.</p>
GAUSS( $\mu, \alpha, n$ )	$\mu * (1 - \alpha) < result < \mu * (1 + \alpha)$	<p>Random number sampled from normal distribution with mean <math>\mu</math> and standard deviation <math>(\alpha * \mu)/n</math></p> <p>The number returned will differ from the mean by at most <math>\alpha * \mu</math></p> <p>A deviation <math>\alpha * \mu</math> will be <math>n</math> standard deviations from the mean.</p>
DDT(x)	$\frac{d}{dt}x(t)$	time derivative of $x$
DDX(f(x),x)	$\frac{\partial}{\partial x}f(x)$	partial derivative of $f(x)$ with respect to $x$
IF(t,x,y)	$x$ if $t$ is true, $y$ otherwise	$t$ is an expression using the relational operators in Table 2.17
INT(x)	$\text{sgn}(x) \lfloor  x  \rfloor$	integer part of the real variable $x$
LIMIT(x,y,z)	$y$ if $x < y$ $x$ if $y < x < z$ $z$ if $x > z$	$x$ limited to range $y$ to $z$
M(x)	$ x $	absolute value of $x$
MIN(x,y)	$\min(x, y)$	minimum of $x$ and $y$
MAX(x,y)	$\max(x, y)$	maximum of $x$ and $y$
PWR(x,y)	$x^y$	$x$ raised to $y$ power
POW(x,y)	$x^y$	$x$ raised to $y$ power
PWRS(x,y)	$x^y$ if $x > 0$ 0 if $x = 0$ $-(-x)^y$ if $x < 0$	sign corrected $x$ raised to $y$ power

Table 2.18: Arithmetic Functions

Function	Meaning	Explanation
RAND()	$0 < result < 1$	random number between 0 and 1 sampled from a uniform distribution
SDT(x)	$\int x(t)dt$	time integral of $x$
SGN(x)	+1 if $x > 0$ 0 if $x = 0$ -1 if $x < 0$	sign value of $x$
SIGN(x,y)	$\text{sgn}(y) x $	sign of $y$ times absolute value of $x$
STP(x)	1 if $x > 0$ 0 otherwise	step function
SQRT(x)	$\sqrt{x}$	square root of $x$
TABLE(x,y,z,*)	$f(x)$ where $f(y) = z$	piecewise linear interpolation, multiple $(y,z)$ pairs can be specified
URAMP(x)	$x$ if $x > 0$ 0 otherwise	ramp function
<b><i>Exponential, logarithmic, and trigonometric functions</i></b>		
ACOS(x)	$\arccos(x)$	result in radians
ACOSH(x)	$\cosh^{-1}(x)$	hyperbolic arccosine of $x$
ARCTAN(x)	$\arctan(x)$	result in radians
ASIN(x)	$\arcsin(x)$	result in radians
ASINH(x)	$\sinh^{-1}(x)$	hyperbolic arcsine of $x$
ATAN(x)	$\arctan(x)$	result in radians
ATANH(x)	$\tanh^{-1}(x)$	hyperbolic arctangent of $x$
ATAN2(x,y)	$\arctan(x/y)$	result in radians
COS(x)	$\cos(x)$	$x$ in radians
COSH(x)	$\cosh(x)$	hyperbolic cosine of $x$
EXP(x)	$e^x$	$e$ to the $x$ power
LN(x)	$\ln(x)$	log base $e$
LOG(x)	$\log(x)$	log base 10
LOG10(x)	$\log(x)$	log base 10
SIN(x)	$\sin(x)$	$x$ in radians
SINH(x)	$\sinh(x)$	hyperbolic sine of $x$
TAN(x)	$\tan(x)$	$x$ in radians
TANH(x)	$\tanh(x)$	hyperbolic tangent of $x$

## Spice Compatible Functions

Table 2.19: SPICE Compatibility Functions

Function	Explanation
SPICE_EXP(V1,V2,TD1,TAU1,TD2,TAU2)	SPICE style transient exponential V1 = initial value V2 = pulsed value TD1 = rise delay time TAU1 = rise time constant TD2 = fall delay time TAU2 = fall time constant
SPICE_PULSE(V1,V2,TD,TR,TF,PW,PER)	SPICE style transient pulse V1 = initial value V2 = pulsed value TD = delay TR = rise time TF = fall time PW = pulse width PER = period
SPICE_SFFM(V0,VA,FC,MDI,FS)	SPICE style transient single frequency FM V0 = offset VA = amplitude FC = carrier frequency MDI = modulation index FS = signal frequency
SPICE_SIN(V0,VA,FREQ,TD,THETA)	SPICE style transient sine wave V0 = offset VA = amplitude FREQ = frequency (hz) TD = delay THETA = damping factor

Information about the restrictions on expressions in specific contexts is given in the subsections that follow.

<sup>1</sup>The random number functions RAND, GAUSS, and AGAUSS return a unique number per use in an expression, but once evaluated this number is constant for the entire run, even across .STEP iterations.

## 2.2.1 Expressions in .PARAM or .GLOBAL\_PARAM Statements

Expressions used in .PARAM statements are the most highly constrained. They must evaluate to a constant at the beginning of a run, and therefore must involve only numerical constants and other previously defined .PARAMS. The value of the parameter will be computed when the netlist is parsed, and will replace the name wherever it is used.

**Example:**        .PARAM SQUARES=5.0

**Example:**        .PARAM SHEETRES=25

**Example:**        .PARAM RESISTANCE={SQUARES\*SHEETRES}

Global parameters are somewhat less constrained. These parameters are allowed to depend on parameters defined in .PARAMS or .GLOBAL\_PARAMS statements, and may contain special variables such as TIME or TEMP. They may not contain any references to solution variables or lead currents.

**Example:**        .PARAM dTdt=.01

**Example:**        .GLOBAL\_PARAM Temperature={27+dTdt\*TIME}

## 2.2.2 Expressions in .PRINT Lines

Expressions on .PRINT lines may contain references to parameters defined in either .PARAM or .GLOBAL\_PARAM statements, device parameters using the syntax <device name>:<parameter name>, and may also contain solution variables.

```
*example with .print expressions
.PARAM RES=50
R1 1 0 {RES}
V1 1 0 sin(0 5 100khz)
.tran 1u 1m
*Print power dissipated through resistor,
*and actual resistance used in the R1
*device
.print tran {V(1)*V(1)/RES} {R1:R}
.end
```

## 2.2.3 Expressions for Device Instance and Model Parameters

Expressions of constants and .PARAM parameters may be used for the values of any device parameters in instance and model lines.

Except in very specific devices, expressions used for device parameter values must evaluate to a time-independent constant, and must not contain dependence on solution variables such as nodal voltages or currents. In these cases, .GLOBAL\_PARAM parameters may also be used as long as they are not time-dependent.

```
*example of use of expressions for device parameters
.PARAM RES=50
.GLOBAL_PARAM theSaturationCurrent=1.5e-14
R1 1 0 {RES}
V1 1 0 sin(0 5 100khz)
D1 1 0 DMODEL
.MODEL D DMODEL IS=theSaturationCurrent

.step theSaturationCurrent 1e-14 5e-14 1e-14
```

Some parameters of specific devices are exceptions to the general rule. These parameters have no restrictions and may depend on any parameters, time, or solution variables in the netlist:

- The V or I instance parameters of the B source.
- The CONTROL instance parameter of the switch (S device).
- The C (capacitance) instance parameter for the capacitor.
- The coupling coefficient instance parameter for the *LINEAR* mutual inductor (K device with no model card specified)

These specific instance parameters may be time-dependent (i.e. they may reference the TIME special variable, but may not depend on any solution variables:

- The TEMP instance parameter of all devices.
- The L (inductance) parameter of the inductor.
- The R (resistance) parameter of the resistor.
- The R, RESISTIVITY, DENSITY, HEATCAPACITY and THERMAL\_HEATCAPACITY parameters of the thermal resistor (resistor level 2).

## 2.2.4 Special PSpice POLY expression

The POLY keyword, available in the E,F,G, H and B dependent sources, is provided to simplify migration of netlists from PSpice to **Xyce**. POLY provides a compact method of specifying polynomial expressions in which the variables in the polynomial are specified followed by an ordered list of polynomial coefficients. All expressions specified with POLY are ultimately translated by **Xyce** into an equivalent, straightforward polynomial expression in a B source. Since a straightforward polynomial expression can be easier to read, there is no real benefit to using POLY except to support legacy netlists imported from PSpice.

There are three different syntax forms for POLY, which can be a source of confusion. The E and G sources (voltage-dependent voltage or current sources) use one form, the F and H sources (current-dependent voltage or current sources) use a second form, and the B source (general nonlinear source) a third form. During input processing, any of the E,F,G or H sources that use nonlinear expressions are first converted into an equivalent B source, and then any B sources that use the POLY shorthand are further converted into standard polynomial expressions. This section describes how the compact form will be translated into the final form that is used internally.

All three formats of POLY express the same three components: a number of variables involved in the expression ( $N$ , the number in parentheses after the POLY keyword), the variables themselves, and an ordered list of coefficients for the polynomial terms. Where they differ is in how the variables are expressed.

### Voltage-controlled sources

The E and G sources are both voltage-controlled, and so their POLY format requires specification of two nodes for each voltage on which the source depends, i.e. the positive and negative nodes from which a voltage drop is computed. There must therefore be twice as many nodes as the number of variables specified in parentheses after the POLY keyword:

```
Epoly 1 2 POLY(3) n1p n1m n2p n2m n3p n3m ...
```

In this example, the voltage between nodes 1 and 2 is determined by a polynomial whose variables are  $V(n1p,n1m)$ ,  $V(n2p,n2m)$ ,  $V(n3p,n3m)$ . Not shown in this example are the polynomial coefficients, which will be described later.

### Current-controlled sources

The F and H sources are both current-controlled, and so their POLY format requires specification of one voltage source name for each current on which the source depends. There must therefore be exactly as many nodes as the number of variables specified in parentheses after the POLY keyword:

```
Fpoly 1 2 POLY(3) V1 V2 V3 ...
```

In this example, the voltage between nodes 1 and 2 is determined by a polynomial whose variables are  $I(V1)$ ,  $I(V2)$ , and  $I(V3)$ . Not shown in this example are the polynomial coefficients, which will be described later.

## B sources

Finally, the most general form of POLY is that used in the general nonlinear dependent source, the B source. In this variant, each specific variable must be named explicitly (i.e. not simply by node name or by voltage source name), because currents and voltages may be mixed as needed.

```
Bpoly 1 2 V={POLY(3) I(V1) V(2,3) V(3) ...}
```

```
Bpoly2 1 2 I={POLY(3) I(V1) V(2,3) V(3) ...}
```

In these examples, the source between nodes 1 and 2 is determined by a polynomial whose variables are I(V1), V(2,3), and V(3). In the first example, the polynomial value determines the voltage between nodes 1 and 2, and in the second the current.

The E, F, G and H formats are all converted internally in a first step to the B format. Thus the following pairs of sources are exactly equivalent:

```
Epoly 1 2 POLY(3) n1p n1m n2p n2m n3p n3m ...
```

```
BEpoly 1 2 V={POLY(3) V(n1p,n1m) V(n2p,n2m) V(n3p,n3m) ...}
```

```
Fpoly 1 2 POLY(3) V1 V2 V3 ...
```

```
BFpoly 1 2 V={POLY(3) I(V1) I(V2) I(V3) ...}
```

After conversion to the B source form, the POLY form is finally converted to a normal expression using the coefficients and variables given.

Coefficients are given in a standard order, and the polynomial is built up by terms until the list of coefficients is exhausted. The first coefficient is the constant term of the polynomial, followed by the coefficients of linear terms, then bi-linear, and so on. For example:

```
Epoly 1 2 POLY(3) n1p n1m n2p n2m n3p n3m 1 .5 .5 .5
```

In this example, the constant term is 1.0, and the coefficients of the three terms linear in the input variables are 0.5. Thus, this E source is precisely equivalent to the general B source:

```
BEstandard 1 2 V={1.0 + .5*V(n1p,n1m) + .5*V(n2p,n2m) + .5*V(n3p,n3m)}
```

The standard ordering for coefficients is:

POLY(N)  $X_1 \dots X_N C_0 C_1 \dots C_N C_{11} \dots C_{1N} C_{21} \dots C_{N1} \dots C_{NN} C_{1^2 1} \dots C_{1^2 N} \dots$

with the polynomial then being:

$$Value = C_0 + \sum_{j=1}^N C_j X_j + \sum_{i=1}^N \sum_{j=1}^N C_{ij} X_i X_j + \sum_{i=1}^N \sum_{j=1}^N C_{i^2 j} X_i^2 X_j + \dots$$

Here we have used the general form  $X_i$  for the  $i^{th}$  variable, which may be either a current or

voltage variable in the general case.

It should be reiterated that the POLY format is provided primarily for support of legacy PSpice netlists that use the feature, and that its compactness may be a disadvantage in readability of the netlist and may be more prone to usage error. **Xyce** users are therefore advised that use of the more straightforward expression format in the B source may be more appropriate when crafting original netlists for use in **Xyce**. Since **Xyce** converts POLY format expressions to the simpler format internally, there is no performance benefit to use of POLY.

## 2.3 Devices

**Xyce** supports many devices, with an emphasis on analog devices, including sources, subcircuits and behavioral models. This section serves as a reference for the devices supported by **Xyce**. Each device is described separately and includes the following information, if applicable:

- a description and an example of the correct netlist syntax.
- the matching model types and their description.
- the matching list of model parameters and associated descriptions.
- the corresponding characteristic equations for the model (as required).
- references to publications on which the model is based.

User-defined models may be implemented using the `.MODEL` (model definition) statement, and macromodels can be created as subcircuits using the `.SUBCKT` (subcircuit) statement.

Please note that the characteristic equations are provided to give a general representation of the device behavior. The actual **Xyce** implementation of the device may be slightly different in order to improve, for example, the robustness of the device.

Table 2.20 gives a summary of the device types and the form of their netlist formats. Each of these is described below in detail.

Table 2.20: Analog Device Quick Reference.

Device Type	Letter	Typical Netlist Format
Nonlinear Dependent Source (B Source)	B	B<name> <+ node> <- node> + <I or V>={<expression>}
Capacitor	C	C<name> <+ node> <- node> [model name] <value> + [IC=<initial value>]
Diode	D	D<name> <anode node> <cathode node> + <model name> [area value]
Voltage Controlled Voltage Source	E	E<name> <+ node> <- node> <+ controlling node> + <- controlling node> <gain>
Current Controlled Current Source	F	F<name> <+ node> <- node> + <controlling V device name> <gain>
Voltage Controlled Current Source	G	G<name> <+ node> <- node> <+ controlling node> + <- controlling node> <transconductance>
Current Controlled Voltage Source	H	H<name> <+ node> <- node> + <controlling V device name> <gain>
Independent Current Source	I	I<name> <+ node> <- node> [[DC] <value>] + [AC [magnitude value] [phase value] ] ] + [transient specification]

Table 2.20: Analog Device Quick Reference.

Device Type	Letter	Typical Netlist Format
Mutual Inductor	K	K<name> <inductor 1> [<ind. n>*] + <linear coupling or model>
Inductor	L	L<name> <+ node> <- node> [model name] <value> + [IC=<initial value>]
JFET	J	J<name> <drain node> <gate node> <source node> + <model name> [area value]
MOSFET	M	M<name> <drain node> <gate node> <source node> + <bulk/substrate node> [SOI node(s)] + <model name> [common model parameter]*
Lossy Transmission Line (LTRA)	O	O<name> <A port (+) node> <A port (-) node> + <B port (+) node> <B port (-) node> + <model name>
Bipolar Junction Transistor (BJT)	Q	Q<name> <collector node> <base node> + <emitter node> [substrate node] + <model name> [area value]
Resistor	R	R<name> <+ node> <- node> [model name] <value> + [L=<length>] [W=<width>]
Voltage Controlled Switch	S	S<name> <+ switch node> <- switch node> + <+ controlling node> <- controlling node> + <model name>
Transmission Line	T	T<name> <A port + node> <A port - node> + <B port + node> <B port - node> + <ideal specification>
Digital Devices	U	U<name> <type> <digital power node> + <digital ground node> [node]* <model name>
Independent Voltage Source	V	V<name> <+ node> <- node> [[DC] <value>] + [AC [magnitude value] [phase value] ] ] + [transient specification]
Subcircuit	X	X<name> [node]* <subcircuit name> + [PARAMS:[<name>=<value>]*]
Current Controlled Switch	W	W<name> <+ switch node> <- switch node> + <controlling V device name> <model name>
Digital Devices, Y Type (deprecated)	Y<type>	Y<type> <name> [node]* <model name>
PDE Devices	YPDE	YPDE <name> [node]* <model name>
Accelerated masses	YACC	YACC <name> <acceleration> <velocity> <position> + [x0=<initial position>] [v0=<initial velocity>]
MESFET	Z	Z<name> <drain node> <gate node> <source node> + <model name> [area value]

## 2.3.1 Voltage Nodes

Devices in a netlist are connected between *nodes*, and all device types in **Xyce** require at least two nodes on each instance line. Node names can consist of any printable characters *except* white space (space, tab, newline), parentheses (“(” or “)” ), braces (“{” or “}”), commas, or the equal sign.

Except for global nodes (below), voltage node names appearing in a subcircuit that are not listed in the subcircuit’s argument list are accessible only to that subcircuit; devices outside the subcircuit cannot connect to local nodes.

### Global nodes

A special syntax is used to designate certain nodes as *global* nodes. Any node whose name starts with the two characters “\$G” is a global node, and such nodes are available to be used in any subcircuit. A typical usage of such global nodes is to define a VDD or VSS signal that all subcircuits need to be able to access, but without having to provide VSS and VDD input nodes to every subcircuit. In this case, a global \$GVDD node would be use for the VDD signal.

The node named 0 is a special global node. Node 0 is always ground, and is accessible to all levels of a hierarchical netlist.

### Subcircuit Nodes

Hierarchical netlists may be created using .SUBCKT [2.1.25] to define common subcircuit types, and X [2.3.26] lines to create instances of those subcircuits. There are two types of nodes associated with such subcircuits, *interface* nodes and *internal* nodes.

Interface nodes are the nodes named on the .SUBCKT line. These are effectively local aliases internal to the subcircuit definition for the node names used on the X instance lines. Internal nodes are nodes inside the subcircuit definition that are strictly local to that subcircuit. Inside a subcircuit, these node names may be used without restriction in device instance lines and expressions on B source lines.

There are some circumstances when it is desirable to access internal nodes of a subcircuit from outside that subcircuit. **Xyce** provides a syntax that allows this to be done in *some* contexts. The primary context in which this is supported is on .PRINT lines, to allow the user to print out signals that are usually local to a subcircuit.

The syntax used by **Xyce** to refer to nodes within a subcircuit is to prefix the name of the node with the full path of subcircuit instances in which the node is contained, with colons (:) separating the instance names. So, to reference a node “A” that is inside a subcircuit instance called “Xnot1” inside another subcircuit instance called “Xmain”, one would refer to “Xmain:Xnot1:A”

The same syntax works on .PRINT lines even if the subcircuit node is one of the interface nodes on the .SUBCKT line, but those nodes can also be accessed by using the names of the nodes at the higher level of circuit hierarchy that are used on its instance line.

```

* Netlist file demonstrating subcircuit node access
V1 1 0 1
X1 1 2 demosubc
X2 2 0 demosubc
.subckt demosubc A B
R1 A C 1
R2 C B 1
.ends


.dc V1 1 5 1

*V(X1:A) and V(1) are the same signal.
*V(X1:C) is the internal C node of the X1 instance
*V(X2:C) is the internal C node of the X2 instance
*V(X1:B), V(X2:A) and V(2) are the same signal
.print DC V(X1:C) V(X2:C) V(X1:A) V(1) V(X1:B) V(X2:A) V(2)
.end

```

Subcircuit nodes may also be accessed from outside of the subcircuit in B source voltage or current expressions, though this usage violates the strict hierarchy of the netlist. The one difference between this usage and .PRINT usage is that it is not possible to use the subcircuit node syntax to access interface nodes. These must be accessed using the node names being used on the instance line, as in the “V(1)” example in the netlist fragment above.

## 2.3.2 Capacitor

<b>Symbol</b>	
<b>Instance Form</b>	C<device name> <(+) node> <(-) node> [model name] [value] + [device parameters]
<b>Model Form</b>	.MODEL <model name> C [model parameters] .MODEL <model name> CAP [model parameters]
<b>Examples</b>	CM12 2 4 5.288e-13 CLOAD 1 0 4.540pF IC=1.5V CFEEDBACK 2 0 CMOD 1.0pF CAGED 2 3 4.0uF D=0.0233 AGE=86200
<b>Parameters and Options</b>	<p>device name The name of the device.</p> <p>(+) node (-) node Polarity definition for a positive voltage across the capacitor. The first node is defined as positive. Therefore, the voltage across the component is the first node voltage minus the second node voltage.</p> <p>model name If model name is omitted, then value is the capacitance in farads. If [model name] is given then the value is determined from the model parameters; see the capacitor value formula below.</p> <p>value Positional specification of device parameter C (capacitance). Alternately, this can be specified as a parameter, C=&lt;value&gt;, or in the (optional) model.</p> <p>device parameters Parameters listed in Table 2.21 may be provided as space separated &lt;parameter&gt;=&lt;value&gt; specifications as needed. Any number of parameters may be specified.</p> <p>model parameters Parameters listed in Table 2.22 may be provided as space separated &lt;parameter&gt;=&lt;value&gt; specifications as needed. Any number of parameters may be specified.</p>

## Comments

Positive current flows through the capacitor from the (+) node to the (-) node. In general, capacitors should have a positive capacitance value (<value> property). In all cases, the capacitance must not be zero.

However, cases exist when a negative capacitance value may be used. This occurs most often in filter designs that analyze an RLC circuit equivalent to a real circuit. When transforming from the real to the RLC equivalent, the result may contain a negative capacitance value.

In a transient run, negative capacitance values may cause the simulation to fail due to instabilities they cause in the time integration algorithms.

For compatibility with PSpice, either C or CAP can be used in a .MODEL statement for a capacitor.

## Device Parameters

Table 2.21: Capacitor Device Instance Parameters

Parameter	Description	Units	Default
AGE	Age of capacitor	hour	0
C	Capacitance	F	1e-06
D	Age degradation coefficient	—	0.0233
IC	Initial voltage drop across device	V	0
L	Semiconductor capacitor width	m	1
TC1	Linear Temperature Coefficient	$^{\circ}\text{C}^{-1}$	0
TC2	Quadratic Temperature Coefficient	$^{\circ}\text{C}^{-2}$	0
TEMP	Device temperature	$^{\circ}\text{C}$	Ambient Temperature
W	Semiconductor capacitor length	m	1e-06

In addition to the parameters shown in the table, the capacitor supports a vector parameter for the temperature correction coefficients. TC1=<linear coefficient> and TC2=<quadratic coefficient> may therefore be specified compactly as TC=<linear coefficient>,<quadratic coefficient>.

## Model Parameters

Table 2.22: Capacitor Device Model Parameters

Parameter	Description	Units	Default
CJ	Junction bottom capacitance	F/m <sup>2</sup>	0
CJSW	Junction sidewall capacitance	F/m	0

Table 2.22: Capacitor Device Model Parameters

Parameter	Description	Units	Default
DEFW	Default device width	m	1e-06
NARROW	Narrowing due to side etching	m	0
TC1	Linear temperature coefficient	°C <sup>-1</sup>	0
TC2	Quadratic temperature coefficient	°C <sup>-2</sup>	0
TNOM	Nominal device temperature	°C	Ambient Tempera- ture

## Capacitor Equations

### Capacitance Value Formula

If [model name] is specified, then the capacitance is given by:

$$C \cdot (1 + \text{TC1} \cdot (T - T_0) + \text{TC2} \cdot (T - T_0)^2)$$

where  $C$  is the base capacitance specified on the device line and is normally positive (though it can be negative, but not zero).  $T_0$  is the nominal temperature (set using **TNOM** option).

### Age-aware Formula

If **AGE** is given, then the capacitance is:

$$C[1 - D \log(\text{AGE})]$$

### Semiconductor Formula

If [model name] and **L** and **W** are given, then the capacitance is:

$$CJ(L - \text{NARROW})(W - \text{NARROW}) + 2 \cdot CJSW(L - W + 2 \cdot \text{NARROW})$$

## 2.3.3 Inductor

**Symbol** 

---

**Instance Form** L<name> <(+) node> <(-) node> [model] <value> [device parameters]

---

**Model Form** .MODEL <model name> L [model parameters]  
.MODEL <model name> IND [model parameters]

---

**Examples** L1 1 5 3.718e-08  
LLOAD 3 6 4.540mH IC=2mA  
Lmodded 3 6 indmod 4.540mH  
.model indmod L (L=.5 TC1=0.010 TC2=0.0094)

---

**Parameters and Options**

(+) node  
(-) node  
Polarity definition for a positive voltage across the inductor. The first node is defined as positive. Therefore, the voltage across the component is the first node voltage minus the second node voltage.

initial value  
The initial current through the inductor during the bias point calculation.

---

**Comments** In general, inductors should have a positive inductance value. The inductance must not be zero.

However, cases exist when a negative value may be used. This occurs most often in filter designs that analyze an RLC circuit equivalent to a real circuit. When transforming from the real to the RLC equivalent, the result may contain a negative inductance value.

If a model name is given, the inductance is modified from the value given on the instance line by the parameters in the model card. See “Inductance Value Formula” below.

When an inductor is named in the list of coupled inductors in a mutual inductor device line (see page 107) , and that mutual inductor is of the nonlinear-core type, the <value> is interpreted as a number of turns rather than as an inductance in Henries.

For compatibility with PSpice, either L or IND can be used in a .MODEL statement for an inductor.

## Device Parameters

Table 2.23: Inductor Device Instance Parameters

Parameter	Description	Units	Default
IC	Initial current through device	A	0
L	Inductance	henry	0
TC1	Linear Temperature Coefficient	$^{\circ}\text{C}^{-1}$	0
TC2	Quadratic Temperature Coefficient	$^{\circ}\text{C}^{-2}$	0
TEMP	Device temperature	$^{\circ}\text{C}$	Ambient Temperature

## Model Parameters

Table 2.24: Inductor Device Model Parameters

Parameter	Description	Units	Default
IC	Initial current through device	A	0
L	Inductance Multiplier	—	1
TC1	First order temperature coeff.	$^{\circ}\text{C}^{-1}$	0
TC2	Second order temperature coeff.	$^{\circ}\text{C}^{-2}$	0
TNOM	Reference temperature	$^{\circ}\text{C}$	27

In addition to the parameters shown in the table, the inductor supports a vector parameter for the temperature correction coefficients. TC1=<linear coefficient> and TC2=<quadratic coefficient> may therefore be specified compactly as TC=<linear coefficient>,<quadratic coefficient>.

## Inductor Equations

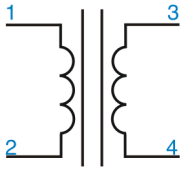
### Inductance Value Formula

If [model name] is specified, then the inductance is given by:

$$\mathbf{L}_{base} \cdot \mathbf{L} \cdot (1 + \mathbf{TC1} \cdot (T - T_0) + \mathbf{TC2} \cdot (T - T_0)^2)$$

where  $\mathbf{L}_{base}$  is the base inductance specified on the device line and is normally positive (though it can be negative, but not zero).  $\mathbf{L}$  is the inductance multiplier specified in the model card.  $T_0$  is the nominal temperature (set using TNOM option).

## 2.3.4 Mutual Inductors

Symbol	
Instance Form	<p>K&lt;name&gt; L&lt;inductor name&gt; [L&lt;inductor name&gt;*]  + &lt;coupling value&gt; [model name]</p>
Model Form	.MODEL <model name> CORE [model parameters]
Examples	<pre>KTUNED L3OUT L4IN .8 KTRNSFRM LPRIMARY LSECNDRY 1 KXFRM L1 L2 L3 L4 .98 KPOT_3C8</pre>
Parameters and Options	<p><b>inductor name</b>  Identifies the inductors to be coupled. The inductors are coupled and in the dot notation the dot is placed on the first node of each inductor. The polarity is determined by the order of the nodes in the L devices and not by the order of the inductors in the K statement.</p> <p><b>coupling value</b>  The coefficient of mutual coupling, which must be between <math>-1.0</math> and <math>1.0</math>.  This coefficient is defined by the equation</p> $\text{<coupling value>} = \frac{M_{ij}}{\sqrt{L_i L_j}}$ <p>where</p> <p><math>L_i</math> is the inductance of the <math>i</math>th named inductor in the K-line</p> <p><math>M_{ij}</math> is the mutual inductance between <math>L_i</math> and <math>L_j</math></p> <p>For transformers of normal geometry, use <math>1.0</math> as the value. Values less than <math>1.0</math> occur in air core transformers when the coils do not completely overlap.</p> <p><b>model name</b>  If model name is present, four things change:</p> <ul style="list-style-type: none"> <li>• The mutual coupling inductor becomes a nonlinear, magnetic core device.</li> <li>• The inductors become windings, so the number specifying inductance now specifies the number of turns.</li> <li>• The list of coupled inductors could be just one inductor.</li> <li>• A model statement is required to specify the model parameters.</li> </ul>

## Model Parameters

Table 2.25: Nonlinear Mutual Inductor Device Model Parameters

Parameter	Description	Units	Default
A	Thermal energy parameter	A/m	1000
ALPHA	Domain coupling parameter	–	5e-05
AREA	Mean magnetic cross-sectional area	cm <sup>2</sup>	0.1
BETAH	Modeling constant	–	0.0001
BETAM	Modeling constant	–	3.125e-05
BHSIUNITS	Flag to report B and H in SI units	–	0
C	Domain flexing parameter	–	0.2
CLIM	Value below which domain flexing parameter will be treated as zero.	–	0.005
CONSTDELVSCALING	Use constant scaling factor to smooth voltage difference over first inductor	V	false
DELVSCALING	Smoothing coefficient for voltage difference over first inductor	V	1000
FACTORMS	Flag to save state variables	–	0
GAP	Effective air gap	cm	0
INCLUDEMEQU	Flag to include the magnetics in the solution.	–	true
K	Domain anisotropy parameter	A/m	500
KIRR	Domain anisotropy parameter	A/m	500
LEVEL	for pspice compatibility – ignored	–	0
MEQNSCALING	M-equation scaling	–	1
MS	Saturation magnetization	A/m	1e+06
MVARSCALING	M-variable scaling.	–	1
OUTPUTSTATEVARS	Flag to save state variables	–	0
PACK	for pspice compatibility – ignored	–	0
PATH	Total mean magnetic path	cm	1
PZEROTOL	Tolerance for nonlinear zero crossing	–	0.1
REQNSCALING	R-equation scaling	–	1
RVARSCALING	R-variable scaling	–	1
TC1	First order temperature coeff.	–	0
TC2	Second order temperature coeff.	–	0
TNOM	Reference temperature	°C	27

Note that **Xyce**'s default value for the GAP parameter as zero. Some simulators will use non-zero values of the GAP as a default. When using netlists from other simulators in **Xyce**, ensure that the

default parameters are consistent.

### Special Notes

The coupling coefficient of the linear mutual inductor (i.e. a mutual inductor without a core model) is permitted to be a time- or solution variable-dependent expression. This is intended to allow simulation of electromechanical devices in which there might be moving coils that interact with fixed coils.

Nonlinear mutual inductors can output  $B(t)$  and  $H(t)$  variables so that one can plot  $B - H$  loops. On the `.print` line the  $B$  and  $H$  variables are accessible using the node output syntax as in `n( non-linear-inductor-name_b )` for  $B$  and `n( non-linear-inductor-name_h )` for  $H$ . A confusing aspect of this is that the non-linear inductor name is the *internal* name used by **Xyce**. For example, if the following non-linear mutual inductor is declared in a netlist:

```
Lp1      3 0 50
Lp2      0 6 50
Lp3      4 0 20
Lp4      0 5 10
```

```
ktrans1  Lp1 Lp2 Lp3 Lp4 1  trans_core
```

then the internal, **Xyce** name of the non-linear mutual inductor is `YMIN!KTRANS1` or `ymin!ktrans1` as the name is not case-sensitive. If the device `ktrans1` were declared within a subcircuit called `sub1` then the full name would be `ymin!sub1!ktrans1`. The reason for this is that both the linear and non-linear mutual inductors are devices that are collections of other devices, inductors in this case. Rather than use one of the few remaining single characters left to signify a new device, **Xyce** uses `Y` devices as an indicator of an extended device set, where the characters after the `Y` denote the device type and then the device name. Here, `ymin` means a `min` device which is a *mutual-inductor, non-linear* device. Thus, to print the  $B$  or  $H$  variable of the non-linear mutual inductor called `ktrans1` one would use `n(ymin!ktrans1_b)` and `n(ymin!ktrans1_h)` respectively for a `.print` line that looks like this:

```
.print tran n(ymin!ktrans1_b) n(ymin!ktrans1_h)
```

Note that while MKS units are used internally in **Xyce**,  $B$  and  $H$  are output by default in the SI units of Gauss for  $B$  and Oersted for  $H$ . To convert  $B$  to units of Tesla divide **Xyce**'s output by 10,000. To convert  $H$  to units of  $A/m$  divide **Xyce**'s output by  $4\pi/1000$ . Additionally, one can set the `.model CORE` parameter `BHSIUNITS` to 1 to force  $B$  and  $H$  to be output in MKS units.

The branch current through any of the inductors making up the mutual inductor can be included on the `.print` line by using the nodal variable syntax `n( non-linear-inductor-name_inductor-name_branch )`.

So continuing with the sample mutual inductor listed earlier, if one wanted the branch current through the sub-inductor `Lp1` one would could use this `.print` statement:

```
.print tran n(ymin!ktrans1_Lp1_branch)
```

Additionally, one can access the  $B$  and  $H$  data via the `.model CORE` line. On the nonlinear mutual inductor's `.model` line set the option `OUTPUTSTATEVARS=1`. This will cause **Xyce** to create a unique file for each nonlinear mutual inductor that uses this `.model` line with a name of the form `Inductor_device_name`. There are five columns of data in this file: time ( $t$ ), magnetic moment ( $M$ ), total current flux ( $R$ ), flux density ( $B$ ) and magnetic field strength ( $H$ ). As with data output on the `.print` line, SI units are used such that  $B$  is output with units of Gauss and  $H$  in Oersted. As mentioned earlier, setting the model flag `BHSIUNITS` to 1 causes the output of  $B$  and  $H$  uses MKS units of Tesla and  $A/m$  respectively.

### Mutual Inductor Equations

The voltage to current relationship for a set of linearly coupled inductors is:

$$V_i = \sum_{j=1}^N c_{ij} \sqrt{L_i L_j} \frac{dI_j}{dt} \quad (2.1)$$

Here,  $V_i$  is the voltage drop across the  $i$ th inductor in the coupled set. The coupling coefficient between a pair of inductors is  $c_{ij}$  with a value typically near unity and  $L$  is the inductance of a given inductor which has units of *Henry's* ( $1 \text{ Henry} = 1H = \text{Volt} \cdot \text{s}/\text{Amp}$ )

For nonlinearly coupled inductors, the above equation is expanded to the form:

$$V_i = \left[ 1 + \left( 1 - \frac{\ell_g}{\ell_t} \right) P(M, I_1 \dots I_N) \right] \sum_{j=1}^N L_{oij} \frac{dI_j}{dt} \quad (2.2)$$

This is similar in form to the linearly coupled inductor equation. However, the coupling has become more complicated as it now depends on the magnetic moment created by the current flow,  $M$ . Additionally, there are geometric factors,  $\ell_g$  and  $\ell_t$  which are the effective air gap and total mean magnetic path for the coupled inductors. The matrix of terms,  $L_{oij}$  is defined as

$$L_{oij} = \frac{\mu_0 A_c N_i N_j}{\ell_t} \quad (2.3)$$

and it represents the physical coupling between inductors  $i$  and  $j$ . In this expression,  $N_i$  is the number of windings around the core of inductor  $i$ ,  $\mu_0$  is the magnetic permeability of free space which has units of Henries per meter and a value of  $4\pi \times 10^{-7}$  and  $A_c$  is the mean magnetic cross-sectional area.

The magnetic moment,  $M$  is defined by:

$$\frac{dM}{dt} = \frac{1}{\ell_t} P \sum_{i=1}^N N_i \frac{dI_i}{dt} \quad (2.4)$$

and the function  $P$  is defined as:

$$P = \frac{cM'_{an} + (1 - c)M'_{irr}}{1 + \left(\frac{\ell_g}{\ell_t} - \alpha\right) cM'_{an} + \frac{\ell_g}{\ell_t}(1 - c)M'_{irr}} \quad (2.5)$$

If  $c < \text{CLIM}$ , then  $c$  is treated as zero in the above equation and **Xyce** simplifies the formulation. In this case, the magnetic-moment equation will not be needed and it will be dropped from the formulation. One can control this behavior by modifying the value of CLIM.

The remaining functions are:

$$M'_{an} = \frac{M_s A}{(A + |H_e|)^2} \quad (2.6)$$

$$H_e = H + \alpha M \quad (2.7)$$

$$H = H_{app} - \frac{\ell_g}{\ell_t} M \quad (2.8)$$

$$H_{app} = \frac{1}{\ell_t} \sum_{i=1}^N N_i I_i \quad (2.9)$$

$$M'_{irr} = \frac{\Delta M \text{sgn}(q) + |\Delta M|}{2(K_{irr} - \alpha|\Delta M|)} \quad (2.10)$$

$$\Delta M = M_{an} - M \quad (2.11)$$

$$M_{an} = \frac{M_s H_e}{A + |H_e|} \quad (2.12)$$

$$q = \text{DELVSCALING} \Delta V \quad (2.13)$$

**Xyce** dynamically modifies DELVSCALING to be  $1000 / \text{Maximum Voltage Drop over the first inductor}$ . This typically produces accurate results for both low voltage and high voltage applications. However, it is possible to use a fixed scaling by setting the model parameter CONSTDELVSCALING to true and then setting DELVSCALING to the desired scaling value.

In **Xyce's** formulation, we define  $R$  as:

$$R = \frac{dH_{app}}{dt} = \frac{1}{\ell_t} \sum_{i=1}^N N_i \frac{dI_i}{dt} \quad (2.14)$$

This simplifies the  $M$  equation to:

$$\frac{dM}{dt} = PR \quad (2.15)$$


**Xyce** then solves for the additional variables  $M$  and  $R$  when modeling a nonlinear mutual inductor device.

To calculate  $B$ - $H$  loops,  $H$  is used as defined above and  $B$  is a derived quantity calculated by:

$$B = \mu_0 (H + M) \quad (2.16)$$

$$= \mu_0 \left[ H_{app} + \left( 1 - \frac{\ell_g}{\ell_t} \right) M \right] \quad (2.17)$$

## 2.3.5 Resistor

<b>Symbol</b>	
<b>Instance Form</b>	R<name> <(+) node> <(-) node> [model name] [value] [device parameters]
<b>Model Form</b>	<pre>.MODEL &lt;model name&gt; R [model parameters] .MODEL &lt;model name&gt; RES [model parameters]</pre>
<b>Examples</b>	<pre>R1 1 2 2K TEMP=27 RLOAD 3 6 RTCMOD 4.540 TEMP=85 .MODEL RTCMOD R (TC1=.01 TC2=-.001) RSEMICON 2 0 RMOD L=1000u W=1u .MODEL RMOD R (RSH=1)</pre>
<b>Parameters and Options</b>	<p>(+) node (-) node Polarity definition for a positive voltage across the resistor. The first node is defined as positive. Therefore, the voltage across the component is the first node voltage minus the second node voltage. Positive current flows from the positive node (first node) to the negative node (second node).</p> <p>model name If [model name] is omitted, then [value] is the resistance in Ohms. If [model name] is given then the resistance is determined from the model parameters; see the resistance value formula below.</p> <p>value Positional specification of device parameter R (resistance). Alternately, this can be specified as a parameter, R=&lt;value&gt;, or in the (optional) model.</p> <p>device parameters Parameters listed in Table 2.26 may be provided as space separated &lt;parameter&gt;=&lt;value&gt; specifications as needed. Any number of parameters may be specified.</p>
<b>Comments</b>	<p>Resistors must have a positive (nonzero) resistance value (R).</p> <p>For compatibility with PSpice, either R or RES can be used in a .MODEL statement for a resistor.</p>

## Device Parameters

Table 2.26: Resistor Device Instance Parameters

Parameter	Description	Units	Default
DTEMP	Device Temperature – For compatibility only. Parameter is NOT used	°C	0
L	Length	m	0
R	Resistance	Ω	1000
TC1	Linear Temperature Coefficient	°C <sup>-1</sup>	0
TC2	Quadratic Temperature Coefficient	°C <sup>-2</sup>	0
TEMP	Device temperature	°C	Ambient Temperature
W	Width	m	0

In addition to the parameters shown in the table, the resistor supports a vector parameter for the temperature correction coefficients. TC1=<linear coefficient> and TC2=<quadratic coefficient> may therefore be specified compactly as TC=<linear coefficient>,<quadratic coefficient>.

## Model Parameters

Table 2.27: Resistor Device Model Parameters

Parameter	Description	Units	Default
DEFW	Default Instance Width	m	1e-05
NARROW	Narrowing due to side etching	m	0
RSH	Sheet Resistance	Ω	0
TC1	Linear Temperature Coefficient	°C <sup>-1</sup>	0
TC2	Quadratic Temperature Coefficient	°C <sup>-2</sup>	0
TNOM	Parameter Measurement Temperature	°C	Ambient Temperature

## Resistor Equations

### Resistance Value Formula

If [model name] is included, then the resistance is:

$$R \cdot (1 + TC1 \cdot (T - T_0) + TC2 \cdot (T - T_0)^2)$$

If **L** and **W** are given, the resistance is:

$$R_{SH} \frac{[L - NARROW]}{[W - NARROW]}$$

## Thermal (level=2) Resistor

**Xyce** supports a thermal resistor model, which is associated with level=2.

### Thermal Resistor Instance Parameters

Table 2.28: Resistor Device Instance Parameters

Parameter	Description	Units	Default
A	Area of conductor	m <sup>2</sup>	0
DENSITY	Resistor material density (unused)	kg/m <sup>3</sup>	0
HEATCAPACITY	Resistor material volumetric heat capacity	J/(m <sup>3</sup> K)	0
L	Length of conductor	m	0
OUTPUTINTVARS	Debug Output switch	–	false
R	Resistance	Ω	1000
RESISTIVITY	Resistor material resistivity	Ω m	0
TEMP	Device temperature	°C	Ambient Tempera- ture
THERMAL_A	Area of material thermally coupled to conductor	m <sup>2</sup>	0
THERMAL_HEATCAPACITY	Volumetric heat capacity of material thermally coupled to conductor	J/(m <sup>3</sup> K)	0
THERMAL_L	Length of material thermally coupled to conductor	m	0
W	Width of conductor	m	0

### Thermal Resistor Model Parameters


Table 2.29: Resistor Device Model Parameters

Parameter	Description	Units	Default
DEFW	Default Instance Width	m	1e-05
DENSITY	Resistor material density (unused)	kg/m <sup>3</sup>	0
HEATCAPACITY	Resistor material volumetric heat capacity	J/(m <sup>3</sup> K)	0
NARROW	Narrowing due to side etching	m	0
RESISTIVITY	Resistor material resistivity	Ω m	0
RSH	Sheet Resistance	Ω	0

Table 2.29: Resistor Device Model Parameters

Parameter	Description	Units	Default
TC1	Linear Temperature Coefficient	$^{\circ}\text{C}^{-1}$	0
TC2	Quadratic Temperature Coefficient	$^{\circ}\text{C}^{-2}$	0
THERMAL_HEATCAPACITY	Volumetric heat capacity of material thermally coupled to conductor	$\text{J}/(\text{m}^3\text{K})$	0
TNOM	Parameter Measurement Temperature	$^{\circ}\text{C}$	Ambient Temperature

## 2.3.6 Diode

<b>Symbol</b>	
<b>Instance Form</b>	D<name> <(+) node> <(-) node> <model name> [area value]
<b>Model Form</b>	.MODEL <model name> D [model parameters]
<b>Examples</b>	DCLAMP 1 0 DMOD D2 15 17 SWITCH 1.5
<b>Parameters and Options</b>	(+) node (-) node The anode and the cathode. area value Scales IS, ISR, IKF, RS, CJO, and IBV, and has a default value of 1. IBV and BV are both specified as positive values.
<b>Comments</b>	The diode is modeled as an ohmic resistance ( $R_S/\text{area}$ ) in series with an intrinsic diode. Positive current is current flowing from the anode through the diode to the cathode.

### Diode Operating Temperature

Model parameters can be assigned unique measurement temperatures using the **TNOM** model parameter.

### Diode level selection

Two distinct implementations of the diode are available. These are selected by using the **LEVEL** model parameter. The default implementation is based on SPICE 3F5, and may be explicitly specified using **LEVEL=1** in the model parameters, but is also selected if no **LEVEL** parameter is specified. The PSpice implementation [2] is obtained by specifying **LEVEL=2**.

The **Xyce** **LEVEL=1** and **LEVEL=2** diodes have a parameter, **IRF**, that allows the user to adjust the reverse current from the basic SPICE implementation. The usual SPICE treatment defines the linear portion of the reverse current in terms of **IS** which is defined by the forward current characteristics. Data shows that often the reverse current is quite far off when determined in this manner. The parameter **IRF** is a multiplier that can be applied to adjust the linear portion of the reverse current.

## Device Parameters

Table 2.30: Diode Device Instance Parameters

Parameter	Description	Units	Default
AREA	Area scaling value (scales IS, ISR, IKF, RS, CJO, and IBV)	–	1
IC		–	0
LAMBERTW	Option to solve diode equations with the Lambert-W function	logical (T/F)	0
OFF	Initial voltage drop across device set to zero	logical (T/F)	0
TEMP	Device temperature	–	Ambient Temperature

## Model Parameters

Table 2.31: Diode Device Model Parameters

Parameter	Description	Units	Default
AF	Flicker noise exponent	–	1
BV	Reverse breakdown "knee" voltage	V	1e+99
CJ	Zero-bias p-n depletion capacitance	F	0
CJO	Zero-bias p-n depletion capacitance	F	0
CJO	Zero-bias p-n depletion capacitance	F	0
EG	Bandgap voltage (barrier height)	eV	1.11
FC	Forward-bias depletion capacitance coefficient	–	0.5
IBV	Reverse breakdown "knee" current	A	0.001
IBVL	Low-level reverse breakdown "knee" current (level 2)	A	0
IKF	High-injection "knee" current (level 2)	A	0
IRF	Reverse current fitting factor	–	1
IS	Saturation current	A	1e-14
ISR	Recombination current parameter (level 2)	A	0
JS	Saturation current	A	1e-14
KF	Flicker noise coefficient	–	0
M	Grading parameter for p-n junction	–	0.5
N	Emission coefficient	–	1
NBV	Reverse breakdown ideality factor (level 2)	–	1
NBVL	Low-level reverse breakdown ideality factor (level 2)	–	1

Table 2.31: Diode Device Model Parameters

Parameter	Description	Units	Default
NR	Emission coefficient for ISR (level 2)	—	2
RS	Parasitic resistance	$\Omega$	0
TBV1	BV temperature coefficient (linear) (level 2)	$^{\circ}\text{C}^{-1}$	0
TBV2	BV temperature coefficient (quadratic) (level 2)	$^{\circ}\text{C}^{-2}$	0
TIKF	IKF temperature coefficient (linear) (level 2)	$^{\circ}\text{C}^{-1}$	0
TNOM		—	Ambient Temperature
TRS1	RS temperature coefficient (linear) (level 2)	$^{\circ}\text{C}^{-1}$	0
TRS2	RS temperature coefficient (quadratic) (level 2)	$^{\circ}\text{C}^{-2}$	0
TT	Transit time	s	0
VB	Reverse breakdown "knee" voltage	V	1e+99
VJ	Potential for p-n junction	V	1
XTI	IS temperature exponent	—	3

## Diode Equations

The equations in this section use the following variables:

- $V_{di}$  = voltage across the intrinsic diode only
- $V_{th}$  =  $k \cdot T/q$  (thermal voltage)
- $k$  = Boltzmann's constant
- $q$  = electron charge
- $T$  = analysis temperature (Kelvin)
- $T_0$  = nominal temperature (set using TNOM option)
- $\omega$  = Frequency (Hz)

Other variables are listed above in the diode model parameters.

### Level=1

The level 1 diode is based on the Spice3f5 level 1 model.

### DC Current (Level=1)

The intrinsic diode current consists of forward and reverse bias regions where

$$I_D = \begin{cases} \text{IS} \cdot \left[ \exp\left(\frac{V_{di}}{\text{NV}_{th}}\right) - 1 \right], & V_{di} > -3.0 \cdot \text{NV}_{th} \\ -\text{IS} \cdot \text{IRF} \cdot \left[ 1.0 + \left( \frac{3.0 \cdot \text{NV}_{th}}{V_{di} \cdot e} \right)^3 \right], & V_{di} < -3.0 \cdot \text{NV}_{th} \end{cases}$$

**IRF** is a **Xyce**-specific parameter that can be used to scale the reverse-biased current to match measured data. It defaults to 1.0, which reduces the model to strict SPICE3F5 compatibility.

When **BV** and an optional parameter **IBV** are explicitly given in the model statement, an exponential model is used to model reverse breakdown (with a “knee” current of **IBV** at a “knee-on” voltage of **BV**). The equation for  $I_D$  implemented by **Xyce** is given by

$$I_D = -\mathbf{IBV}_{\text{eff}} \cdot \exp\left(-\frac{\mathbf{BV}_{\text{eff}} + V_{di}}{\mathbf{NV}_{th}}\right), \quad V_{di} \leq \mathbf{BV}_{\text{eff}},$$

where  $\mathbf{BV}_{\text{eff}}$  and  $\mathbf{IBV}_{\text{eff}}$  are chosen to satisfy the following constraints:

1. Continuity of  $I_D$  between reverse bias and reverse breakdown regions (i.e., continuity of  $I_D$  at  $V_{di} = -\mathbf{BV}_{\text{eff}}$ ):

$$\mathbf{IBV}_{\text{eff}} = \mathbf{IRF} \cdot \mathbf{IS} \left(1 - \left(\frac{3.0 \cdot \mathbf{NV}_{th}}{e \cdot \mathbf{BV}_{\text{eff}}}\right)^3\right)$$

2. “Knee-on” voltage/current matching:

$$\mathbf{IBV}_{\text{eff}} \cdot \exp\left(-\frac{\mathbf{BV}_{\text{eff}} - \mathbf{BV}}{\mathbf{NV}_{th}}\right) = \mathbf{IBV}$$

Substituting the first expression into the second yields a single constraint on  $\mathbf{BV}_{\text{eff}}$  which cannot be solved for directly. By performing some basic algebraic manipulation and rearranging terms, the problem of finding  $\mathbf{BV}_{\text{eff}}$  which satisfies the above two constraints can be cast as finding the (unique) solution of the equation

$$\mathbf{BV}_{\text{eff}} = f(\mathbf{BV}_{\text{eff}}), \quad (2.18)$$

where  $f(\cdot)$  is the function that is obtained by solving for the  $\mathbf{BV}_{\text{eff}}$  term which appears in the exponential in terms of  $\mathbf{BV}_{\text{eff}}$  and the other parameters. **Xyce** solves Eqn. 2.18 by performing the so-called *Picard Iteration* procedure [6], i.e. by producing successive estimates of  $\mathbf{BV}_{\text{eff}}$  (which we will denote as  $\mathbf{BV}_{\text{eff}}^k$ ) according to

$$\mathbf{BV}_{\text{eff}}^{k+1} = f(\mathbf{BV}_{\text{eff}}^k)$$

starting with an initial guess of  $\mathbf{BV}_{\text{eff}}^0 = \mathbf{BV}$ . The current iteration procedure implemented in **Xyce** can be shown to guarantee at least six significant digits of accuracy between the numerical estimate of  $\mathbf{BV}_{\text{eff}}$  and the true value.

In addition to the above, **Xyce** also requires that  $\mathbf{BV}_{\text{eff}}$  lie in the range  $\mathbf{BV} \geq \mathbf{BV}_{\text{eff}} \geq 3.0\mathbf{NV}_{th}$ . In terms of **IBV**, this is equivalent to enforcing the following two constraints:

$$\mathbf{IRF} \cdot \mathbf{IS} \left(1 - \left(\frac{3.0 \cdot \mathbf{NV}_{th}}{e \cdot \mathbf{BV}}\right)^3\right) \leq \mathbf{IBV} \quad (2.19)$$

$$\mathbf{IRF} \cdot \mathbf{IS} (1 - e^{-3}) \exp \left( \frac{-3.0 \cdot \mathbf{NV}_{th} + \mathbf{BV}}{\mathbf{NV}_{th}} \right) \geq \mathbf{IBV} \quad (2.20)$$

**Xyce** first checks the value of **IBV** to ensure that the above two constraints are satisfied. If Eqn. 2.19 is violated, **Xyce** sets **IBV<sub>eff</sub>** to be equal to the left-hand side of Eqn. 2.19 and, correspondingly, sets **BV<sub>eff</sub>** to  $-3.0 \cdot \mathbf{NV}_{th}$ . If Eqn. 2.20 is violated, **Xyce** sets **IBV<sub>eff</sub>** to be equal to the left-hand side of Eqn. 2.20 and, correspondingly, sets **BV<sub>eff</sub>** to **BV**.

#### Capacitance (Level=1)

The p-n diode capacitance consists of a depletion layer capacitance  $C_d$  and a diffusion capacitance  $C_{dif}$ . The first is given by

$$C_d = \begin{cases} \mathbf{CJ} \cdot \mathbf{AREA} \left(1 - \frac{V_{di}}{\mathbf{VJ}}\right)^{-M}, & V_{di} \leq \mathbf{FC} \cdot \mathbf{VJ} \\ \frac{\mathbf{CJ} \cdot \mathbf{AREA}}{\mathbf{F2}} \left(\mathbf{F3} + M \frac{V_{di}}{\mathbf{VJ}}\right), & V_{di} > \mathbf{FC} \cdot \mathbf{VJ} \end{cases}$$

The diffusion capacitance (sometimes referred to as the transit time capacitance) is

$$C_{dif} = \mathbf{TT} G_d = \mathbf{TT} \frac{dI_D}{dV_{di}}$$

where  $G_d$  is the junction conductance.

#### Temperature Effects (Level=1)

The diode model contains explicit temperature dependencies in the ideal diode current, the generation/recombination current and the breakdown current. Further temperature dependencies are present in the diode model via the saturation current  $I_S$ , the depletion layer junction capacitance  $\mathbf{CJ}$ , the junction potential  $V_J$ .


$$\begin{aligned} V_t(T) &= \frac{kT}{q} \\ V_{tnom}(T) &= \frac{k\mathbf{TNOM}}{q} \\ E_g(T) &= E_{g0} - \frac{\alpha T^2}{\beta + T} \\ E_{gNOM}(T) &= E_{g0} - \frac{\alpha \mathbf{TNOM}^2}{\mathbf{TNOM} + \beta} \\ arg1(T) &= -\frac{E_g(T)}{2kT} + \frac{E_{g300}}{2kT_0} \\ arg2(T) &= -\frac{E_{gNOM}(T)}{2k\mathbf{TNOM}} + \frac{E_{g300}}{2kT_0} \\ pbfact1(T) &= -2.0 \cdot V_t(T) \left( 1.5 \cdot \ln \left( \frac{T}{T_0} \right) + q \cdot arg1(T) \right) \end{aligned}$$

$$\begin{aligned}
pbfact2(T) &= -2.0 \cdot V_{tnom}(T) \left( 1.5 \cdot \ln \left( \frac{\mathbf{TNOM}}{T_0} \right) + q \cdot arg2(T) \right) \\
pbo(T) &= (\mathbf{VJ} - pbfact2(T)) \frac{T_0}{\mathbf{TNOM}} \\
V_J(T) &= pbfact1(T) + \frac{T}{T_0} pbo(T) \\
gma_{old}(T) &= \frac{\mathbf{VJ} - pbo(T)}{pbo(T)} \\
gma_{new}(T) &= \frac{V_J(T) - pbo(T)}{pbo(T)} \\
CJ(T) &= \mathbf{CJ0} \frac{1.0 + \mathbf{M} (4.0 \times 10^{-4} (T - T_0) - gma_{new}(T))}{1.0 + \mathbf{M} (4.0 \times 10^{-4} (\mathbf{TNOM} - T_0) - gma_{old}(T))} \\
I_S(T) &= \mathbf{IS} \cdot \exp \left( \left( \frac{T}{\mathbf{TNOM}} - 1.0 \right) \cdot \frac{\mathbf{EG}}{\mathbf{NV}_t(T)} + \frac{\mathbf{XTI}}{\mathbf{N}} \cdot \ln \left( \frac{T}{\mathbf{TNOM}} \right) \right)
\end{aligned}$$

where, for silicon,  $\alpha = 7.02 \times 10^{-4} \text{ eV/K}$ ,  $\beta = 1108 \text{ K}$  and  $E_{g0} = 1.16 \text{ eV}$ .

For a more thorough description of p-n junction physics, see [9]. For a thorough description of the U.C. Berkeley SPICE models see Reference [11].

# 2.3.7 Independent Current Source

Symbol	
Instance Form	<code>I&lt;name&gt; &lt;(+) node&gt; &lt;(-) node&gt; [ [DC] &lt;value&gt; ]</code> <code>+ [AC [magnitude value [phase value] ] ]</code> <code>+ [transient specification]</code>
Examples	<code>ISLOW 1 22 SIN(0.5 1.0ma 1kHz 1ms)</code> <code>IPULSE 1 3 PULSE(-1 1 2ns 2ns 2ns 50ns 100ns)</code>
Parameters and Options	<p>transient specification</p> <p>There are five predefined time-varying functions for sources:</p> <p><code>PULSE &lt;parameters&gt;</code> Pulse waveform</p> <p><code>SIN &lt;parameters&gt;</code> Sinusoidal waveform</p> <p><code>EXP &lt;parameters&gt;</code> Exponential waveform</p> <p><code>PWL &lt;parameters&gt;</code> Piecewise linear waveform</p> <p><code>SFFM &lt;parameters&gt;</code> Frequency-modulated waveform</p>
Comments	Positive current flows from the positive node through the source to the negative node. The default value is zero for the DC, AC, and transient values. None, any, or all of the DC, AC, and transient values can be specified. The AC phase value is in degrees.

## Transient Specifications

This section outlines the available transient specifications.  $\Delta t$  and  $T_F$  are the time step size and simulation end-time, respectively.

### Pulse

`PULSE(I1 I2 TD TR TF PW PER)`

Table 2.32: Pulse Parameters

Parameter	Description	Units	Default
I1	Initial Value	amp	–
I2	Pulse Value	amp	–

Table 2.32: Pulse Parameters

Parameter	Description	Units	Default
TD	Delay Time	s	0.0
TR	Rise Time	s	$\Delta t$
TF	Fall Time	s	$\Delta t$
PW	Pulse Width	s	$T_F$
PER	Period	s	$T_F$

### Sine

SIN(I0 IA FREQ TD THETA)

Table 2.33: Sine Parameters

Parameter	Description	Units	Default
I0	Offset	amp	–
IA	Amplitude	amp	–
FREQ	Frequency	$s^{-1}$	0.0
TD	Delay	s	$\Delta t$
THETA	Attenuation Factor	s	$\Delta t$

The waveform is shaped according to the following equations:

$$I = \begin{cases} I_0, & 0 < t < T_D \\ I_0 + I_A \sin[2\pi \cdot \mathbf{FREQ} \cdot (t - T_D)] \exp[-(t - T_D) \cdot \mathbf{THETA}], & T_D < t < T_F \end{cases}$$

### Exponent

EXP(I1 I2 TD1 TAU1 TD2 TAU2)

Table 2.34: Exponent Parameters

Parameter	Description	Units	Default
I1	Initial Amplitude	amp	N/A
I2	Amplitude	amp	N/A
TD1	Rise Delay Time	s	0.0
TAU1	Rise Time Constant	s	$\Delta t$
TD2	Delay Fall Time	s	$TD1 + \Delta t$
TAU2	Fall Time Constant	s	$\Delta t$

The waveform is shaped according to the following equations:

$$I = \begin{cases} I_1, & 0 < t < TD1 \\ I_1 + (I_2 - I_1)\{1 - \exp[-(t - TD1)/TAU1]\}, & TD1 < t < TD2 \\ I_1 + (I_2 - I_1)\{1 - \exp[-(t - TD1)/TAU1]\} \\ \quad + (I_1 - I_2)\{1 - \exp[-(t - TD2)/TAU2]\}, & TD2 < t < T_2 \end{cases}$$

### Piecewise Linear

PWL T0 IO [Tn In]\*

PWL FILE "<name>" [TD=<timeDelay>] [R=<repeatTime>]

Table 2.35: Piecewise Linear Parameters

Parameter	Description	Units	Default
$T_n$	Time at Corner	s	none
$I_n$	Current at Corner	amp	none
TD	Time Delay	s	0
R	Repeat Time	s	none

When the FILE option is given, **Xyce** will read the corner points from the file specified in the <name> field. This file should be a plain ASCII text file with time/current pairs. There should be one pair per line, and the time and current values should be separated by whitespace or commas.

TD has units of seconds, and specifies the length of time to delay the start of PWL waveform. The default is to have no delay, and TD is an optional parameter.

The Repeat Time (R) is an optional parameter. If R is omitted then the waveform will not repeat. If R is included then the waveform will repeat until the end of the simulation. As examples, R=0 means repeat the PWL waveform from time=0. In general, R=<repeatTime> means repeat the waveform from time equal to <repeatTime> seconds in the waveform specification, where <repeatTime> must be greater than or equal to 0 and less than  $T_n$ . If the R parameter is used then it must have a value. The specification PWL FILE "<name>" R is illegal in **Xyce** as a shorthand for R=0. Finally, the **Xyce** syntax for PWL sources is not compatible with the PSpice REPEAT syntax for PWL sources.

### Frequency Modulated

SFFM (IOFF IAMPL FC MOD FM)

Table 2.36: Frequency Modulated Parameters

Parameter	Description	Units	Default
IOFF	Offset Current	amp	none

Table 2.36: Frequency Modulated Parameters

Parameter	Description	Units	Default
IAMPL	Peak Current Amplitude	amp	none
FC	Carrier Frequency	hertz	1/TSTOP
MOD	Modulation Index	-	0
FM	Modulation Frequency	hertz	1/TSTOP

**TSTOP** is the final time, as entered into the transient (.TRANS) command. The waveform is shaped according to the following equation:

$$I = \text{ioff} + \text{iamp1} \cdot \sin(2\pi \cdot \text{fc} \cdot \text{TIME} + \text{mod} \cdot \sin(2\pi \cdot \text{fm} \cdot \text{TIME}))$$

where **TIME** is the current simulation time.

## 2.3.8 Independent Voltage Source



### Symbol

---

**Instance Form**    `V<name> <(+) node> <(-) node> [ [DC] <value> ]`  
                          `+ [AC [magnitude value [phase value] ] ] [transient specification]`

---

**Examples**            `VSLOW 1 22 SIN(0.5 1.0ma 1kHz 1ms)`  
                          `VPULSE 1 3 PULSE(-1 1 2ns 2ns 2ns 50ns 100ns)`

---

### Parameters and Options

transient specification

There are five predefined time-varying functions for sources:

PULSE <parameters> Pulse waveform

SIN <parameters> Sinusoidal waveform

EXP <parameters> Exponential waveform

PWL <parameters> Piecewise linear waveform

SFFM <parameters> Frequency-modulated waveform

---

**Comments**            Positive current flows from the positive node through the source to the negative node. None, any, or all of the DC, AC, and transient values can be specified. The AC phase value is in degrees.

## Transient Specifications

This section outlines the available transient specifications.  $\Delta t$  and  $T_F$  are the time step size and simulation end-time, respectively.

### Pulse

`PULSE(V1 V22 TD TR TF PW PER)`

Table 2.37: Pulse Parameters

Parameter	Description	Units	Default
V1	Initial Value	Volt	–
V2	Pulse Value	Volt	–
TD	Delay Time	s	0.0
TR	Rise Time	s	$\Delta t$

Table 2.37: Pulse Parameters

Parameter	Description	Units	Default
TF	Fall Time	s	$\Delta t$
PW	Pulse Width	s	$T_F$
PER	Period	s	$T_F$

### Sine

SIN(V0 VA FREQ TD THETA)

Table 2.38: Sine Parameters

Parameter	Description	Units	Default
V0	Offset	Volt	–
VA	Amplitude	Volt	–
FREQ	Frequency	$s^{-1}$	0.0
TD	Delay	s	$\Delta t$
THETA	Attenuation Factor	s	$\Delta t$

The waveform is shaped according to the following equations:

$$V = \begin{cases} V_0, & 0 < t < T_D \\ V_0 + V_A \sin[2\pi \cdot \mathbf{FREQ} \cdot (t - T_D)] \exp[-(t - T_D) \cdot \mathbf{THETA}], & T_D < t < T_F \end{cases}$$

### Exponent

EXP(V1 V2 TD1 TAU1 TD2 TAU2)

Table 2.39: Exponent Parameters

Parameter	Description	Units	Default
V1	Initial Amplitude	Volt	N/A
V2	Amplitude	Volt	N/A
TD1	Rise Delay Time	s	0.0
TAU1	Rise Time Constant	s	$\Delta t$
TD2	Delay Fall Time	s	$TD1 + \Delta t$
TAU2	Fall Time Constant	s	$\Delta t$

The waveform is shaped according to the following equations:

$$V = \begin{cases} V_1, & 0 < t < TD1 \\ V_1 + (V_2 - V_1)\{1 - \exp[-(t - TD1)/TAU1]\}, & TD1 < t < TD2 \\ V_1 + (V_2 - V_1)\{1 - \exp[-(t - TD1)/TAU1]\} \\ \quad + (V_1 - V_2)\{1 - \exp[-(t - TD2)/TAU2]\}, & TD2 < t < T_2 \end{cases}$$

### Piecewise Linear

PWL T0 V0 [Tn Vn]\*

PWL FILE "<name>" [TD=<timeDelay>] [R=<repeatTime>]

Table 2.40: Piecewise Linear Parameters

Parameter	Description	Units	Default
$T_n$	Time at Corner	s	none
$V_n$	Voltage at Corner	Volt	none
TD	Time Delay	s	0
R	Repeat Time	s	none

When the FILE option is given, **Xyce** will read the corner points from the file specified in the <name> field. This file should be a plain ASCII text file with time/voltage pairs. There should be one pair per line, and the time and voltage values should be separated by whitespace or commas.

TD has units of seconds, and specifies the length of time to delay the start of PWL waveform. The default is to have no delay, and TD is an optional parameter.

The Repeat Time (R) is an optional parameter. If R is omitted then the waveform will not repeat. If R is included then the waveform will repeat until the end of the simulation. As examples, R=0 means repeat the PWL waveform from time=0. In general, R=<repeatTime> means repeat the waveform from time equal to <repeatTime> seconds in the waveform specification, where <repeatTime> must be greater than or equal to 0 and less than  $T_n$ . If the R parameter is used then it must have a value. The specification PWL FILE "<name>" R is illegal in **Xyce** as a shorthand for R=0. Finally, the **Xyce** syntax for PWL sources is not compatible with the PSpice REPEAT syntax for PWL sources.

### Frequency Modulated

SFFM (VOFF VAMPL FC MOD FM)

Table 2.41: Frequency Modulated Parameters

Parameter	Description	Units	Default
VOFF	Offset Current	Volt	none

Table 2.41: Frequency Modulated Parameters

Parameter	Description	Units	Default
VAMPL	Peak Current Amplitude	Volt	none
FC	Carrier Frequency	hertz	1/TSTOP
MOD	Modulation Index	-	0
FM	Modulation Frequency	hertz	1/TSTOP

**TSTOP** is the final time, as entered into the transient (.TRANs) command. The waveform is shaped according to the following equation:

$$V = \mathbf{voff} + \mathbf{vAMPL} \cdot \sin(2\pi \cdot \mathbf{fc} \cdot \mathbf{TIME} + \mathbf{mod} \cdot \sin(2\pi \cdot \mathbf{fm} \cdot \mathbf{TIME}))$$

where **TIME** is the current simulation time.

## 2.3.9 Voltage Controlled Voltage Source



Symbol

---

**Instance Form**

```
E<name> <(+) node> <(-) node> <(+) controlling node>
+ <(-) controlling node> <gain>
E<name> <(+) node> <(-) node> VALUE = <expression>
E<name> <(+) node> <(-) node> TABLE <expression> =
+ < <input value>,<output value> >*
E<name> <(+) node> <(-) node> POLY(<value>)
+ [<+ control node> <- control node>]*
+ [<polynomial coefficient value>]*
```

---

**Examples**

```
EBUFFER 1 2 10 11 5.0
ESQROOT 5 0 VALUE = 5V*SQRT(V(3,2))
ET2 2 0 TABLE V(ANODE,CATHODE) = (0,0) (30,1)
EP1 5 1 POLY(2) 3 0 4 0 0 .5 .5
```

---

### Parameters and Options

(+) node  
(-) node  
Output nodes. Positive current flows from the (+) node through the source to the (-) node.

(+) controlling node  
(-) controlling node  
Node pairs that define a set of controlling voltages. A given node may appear multiple times and the output and controlling nodes may be the same.

---

**Comments**

In the first form, a specified voltage drop between controlling nodes is multiplied by the gain to determine the voltage drop across the output nodes.

The second through fourth forms allow nonlinear controlled sources using the VALUE, TABLE, or POLY keywords, respectively, and are used in analog behavioral modeling. They are provided primarily for netlist compatibility with other simulators. These three forms are automatically converted within **Xyce** to its principal ABM device, the B nonlinear dependent source device. See the **Xyce** User's Guide for more information on analog behavioral modeling. For details concerning the use of the POLY format, see section 2.2.4.

## 2.3.10 Current Controlled Current Source



### Symbol

---

**Instance Form**    F<name> <(+) node> <(-) node>  
                      + <controlling V device name> <gain>  
                      F<name> <(+) node> <(-) node> POLY(<value>)  
                      + <controlling V device name> \*  
                      + < <polynomial coefficient value> > \*

---

**Examples**            FSENSE 1 2 VSENSE 10.0  
                      FAMP 13 0 POLY(1) VIN 0 500  
                      FNONLIN 100 101 POLY(2) VCINTRL1 VCINTRL2 0.0 13.6 0.2 0.005

---

### Parameters and Options

(+) node  
(-) node  
      Output nodes. Positive current flows from the (+) node through the source to the (-) node.

controlling V device  
      The controlling voltage source which must be an independent voltage source (V device).

---

**Comments**            In the first form, a specified current through a controlling device is multiplied by the gain to determine this device's output current.

                      The second form using the POLY keyword is used in analog behavioral modeling. This form is automatically converted within **Xyce** to its principal ABM device, the B nonlinear dependent source device. See the **Xyce** User's Guide for more information on analog behavioral modeling. For details concerning the use of the POLY format, see section 2.2.4.

## 2.3.11 Current Controlled Voltage Source

The syntax of this device is exactly the same as for a Current Controlled Current Source. For a Current-Controlled Voltage Source just substitute an H for the F. The H device generates a voltage, whereas the F device generates a current.



### Symbol

---

<b>Instance Form</b>	<pre>H&lt;name&gt; &lt;(+) node&gt; &lt;(-) node&gt; + &lt;controlling V device name&gt; &lt;gain&gt; H&lt;name&gt; &lt;(+) node&gt; &lt;(-) node&gt; VALUE= &lt;expression&gt; H&lt;name&gt; &lt;(+) node&gt; &lt;(-) node&gt; TABLE &lt;expression&gt; = + [&lt;input value&gt;, &lt;output value&gt;]* H&lt;name&gt; &lt;(+) node&gt; &lt;(-) node&gt; POLY(&lt;value&gt;) + &lt;controlling V device name&gt;* + &lt; &lt;polynomial coefficient value&gt; &gt;*</pre>
----------------------	--

---

<b>Examples</b>	<pre>HSENSE 1 2 VSENSE 10.0 HAMP 13 0 POLY(1) VIN 0 500 HNONLIN 100 101 POLY(2) VCNTRL1 VCINTRL2 0.0 13.6 0.2 0.005</pre>
-----------------	---

---

<b>Comments</b>	<p>In the first form, the current through a specified voltage source controls is multiplied by a constant to obtain the voltage-source output.</p> <p>The second through fourth forms using the VALUE, TABLE, or POLY keywords, respectively, are used in analog behavioral modeling. They are provided primarily for netlist compatibility with other simulators. These three forms are automatically converted within <b>Xyce</b> to its principal ABM device, the B nonlinear dependent source device. See the <b>Xyce</b> User's Guide for more information on analog behavioral modeling. For details concerning the use of the POLY format, see section 2.2.4.</p>
-----------------	--

## 2.3.12 Voltage Controlled Current Source



Symbol

---

**Instance Form**

```
G<name> <(+) node> <(-) node> <(+) controlling node>
+ <(-) controlling node> <transconductance>
G<name> <(+) <node> <(-) node> VALUE = <expression>
G<name> <(+) <node> <(-) node> TABLE <expression> =
+ < <input value>,<output value> >*
G<name> <(+) <node> <(-) node> POLY(<value>)
+ [<+ controlling node> <- controlling node>]*
+ [<polynomial coefficient>]*
```

---

**Examples**

```
GBUFFER 1 2 10 11 5.0
GPSK 11 6 VALUE = 5MA*SIN(6.28*10kHz*TIME+V(3))
GA2 2 0 TABLE V(5) = (0,0) (1,5) (10,5) (11,0)
```

---

### Parameters and Options

(+) node  
(-) node  
Output nodes. Positive current flows from the (+) node through the source to the (-) node.

(+) controlling node  
(-) controlling node  
Node pairs that define a set of controlling voltages. A given node may appear multiple times and the output and controlling nodes may be the same.

---

**Comments**

In the first form, the voltage drop between the controlling nodes is multiplied by the transconductance to obtain the current-source output of the G device.

The second through fourth forms using the VALUE, TABLE, and POLY keywords, respectively, are used in analog behavioral modeling. They are provided primarily for netlist compatibility with other simulators. These two forms are automatically converted within **Xyce** to its principal ABM device, the B nonlinear dependent source device. See the **Xyce** User's Guide for more information on analog behavioral modeling. For details concerning the use of the POLY format, see section 2.2.4.

## 2.3.13 Nonlinear Dependent Source

---

**Instance Form**    B<name> <(+) node> <(-) node> V=ABM expression  
                      B<name> <(+) node> <(-) node> I=ABM expression

---

**Examples**        B1 2 0 V={sqrt(V(1))}  
                      B2 4 0 V={V(1)\*TIME}  
                      B3 4 2 I={I(V1) + V(4,2)/100}  
                      B4 5 0 V={Table V(5)=(0,0) (1.0,2.0) (2.0,3.0) (3.0,10.0)}

---

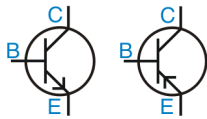
**Comments**        The nonlinear dependent source device, also known as the B-source device, is used in analog behavioral modeling (ABM). The (+) and (-) nodes are the output nodes. Positive current flows from the (+) node through the source to the (-) node.

See the “Analog Behavioral Modeling” chapter of the **Xyce** User’s Guide for guidance on using the Bsource device and ABM expressions, and the Expressions section 2.2 for complete documentation of expressions and expression operators.

**Note: the braces surrounding all expressions are required.**

# 2.3.14 Bipolar Junction Transistor (BJT)

## Symbol



## Instance Form

Q<name> <collector node> <base node> <emitter node>  
+ [substrate node] <model name> [area value]

Q<name> <collector node> <base node> <emitter node>  
+ <thermal node> <VBIC model name>

## Model Form

.MODEL <model name> NPN [model parameters]  
.MODEL <model name> PNP [model parameters]

## Examples

```
Q2 10 2 9 PNP1
Q12 14 2 0 1 NPN2 2.0
Q6 VC 4 11 [SUB] LAXPNP
Q6 Coll Base Emit DT VBICMODEL1
```

## Parameters and Options

substrate node  
Optional and defaults to ground. Since **Xyce** permits alphanumeric node names and because there is no easy way to make a distinction between these and the model names, the name (not a number) used for the substrate node must be enclosed in square brackets [ ]. Otherwise, nodes would be interpreted as model names. See the fourth example above.

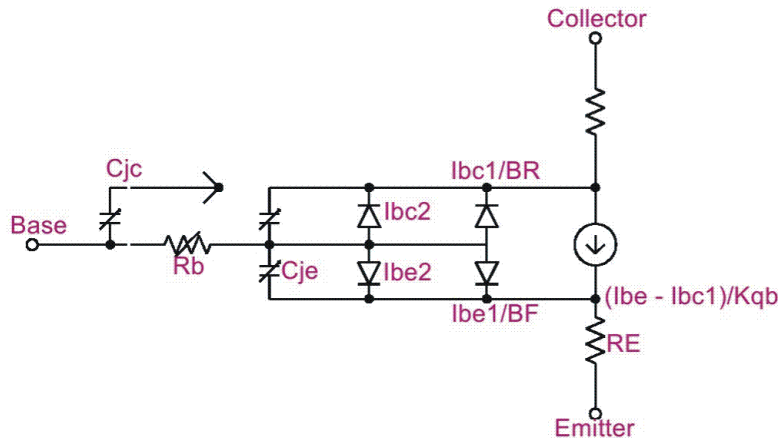
area value  
The relative device area with a default value of 1.

## Comments

The BJT is modeled as an intrinsic transistor using ohmic resistances in series with the collector (RC/area), with the base (value varies with current, see BJT equations) and with the emitter (RE/area). For model parameters with optional names, such as VAF and VA (the optional name is in parentheses), either may be used. For model types NPN and PNP, the isolation junction capacitance is connected between the intrinsic-collector and substrate nodes. This is the same as in SPICE and works well for vertical IC transistor structures.

**The VBIC model requires a slightly different form of the instance line than does the level 1 BJT; this variant of the Q line is shown in the fourth example above.** VBIC instance lines have four required nodes, the first three are

the normal collector, base, and emitter, and the fourth node is for electrothermal effects. This fourth node, named “dt” in the VBIC literature, is the difference between the device temperature including self-heating and the baseline temperature of the device. The base temperature of the device is the sum of the ambient temperature of the simulation and the DTEMP model parameter. It is common to tie this “dt” node to ground using a large-value resistor and to use the node only for output to observe the device heating, but it can also be used to couple the thermal effects of several VBIC models.



**Figure 2.1.** BJT model schematic. Adapted from reference [2].

## BJT Level selection

**Xyce** supports the level 1 BJT model, which is based on the documented standard SPICE 3F5 BJT model, but was coded independently at Sandia. It is mostly based on the classic Gummel-Poon BJT model [7].

A version of the VBIC model is provided as BJT level 10. This is the 3-terminal, electrothermal, constant phase model of VBIC version 1.2 [8].

The VBIC model supports both PNP and NPN transistors, and may therefore be used with model cards of type PNP and NPN.

An experimental release of the FBH HBT\_X model version 2.1[9] is provided as BJT level 23.

## BJT Operating Temperature

Model parameters may be assigned unique measurement temperatures using the **TNOM** model parameter. See BJT model parameters for more information.

## Level=1 Instance Parameters

Table 2.42 gives the available instance parameters for the level 1 BJT.

Table 2.42: Bipolar Junction Transistor Device Instance Parameters

Parameter	Description	Units	Default
AREA	Relative device area	—	1
IC1	Vector of initial values: Vbe, Vce. Vbe=IC1	V	0
IC2	Vector of initial values: Vbe, Vce. Vce=IC2	V	0
LAMBERTW	Flag for toggling the use of the lambert-W function instead of exponentials.	logical (T/F)	false
OFF	Initial condition of no voltage drops accross device	logical (T/F)	false
TEMP	Device temperature	°C	Ambient Temperature

### Level=1 Model Parameters

Table 2.43 gives the available model parameters for the level 1 BJT.

Table 2.43: Bipolar Junction Transistor Device Model Parameters

Parameter	Description	Units	Default
AF	Flicker noise exponent	—	1
BF	Ideal maximum foward beta	—	100
BFM	Ideal maximum foward beta	—	100
BR	Ideal maximum reverse beta	—	1
BRM	Ideal maximum reverse beta	—	1
BV	Reverse early voltage	V	0
C2	Coefficient for base-emitter leak current.	—	0
C4	Coefficient for base-collector leak current.	—	0
CCS	Substrate zero-bias p-n capacitance	F	0
CDIS	Fraction of CJC connected internally to RB	—	1
CJC	Base-collector zero-bias p-n capacitance	F	0
CJE	Base-emitter zero-bias p-n capacitance	F	0
CJS	Substrate zero-bias p-n capacitance	F	0
CSUB	Substrate zero-bias p-n capacitance	F	0
EG	Bandgap voltage (barrier highth)	eV	1.11
ESUB	Substrate p-n grading factor	—	0
FC	Foward-bias depletion capacitor coefficient	—	0.5
IK	Corner for foward-beta high-current roll-off	A	0

Table 2.43: Bipolar Junction Transistor Device Model Parameters

Parameter	Description	Units	Default
IKF	Corner for forward-beta high-current roll-off	A	0
IKR	Corner for reverse-beta high-current roll-off	A	0
IOB	Current at which RB falls off by half	A	0
IRB	Current at which RB falls off by half	A	0
IS	Transport saturation current	A	1e-16
ISC	Base-collector leakage saturation current	A	0
ISE	Base-emitter leakage saturation current	A	0
ITF	Transit time dependency on IC	—	0
JBF	Corner for forward-beta high-current roll-off	A	0
JBR	Corner for reverse-beta high-current roll-off	A	0
JLC	Base-collector leakage saturation current	A	0
JLE	Base-emitter leakage saturation current	A	0
JRB	Current at which RB falls off by half	A	0
JTF	Transit time dependency on IC	—	0
KF	Flicker noise coefficient	—	0
MC	Base-collector p-n grading factor	—	0.33
ME	Base-emitter p-n grading factor	—	0.33
MJC	Base-collector p-n grading factor	—	0.33
MJE	Base-emitter p-n grading factor	—	0.33
MJS	Substrate p-n grading factor	—	0
MS	Substrate p-n grading factor	—	0
NC	Base-collector leakage emission coefficient	—	2
NE	Base-emitter leakage emission coefficient	—	1.5
NF	Forward current emission coefficient	—	1
NK	High current rolloff coefficient	—	0.5
NKF	High current rolloff coefficient	—	0.5
NLE	Base-emitter leakage emission coefficient	—	1.5
NR	Reverse current emission coefficient	—	1
PC	Base-collector built-in potential	V	0.75
PE	Base-emitter built-in potential	V	0.75
PS	Substrate built-in potential	V	0.75
PSUB	Substrate built-in potential	V	0.75
PT	Temperature exponent for IS. (synonymous with XTI)	—	3

Table 2.43: Bipolar Junction Transistor Device Model Parameters

Parameter	Description	Units	Default
PTF	Excess Phase at $1/(2\pi \cdot TF)$ Hz	degree	0
RB	Zero-bias (maximum) base resistance	$\Omega$	0
RBM	Maximum base resistance	$\Omega$	0
RC	Collector ohmic resistance	$\Omega$	0
RE	Emitter ohmic resistance	$\Omega$	0
TB	Foward and reverse beta temperature coefficient	—	0
TCB	Foward and reverse beta temperature coefficient	—	0
TEMPMODEL	Specifies the type of parameter interpolation over temperature	—	'NONE'
TF	Ideal foward transit time	s	0
TNOM	Parameter measurement temperature	$^{\circ}\text{C}$	Ambient Temperature
TR	Ideal reverse transit time	s	0
VA	Foward early voltage	V	0
VAF	Foward early voltage	V	0
VAR	Reverse early voltage	V	0
VB	Reverse early voltage	V	0
VBF	Foward early voltage	V	0
VJC	Base-collector built-in potential	V	0.75
VJE	Base-emitter built-in potential	V	0.75
VJS	Substrate built-in potential	V	0.75
VRB	Reverse early voltage	V	0
VTF	Transit time dependancy on Vbc	V	0
XCJC	Fraction of CJC connected internally to RB	—	1
XTB	Foward and reverse beta temperature coefficient	—	0
XTF	Transit time bias dependence coefficient	—	0
XTI	Temperature exponent for IS. (synonymous with PT)	—	3

### Level=10 instance parameters

The VBIC (level 10 transistor) supports a single instance parameter, *M* (Multiplicity). This parameter emulates an integer number of identical VBIC transistors connected in parallel. At this time, the VBIC is the only Q device that supports a multiplicity instance parameter. The level 1 Q device instead supports an AREA instance parameter that can be used for the same purpose.

## Level=10 model parameters

Table 2.44 gives the available device instance parameters and 2.45 gives the available model parameters for the level 10 BJT.

Table 2.44: VBIC 3T et cf v1.2 Device Instance Parameters

Parameter	Description	Units	Default
M	Number of devices in parallel	–	1

Table 2.45: VBIC 3T et cf v1.2 Device Model Parameters

Parameter	Description	Units	Default
AFN	Base-Emitter Flicker Noise coefficient (unused)	–	1
AJC	Base-Collector capacitor smoothing factor	–	-0.5
AJE	Base-Emitter capacitor smoothing factor	–	-0.5
AJS	Substrate-collector capacitor smoothing factor (unused)	–	-0.5
ART		–	0.1
AVC1	B-C weak avalanche parameter	$V^{-1}$	0
AVC2	B-C weak avalanche parameter	$V^{-1}$	0
BFN	B-E flicker noise dependence (unused)	–	1
CBC0	Extrinsic B-C overlap capacitance	F	0
CBE0	Extrinsic B-E overlap capacitance	F	0
CCS0	(unused)	–	0
CJC	B-C zero-bias capacitance	F	0
CJCP	S-C zero-bias capacitance	F	0
CJE	B-E zero-bias capacitance	F	0
CJEP	S-E zero-bias capacitance	F	0
CTH	Thermal capacitance	F	0
DEAR	Activation energy for ISRR	–	0
DTEMP	Device temperature (use 0.0 for ambient)	–	0
EA	Activation energy for IS	eV	1.12
EAIC	Activation energy for IBCI	eV	1.12
EAIE	Activation energy for IBEI	eV	1.12
EAIS	Activation energy for IBCIP	eV	1.12
EANC	Activation energy for IBCN	eV	1.12
EANE	Activation energy for IBEN	eV	1.12
EANS	Activation energy for IBCNP	eV	1.12
EAP	Activation energy for ISP	–	1.12

Table 2.45: VBIC 3T et cf v1.2 Device Model Parameters

Parameter	Description	Units	Default
EBBE	(unused)	–	0
FC	Forward-bias depletion capacitance limit	–	0.9
GAMM	Epi doping parameter	–	0
HRCF	High current RC factor	–	0
IBBE		–	1e-06
IBCI	Ideal B-C saturation current	A	1e-16
IBCIP	Ideal parasitic B-C saturation current	A	0
IBCN	Nonideal B-C saturation current	A	0
IBCNP	Nonideal parasitic B-C saturation current	A	0
IBEI	Ideal B-E saturation current	A	1e-18
IBEIP	Ideal parasitic B-E saturation current	A	0
IBEN	Nonideal B-E saturation current	A	0
IBENP	Nonideal parasitic B-E saturation current	A	0
IKF	Forward knee current	A	0
IKP	Parasitic knee current	A	0
IKR	Reverse knee current	A	0
IS	Transport saturation current	A	1e-16
ISP	Parasitic transport saturation current	A	0
ISRR	Saturation current for reverse operation	–	1
ITF	Coefficient of $t_f$ dependence on $I_c$	–	0
KFN	B-E flicker (1/f) noise coefficient (unused)	–	0
MC	B-C grading coefficient	–	0.33
ME	B-E grading coefficient	–	0.33
MS	S-C grading coefficient	–	0.33
NBBE		–	1
NCI	Ideal B-C emission coefficient	–	1
NCIP	Ideal parasitic B-C emission coefficient	–	1
NCN	Non-ideal B-C emission coefficient	–	2
NCNP	Non-ideal parasitic B-C emission coefficient	–	2
NEI	Ideal B-E emission coefficient	–	1
NEN	Non-ideal B-E emission coefficient	–	2
NF	Forward emission coefficient	–	1
NFP	Parasitic forward emission coefficient	–	1
NKF		–	0.5
NR	Reverse emission coefficient	–	1

Table 2.45: VBIC 3T et cf v1.2 Device Model Parameters

Parameter	Description	Units	Default
PC	B-C built-in potential	–	0.75
PE	B-E built-in potential	–	0.75
PS	S-C built-in potential	–	0.75
QBM		–	0
QCO	Epi charge parameter	C	0
QTF	Variation of $t_f$ with base width modulation	–	0
RBI	Intrinsic base resistance	$\Omega$	0
RBP	Parasitic base resistance	$\Omega$	0
RBX	Extrinsic base resistance	$\Omega$	0
RCI	Intrinsic Collector resistance	$\Omega$	0
RCX	Extrinsic Collector resistance	$\Omega$	0
RE	Emitter resistance	$\Omega$	0
RS	Substrate resistance	$\Omega$	0
RTH	Thermal resistance, must be given for self-heating	$\Omega$	0
TAVC	Temperature coefficient of $A_{vc2}$	–	0
TD	Forward excess-phase delay time (unused in this version)	–	0
TF	Forward transit time	s	0
TNBBE		–	0
TNF	Temperature coefficient of $N_f$	–	0
TNOM	Nominal temperature	$^{\circ}\text{C}$	27
TR	Reverse transit time	–	0
TVBBE1		–	0
TVBBE2		–	0
VBBE		–	0
VEF	Forward Early voltage	V	0
VER	Reverse Early voltage	V	0
VERS	Version of this VBIC model	–	1.2
VO	Epi drift saturation voltage	V	0
VREV		–	0
VRT		–	0
VTF	Coefficient of $t_f$ dependence on $V_{bc}$	–	0
WBE	Portion of $I_{bei}$ from $V_{bei}$	–	1
WSP	Portion of $I_{ccp}$ from $V_{bep}$	–	1

Table 2.45: VBIC 3T et cf v1.2 Device Model Parameters

Parameter	Description	Units	Default
XII	Temperature exponent of Ibei, IbcI, Ibeip, and Ibcip	–	3
XIKF		–	0
XIN	Temperature exponent of Iben, Ibcn, Ibenp, and Ibcnp	–	3
XIS	Temperature exponent of IS	–	3
XISR	Temperature exponent of ISRR	–	0
XRBI		–	0
XRBP		–	0
XR BX		–	0
XRCI		–	0
XRCX		–	0
XRE	Temperature exponent of re	–	0
XRS	Temperature exponent of rs	–	0
XTF	Coefficient of tf with bias dependence	–	0
XVO	Temperature exponent of vo	–	0

### Level=23 instance parameters

Table 2.46 lists the parameters for the level 23 BJT (FBH HBT\_X model) available on the instance line.

Table 2.46: FBH HBT\_X v2.1 Device Instance Parameters

Parameter	Description	Units	Default
L	Length of emitter fingers	m	3e-05
N	Number of emitter fingers	–	1
TEMP	Device operating temperature	°C	25
W	Width of emitter fingers	m	3e-06

### Level=23 model parameters

Table 2.47: FBH HBT\_X v2.1 Device Model Parameters

Parameter	Description	Units	Default
AHC		–	0
BF		–	100
BR		–	1

Table 2.47: FBH HBT\_X v2.1 Device Model Parameters

Parameter	Description	Units	Default
BVCEO		—	0
BVEBO		—	0
CJC		—	1e-15
CJE		—	1e-15
CMIN		—	1e-16
CPB		—	0
CPC		—	0
CQ		—	0
CTH		—	7e-07
DEBUG		—	0
DEBUGPLUS		—	0
IKF		—	0
IKR		—	0
JO		—	0.001
JK		—	0.0004
JSC		—	0
JSE		—	0
JSEE		—	0
JSF		—	2e-23
JSR		—	2e-17
KBETA		—	0
KC		—	0
KJC		—	1
LB		—	0
LC		—	0
LE		—	0
MC		—	0
MJC		—	0.5
MJE		—	0.5
MODE		—	1
NC		—	0
NE		—	0
NEE		—	0
NF		—	1
NOISE		—	1

Table 2.47: FBH HBT\_X v2.1 Device Model Parameters

Parameter	Description	Units	Default
NR		—	1
RB		—	1
RB2		—	1
RBBXX		—	1e+06
RBXX		—	1e+06
RC		—	1
RCIO		—	0.001
RCXX		—	1e+06
RE		—	1
RJK		—	0.001
RTH		—	0.1
TF		—	1e-12
TFT		—	0
THCS		—	0
TNOM		—	20
TR		—	1e-15
TRX		—	1e-15
VAF		—	0
VAR		—	0
VCES		—	0.001
VG		—	1.3
VGB		—	0
VGBB		—	0
VGC		—	0
VGR		—	0
VJC		—	1.3
VJE		—	1.3
XCJC		—	0.5
XJO		—	1

## BJT Equations

The BJT implementation within **Xyce** is based on [10]. The equations in this section describe an NPN transistor. For the PNP device, reverse the signs of all voltages and currents. The equations

use the following variables:

$V_{be}$	=	intrinsic base-intrinsic emitter voltage
$V_{bc}$	=	intrinsic base-intrinsic collector voltage
$V_{bs}$	=	intrinsic base-substrate voltage
$V_{bw}$	=	intrinsic base-extrinsic collector voltage (quasi-saturation only)
$V_{bx}$	=	extrinsic base-intrinsic collector voltage
$V_{ce}$	=	intrinsic collector-intrinsic emitter voltage
$V_{js}$	=	(NPN) intrinsic collector-substrate voltage (PNP) intrinsic substrate-collector voltage
$V_t$	=	$kT/q$ (thermal voltage)
$V_{th}$	=	threshold voltage
$k$	=	Boltzmann's constant
$q$	=	electron charge
$T$	=	analysis temperature (K)
$T_0$	=	nominal temperature (set using TNOM option)

Other variables are listed above in BJT Model Parameters.

### DC Current

The BJT model is based on the Gummel and Poon model [11] where the different terminal currents are written

$$\begin{aligned}
 I_e &= -I_{cc} - I_{be} + I_{re} + (C_{dife} + C_{de}) \frac{dV_{be}}{dt} \\
 I_c &= -I_{cc} + I_{bc} - I_{rc} - (C_{difc} + C_{dc}) \frac{dV_{bc}}{dt} \\
 I_b &= I_e - I_c
 \end{aligned}$$

Here,  $C_{dife}$  and  $C_{difc}$  are the capacitances related to the hole charges per unit area in the base,  $Q_{dife}$  and  $Q_{difc}$ , affiliated with the electrons introduced across the emitter-base and collector-base junctions, respectively. Also,  $C_{be}$  and  $C_{bc}$  are the capacitances related to donations to the hole charge of the base,  $Q_{be}$  and  $Q_{bc}$ , affiliated with the differences in the depletion regions of the emitter-base and collector-base junctions, respectively. The intermediate currents used are defined as

$$\begin{aligned}
 -I_{be} &= \frac{IS}{BF} \left[ \exp \left( \frac{V_{be}}{N F V_{th}} \right) - 1 \right] \\
 -I_{cc} &= \frac{Q_{bo}}{Q_b} IS \left[ \exp \left( \frac{V_{be}}{N F V_{th}} \right) - \exp \left( \frac{V_{bc}}{N F V_{th}} \right) \right] \\
 -I_{bc} &= \frac{IS}{BR} \left[ \exp \left( \frac{V_{bc}}{N R V_{th}} \right) - 1 \right]
 \end{aligned}$$

$$I_{re} = \text{ISE} \left[ \exp \left( \frac{V_{be}}{\text{NE}V_{th}} \right) - 1 \right]$$

$$I_{rc} = \text{ISC} \left[ \exp \left( \frac{V_{bc}}{\text{NC}V_{th}} \right) - 1 \right]$$

where the last two terms are the generation/recombination currents related to the emitter and collector junctions, respectively. The charge  $Q_b$  is the majority carrier charge in the base at large injection levels and is a key difference in the Gummel-Poon model over the earlier Ebers-Moll model. The ratio  $Q_b/Q_{bo}$  (where  $Q_{bo}$  represents the zero-bias base charge, i.e. the value of  $Q_b$  when  $V_{be} = V_{bc} = 0$ ) as computed by **Xyce** is given by

$$\frac{Q_b}{Q_{bo}} = \frac{q_1}{2} \left( 1 + \sqrt{1 + 4q_2} \right)$$

where

$$q_1 = \left( 1 - \frac{V_{be}}{\text{VAR}} - \frac{V_{bc}}{\text{VAF}} \right)^{-1}$$

$$q_2 = \frac{\text{IS}}{\text{IKF}} \left[ \exp \left( \frac{V_{be}}{\text{NF}V_{th}} \right) - 1 \right] + \frac{\text{IS}}{\text{IKR}} \left[ \exp \left( \frac{V_{bc}}{\text{NR}V_{th}} \right) - 1 \right]$$

### Capacitance Terms

The capacitances listed in the above DC  $I - V$  equations each consist of a depletion layer capacitance  $C_d$  and a diffusion capacitance  $C_{dif}$ . The first is given by

$$C_d = \begin{cases} \text{CJ} \left( 1 - \frac{V_{di}}{\text{VJ}} \right)^{-\text{M}} & V_{di} \leq \text{FC} \cdot \text{VJ} \\ \text{CJ} (1 - \text{FC})^{-(1+\text{M})} \left[ 1 - \text{FC}(1 + \text{M}) + \text{M} \frac{V_{di}}{\text{VJ}} \right] & V_{di} > \text{FC} \cdot \text{VJ} \end{cases}$$

where  $\text{CJ} = \text{CJE}$  for  $C_{de}$ , and where  $\text{CJ} = \text{CJC}$  for  $C_{dc}$ . The diffusion capacitance (sometimes referred to as the transit time capacitance) is

$$C_{dif} = \text{TT}G_d = \text{TT} \frac{dI}{dV_{di}}$$

where  $I$  is the diode DC current given,  $G_d$  is the corresponding junction conductance, and where  $\text{TT} = \text{TF}$  for  $C_{dif_e}$  and  $\text{TT} = \text{TR}$  for  $C_{dif_c}$ .

### Temperature Effects

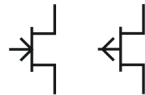
SPICE temperature effects are default, but all levels of the BJT have a more advanced temperature compensation available. By specifying `TEMPMODEL=QUADRATIC` in the netlist, parameters can be

interpolated quadratically between measured values extracted from data. In the BJT, IS and ISE are interpolated logarithmically because they can change over an order of magnitude or more for temperature ranges of interest. See the Section 2.1.14 for more details on how to include quadratic temperature effects.

For further information on BJT models, see [11]. For a thorough description of the U.C. Berkeley SPICE models see Reference [12].

## 2.3.15 Junction Field-Effect Transistor (JFET)

### Symbol



---

<b>Instance Form</b>	J<name> <drain node> <gate node> <source node> <model name> + [area value] [device parameters]
----------------------	--

---

<b>Examples</b>	JIN 100 1 0 JFAST J13 22 14 23 JNOM 2.0 J1 1 2 0 2N5114
-----------------	---

---

<b>Model Form</b>	.MODEL <model name> NJF [model parameters] .MODEL <model name> PJF [model parameters]
-------------------	--

---

### Parameters and Options

drain node	Node connected to drain.
gate node	Node connected to gate.
source node	Node connected to source.
source node	Name of model defined in .MODEL line.
area value	The JFET is modeled as an intrinsic FET using an ohmic resistance ( $R_D/\text{area}$ ) in series with the drain and another ohmic resistance ( $R_S/\text{area}$ ) in series with the source. area is an area factor with a default of 1.
device parameters	Parameters listed in Table 2.48 may be provided as space separated <parameter>=<value> specifications as needed. Any number of parameters may be specified.

---

<b>Comments</b>	The JFET was first proposed and analyzed by Shockley. The SPICE-compatible JFET model is an approximation to the Shockley analysis that employs an adjustable parameter B. Both the Shockley formulation and the SPICE approximation are available in Xyce.
-----------------	---

## Device Parameters

Table 2.48: JFET Device Instance Parameters

Parameter	Description	Units	Default
AREA	Device area	m <sup>2</sup>	1
TEMP	Device temperature	–	Ambient Temperature

## Model Parameters

Table 2.49: JFET Device Model Parameters

Parameter	Description	Units	Default
AF	Flicker noise exponent	–	1
B	Doping tail parameter (level 1)	V <sup>-1</sup>	1
BETA	Transconductance parameter	A/V <sup>2</sup>	0.0001
CGD	Zero-bias gate-drain junction capacitance	F	0
CGS	Zero-bias gate-source junction capacitance	F	0
DELTA	Saturation voltage parameter (level 2)	V	0
FC	Coefficient for forward-bias depletion capacitance	F	0.5
IS	Gate junction saturation current	A	1e-14
KF	Flicker noise coefficient	–	0.05
LAMBDA	Channel length modulation	V <sup>-1</sup>	0
PB	Gate junction potential	V	1
RD	Drain ohmic resistance	Ω	0
RS	Source ohmic resistance	Ω	0
TEMPMODEL	Specifies the type of parameter interpolation over temperature	–	'NONE'
THETA	Mobility modulation parameter (level 2)	V <sup>-1</sup>	0
TNOM		–	Ambient Temperature
VTO	Threshold voltage	V	-2

## Device Parameters

Table 2.50: JFET Device Instance Parameters

Parameter	Description	Units	Default
AREA	Device area	m <sup>2</sup>	1
TEMP	Device temperature	–	Ambient Temperature

## Model Parameters

Table 2.51: JFET Device Model Parameters

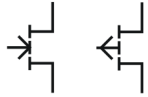
Parameter	Description	Units	Default
AF	Flicker noise exponent	–	1
B	Doping tail parameter (level 1)	V <sup>-1</sup>	1
BETA	Transconductance parameter	A/V <sup>2</sup>	0.0001
CGD	Zero-bias gate-drain junction capacitance	F	0
CGS	Zero-bias gate-source junction capacitance	F	0
DELTA	Saturation voltage parameter (level 2)	V	0
FC	Coefficient for forward-bias depletion capacitance	F	0.5
IS	Gate junction saturation current	A	1e-14
KF	Flicker noise coefficient	–	0.05
LAMBDA	Channel length modulation	V <sup>-1</sup>	0
PB	Gate junction potential	V	1
RD	Drain ohmic resistance	Ω	0
RS	Source ohmic resistance	Ω	0
TEMPMODEL	Specifies the type of parameter interpolation over temperature	–	'NONE'
THETA	Mobility modulation parameter (level 2)	V <sup>-1</sup>	0
TNOM		–	Ambient Temperature
VTO	Threshold voltage	V	-2

## JFET Level selection

**Xyce** supports two JFET models. LEVEL=1, the default, is the SPICE 3f5 treatment. This model employs a doping profile parameter B. When B=1, the original SPICE square law is exactly implemented, and when B=0.6 the model is close to that of Shockley.

When LEVEL=2 is selected, the Shockley model is used with some additional physics effects: channel length modulation and the effect of gate electric field on mobility. An additional parameter, DELTA, is added to the LEVEL 2 model that allows the user to adjust the saturation voltage.

## 2.3.16 Metal-Semiconductor FET (MESFET)

Symbol	
Instance Form	<code>Z&lt;name&gt; &lt; drain node&gt; &lt;gate node&gt; &lt;source node&gt; &lt;model name&gt;</code> <code>+ [area value] [device parameters]</code>
Model Form	<code>.MODEL &lt;model name&gt; NMF [model parameters]</code> <code>.MODEL &lt;model name&gt; PMF [model parameters]</code>
Examples	<code>Z1 2 3 0 MESMOD AREA=1.4</code> <code>Z1 7 2 3 ZM1</code>
Parameters and Options	<p><code>drain node</code> Node connected to drain.</p> <p><code>gate node</code> Node connected to gate.</p> <p><code>source node</code> Node connected to source.</p> <p><code>source node</code> Name of model defined in .MODEL line.</p> <p><code>area value</code> The MESFET is modeled as an intrinsic FET using an ohmic resistance (<math>RD/area</math>) in series with the drain and another ohmic resistance (<math>RS/area</math>) in series with the source. <code>area value</code> is a scaling factor with a default of 1.</p> <p><code>device parameters</code> Parameters listed in Table 2.52 may be provided as space separated <code>&lt;parameter&gt;=&lt;value&gt;</code> specifications as needed. Any number of parameters may be specified.</p>
Comments	Although MESFETs can be made of Si, such devices are not as common as GaAs MESFETS. And since the mobility of electrons is much higher than holes in GaAs, nearly all commercial devices are n-type MESFETS.

## Device Parameters

Table 2.52: MESFET Device Instance Parameters

Parameter	Description	Units	Default
AREA	device area	m <sup>2</sup>	1
TEMP	Device temperature	–	Ambient Temperature

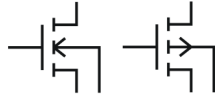
## Model Parameters

Table 2.53: MESFET Device Model Parameters

Parameter	Description	Units	Default
AF	Flicker noise exponent	–	1
ALPHA	Saturation voltage parameter	V <sup>-1</sup>	2
B	Doping tail parameter	V <sup>-1</sup>	0.3
BETA	Transconductance parameter	A/V <sup>2</sup>	0.0025
CGD	Zero-bias gate-drain junction capacitance	F	0
CGS	Zero-bias gate-source junction capacitance	F	0
FC	Coefficient for forward-bias depletion capacitance	F	0.5
IS	Gate junction saturation current	A	1e-14
KF	Flicker noise coefficient	–	0.05
LAMBDA	Channel length modulation	V <sup>-1</sup>	0
PB	Gate junction potential	V	1
RD	Drain ohmic resistance	Ω	0
RS	Source ohmic resistance	Ω	0
TEMPMODEL	Specifies the type of parameter interpolation over temperature	–	'NONE'
TNOM		–	Ambient Temperature
VTO	Threshold voltage	V	0

## 2.3.17 MOS Field Effect Transistor (MOSFET)

Symbol



Instance Form

```
M<name> <drain node> <gate node> <source node>
+ <bulk/substrate node> <model name>
+ [L=<value>] [W=<value>]
+ [AD=<value>] [AS=<value>]
+ [PD=<value>] [PS=<value>]
+ [NRD=<value>] [NRS=<value>]
+ [M=<value>] [IC=<value, ...>]
```

Special Form  
(BSIMSOI)

```
M<name> <drain node> <gate node> <source node>
+ <substrate node (E)>
+ [<External body contact (P)>]
+ [<internal body contact (B)>]
+ [<temperature node (T)>]
+ <model name>
+ [L=<value>] [W=<value>]
+ [AD=<value>] [AS=<value>]
+ [PD=<value>] [PS=<value>]
+ [NRD=<value>] [NRS=<value>] [NRB=<value>]
+ [BJTOFF=<value>]
+ [IC=<val>,<val>,<val>,<val>,<val>]
+ [RTH0=<val>] [CTH0=<val>]
+ [NBC=<val>] [NSEG=<val>] [PDBCP=<val>] [PSBCP=<val>]
+ [AGBCP=<val>] [AEBCP=<val>] [VBSUSR=<val>] [TNODEOUT]
+ [FRBODY=<val>] [M=<value>]
```

Model Form

```
.MODEL <model name> NMOS [model parameters]
.MODEL <model name> PMOS [model parameters]
```

Examples

```
M5 4 12 3 0 PNOM L=20u W=10u
M3 5 13 10 0 PSTRONG
M6 7 13 10 0 PSTRONG M=2
M8 10 12 100 100 NWEAK L=30u W=20u
+ AD=288p AS=288p PD=60u PS=60u NRD=14 NRS=24
```

Parameters and  
Options

L The MOSFET channel length and width that are decreased to get the actual channel length and width. They may be given in the device

.MODEL or .OPTIONS statements. The value in the device statement overrides the value in the model statement, which overrides the value in the .OPTIONS statement. If L or W values are not given, their default value is 100  $\mu\text{m}$ .

AD

AS The drain and source diffusion areas. Defaults for AD and AS can be set in the .OPTIONS statement. If AD or AS defaults are not set, their default value is 0.

PD

PS The drain and source diffusion perimeters. Their default value is 0.

NRD

NRS Multipliers (in units of  $\square$ ) that can be multiplied by RSH to yield the parasitic (ohmic) resistances of the drain (RD) and source (RS), respectively. NRD, NRS default to 0.

Consider a square sheet of resistive material. Analysis shows that the resistance between two parallel edges of such a sheet depends upon its composition and thickness, but is independent of its size as long as it is square. In other words, the resistance will be the same whether the square's edge is 2 mm, 2 cm, or 2 m. For this reason, the *sheet resistance* of such a layer, abbreviated RSH, has units of Ohms per square, written  $\Omega/\square$ .

M If specified, the value is used as a number of parallel MOSFETs to be simulated. For example, if M=2 is specified, **Xyce** simulates two identical mosfets connected to the same nodes in parallel.

IC The BSIM3 (model level 9), BSIM4 (model level 14) and BSIMSOI (model level 10) allow one to specify the initial voltage difference across nodes of the device during the DC operating point calculation. For the BSIM3 and BSIM4 the syntax is  $IC=V_{ds}, V_{gs}, V_{bs}$  where  $V_{ds}$  is the voltage difference between the drain and source,  $V_{gs}$  is the voltage difference between the gate and source and  $V_{bs}$  is the voltage difference between the body and source. The BSIMSOI device's initial condition syntax is  $IC=V_{ds}, V_{gs}, V_{bs}, V_{es}, V_{ps}$  where the two extra terms are the voltage difference between the substrate and source, and the external body and source nodes respectively. Note that for any of these lists of voltage differences, fewer than the full number of options may be specified. For example,  $IC=5.0$  specifies an initial condition on  $V_{ds}$  but does not specify any initial conditions on the other nodes. Therefore, one cannot specify  $V_{gs}$  without specifying  $V_{ds}$ , etc.

It is illegal to specify initial conditions on any nodes that are tied together. **Xyce** attempts to catch such errors, but complex circuits may stymie this error trap.

---

## BSIMSOI Options

There are a large number of extra instance parameters and optional nodes available for the BSIMSOI (level 10) MOSFET.

#### substrate node

The fourth node of the BSIMSOI device is always the substrate node, which is referred to as the E node.

#### external body contact node

If given, the fifth node is the external body contact node, P. It is connected to the internal body node through a body tie resistor. If P is not given, the internal body node is not accessible from the netlist and floats.

If there are only five nodes specified and TNODEOUT is also specified, the fifth node is the temperature node instead.

#### internal body contact node

If given, the sixth node is the internal body contact node, B. It is connected to the external body node through a body tie resistor. If B is not given and P is given, the internal body node is not accessible from the netlist, but is still tied to the external body contact through the tie resistance.

If there are only six nodes specified and TNODEOUT is also specified, the sixth node is the temperature node instead.

#### temperature node

If the parameter TNODEOUT is specified, the final node (fifth, sixth, or seventh) is interpreted as a temperature node. The temperature node is intended for thermal coupling simulation.

#### BJTOFF

Turns off the parasitic BJT currents.

IC The IC parameter allows specification of the five junction initial conditions,  $V_{ds}$ ,  $V_{gs}$ ,  $V_{bs}$ ,  $V_{es}$  and  $V_{ps}$ .  $V_{ps}$  is ignored in a four-terminal device.

#### RTH0

Thermal resistance per unit width. Taken from model card if not given.

#### CTH0

Thermal capacitance per unit width. Taken from model card if not given.

NBC Number of body contact isolation edges.

#### NSEG

Number of segments for channel width partitioning.

#### PDBCP

Parasitic perimeter length for body contact at drain side.

#### PSBCP

Parasitic perimeter length for body contact at source side.

#### AGBCP

Parasitic gate-to-body overlap area for body contact.

AEBCP	Parasitic body-to-substrate overlap area for body contact.
VBSUSR	Optional initial value of VBS specified by user for use in transient analysis. (unused in <b>Xyce</b> ).
FRBODY	Layout-dependent body resistance coefficient.

---

**Comments**      The simulator provides three MOSFET device models, which differ in the formulation of the I-V characteristic. The **LEVEL** parameter selects among different models as shown below.

## MOSFET Operating Temperature

Model parameters may be assigned unique measurement temperatures using the **TNOM** model parameter. See the MOSFET model parameters for more information.

## Instance Parameters

Tables 2.54, 2.56, 2.58, 2.60, 2.62 and 2.64 give the available instance parameters for the levels 1,2,3,6,9 and 10 MOSFETs, respectively.

In addition to the parameters shown in the tables, where a list of numbered initial condition parameters are shown, the MOSFETs support a vector parameter for the initial conditions. **IC1** and **IC2** may therefore be specified compactly as **IC=<ic1>,<ic2>**.

## Model Parameters

Tables 2.55, 2.57, 2.59, 2.61, 2.63, and 2.65 give the available model parameters for the levels 1,2,3,6,9 and 10 MOSFETs, respectively.

For a thorough description of MOSFET models see [12, 13, 14, 15, 16, 17, 18, 19, 20, 21].

### All MOSFET models

The parameters shared by all MOSFET model levels are principally parasitic element values (e.g., series resistance, overlap capacitance, etc.).

### Model levels 1 and 3

The DC behaviors of the level 1 and 3 MOSFET models are defined by the parameters **VTO**, **KP**, **LAMBDA**, **PHI**, and **GAMMA**. The simulator calculates these if the process parameters (e.g., **TOX**, and **NSUB**) are specified, but these are always overridden by any user-defined values. The **VTO** value is positive (negative) for modeling the enhancement mode and negative (positive) for the depletion mode of N-channel (P-channel) devices.

For MOSFETs, the capacitance model enforces charge conservation, influencing just the Level 1 and 3 models.

Effective device parameter lengths and widths are calculated as follows:

$$P_i = P_0 + P_L/L_e + P_W/W_e$$

where

$$\begin{aligned} L_e &= \text{effective length} = L - (2 \cdot LD) \\ W_e &= \text{effective width} = W - (2 \cdot WD) \end{aligned}$$

See **.MODEL** (model definition) for more information.

#### Model level 9 (BSIM3 version 3.2.2)

The University of California, Berkeley BSIM3 model is a physical-based model with a large number of dependencies on essential dimensional and processing parameters. It incorporates the key effects that are critical in modeling deep-submicrometer MOSFETs. These include threshold voltage reduction, nonuniform doping, mobility reduction due to the vertical field, bulk charge effect, carrier velocity saturation, drain-induced barrier lowering (DIBL), channel length modulation (CLM), hot-carrier-induced output resistance reduction, subthreshold conduction, source/drain parasitic resistance, substrate current induced body effect (SCBE) and drain voltage reduction in LDD structure.

The BSIM3 Version 3.2.2 model is a deep submicron MOSFET model with several major enhancements over earlier versions. These include a single I-V formula used to define the current and output conductance for operating regions, improved narrow width device modeling, a superior capacitance model with improved short and narrow geometry models, a new relaxation-time model to better transient modeling and enhanced model fitting of assorted W/L ratios using a single parameter set. This version preserves the large number of integrated dependencies on dimensional and processing parameters of the Version 2 model. For further information, see Reference [13].

#### Additional notes

1. If any of the following BSIM3 3.2.2 model parameters are not specified, they are computed via the following:

If **VTHO** is not specified, then:

$$\mathbf{VTHO} = \mathbf{VFB} + \phi_s \mathbf{K1} \sqrt{\phi_s}$$

where:

$$\mathbf{VFB} = -1.0$$

If **VTHO** is given, then:

$$\mathbf{VFB} = \mathbf{VTHO} - \phi_s + \mathbf{K1} \sqrt{\phi_s}$$

$$\begin{aligned}\mathbf{VBX} &= \phi_s - \frac{q \cdot \mathbf{NCH} \cdot \mathbf{XT}^2}{2\varepsilon_{si}} \\ \mathbf{CF} &= \left( \frac{2\varepsilon_{ox}}{\pi} \right) \ln \left( 1 + \frac{1}{4 \times 10^7 \cdot \mathbf{TOX}} \right)\end{aligned}$$

where:

$$E_g(T) = \text{the energy bandgap at temperature } T = 1.16 - \frac{T^2}{7.02 \times 10^4(T + 1108)}$$

2. If **K1** and **K2** are not given then they are computed via the following:

$$\begin{aligned}\mathbf{K1} &= \mathbf{GAMMA2} - 2 \cdot \mathbf{K2} \sqrt{\phi_s - \mathbf{VBM}} \\ \mathbf{K2} &= \frac{(\mathbf{GAMMA1} - \mathbf{GAMMA2})(\sqrt{\phi_s - \mathbf{VBX}} - \sqrt{\phi_s})}{2\sqrt{\phi_s}(\sqrt{\phi_s - \mathbf{VBM}} - \sqrt{\phi_s}) + \mathbf{VBM}}\end{aligned}$$

where:

$$\begin{aligned}\phi_s &= 2V_t \ln \left( \frac{\mathbf{NCH}}{n_i} \right) \\ V_t &= kT/q \\ n_i &= 1.45 \times 10^{10} \left( \frac{T}{300.15} \right)^{1.5} \exp \left( 21.5565981 - \frac{E_g(T)}{2V_t} \right)\end{aligned}$$

3. If **NCH** is not specified and **GAMMA1** is, then:

$$\mathbf{NCH} = \frac{\mathbf{GAMMA1}^2 \times \mathbf{COX}^2}{2q\varepsilon_{si}}$$

If **GAMMA1** and **NCH** are *not* specified, then **NCH** defaults to  $1.7 \times 10^{23} \text{ m}^{-3}$  and **GAMMA1** is computed using **NCH**:

$$\mathbf{GAMMA1} = \frac{\sqrt{2q\varepsilon_{si} \cdot \mathbf{NCH}}}{\mathbf{COX}}$$

If **GAMMA2** is not specified, then:

$$\mathbf{GAMMA2} = \frac{\sqrt{2q\varepsilon_{si} \cdot \mathbf{NSUB}}}{\mathbf{COX}}$$

4. If **CGSO** is not specified and **DLC** > 0, then:

$$\mathbf{CGSO} = \begin{cases} 0, & ((\mathbf{DLC} \cdot \mathbf{COX}) - \mathbf{CGSL}) < 0 \\ 0.6 \cdot \mathbf{XJ} \cdot \mathbf{COX}, & ((\mathbf{DLC} \cdot \mathbf{COX}) - \mathbf{CGSL}) \geq 0 \end{cases}$$

5. If **CGDO** is not specified and **DLC** > 0, then:

$$\mathbf{CGDO} = \begin{cases} 0, & ((\mathbf{DLC} \cdot \mathbf{COX}) - \mathbf{CGSL}) < 0 \\ 0.6 \cdot \mathbf{XJ} \cdot \mathbf{COX}, & ((\mathbf{DLC} \cdot \mathbf{COX}) - \mathbf{CGSL}) \geq 0 \end{cases}$$

## Model level 10 (BSIMSOI version 3.2)

The BSIMSOI is an international standard model for SOI (silicon on insulator) circuit design and is formulated on top of the BSIM3v3 framework. A detailed description can be found in the BSIMSOI 3.1 User's Manual [22] and the BSIMSOI 3.2 release notes [23].

This version (v3.2) of the BSIMSOI includes three depletion models; the partially depleted BSIM-SOI PD ( $\text{soiMod}=0$ ), the fully depleted BSIMSOI FD ( $\text{soiMod}=2$ ), and the unified SOI model ( $\text{soiMod}=1$ ).

BSIMPD is the Partial-Depletion (PD) mode of the BSIMSOI. A typical PD SOI MOSFET is formed on a thin SOI film which is layered on top of a buried oxide. BSIMPD has the following features and enhancements:

- Real floating body simulation of both I-V and C-V. The body potential is determined by the balance of all body current components.
- An improved parasitic bipolar current model. This includes enhancements in the various diode leakage components, second order effects (high-level injection and Early effect), diffusion charge equation, and temperature dependence of the diode junction capacitance.
- An improved impact-ionization current model. The contribution from BJT current is also modeled by the parameter  $F_{bji}$ .
- A gate-to-body tunneling current model, which is important to thin-oxide SOI technologies.
- Enhancements in the threshold voltage and bulk charge formulation of the high positive body bias regime.
- Instance parameters ( $P_{dbc}$ ,  $P_{sbc}$ ,  $A_{gbc}$ ,  $A_{ebc}$ ,  $N_{bc}$ ) are provided to model the parasitics of devices with various body-contact and isolation structures.
- An external body node (the 6th node) and other improvements are introduced to facilitate the modeling of distributed body resistance.
- Self heating. An external temperature node (the 7th node) is supported to facilitate the simulation of thermal coupling among neighboring devices.
- A unique SOI low frequency noise model, including a new excess noise resulting from the floating body effect.
- Width dependence of the body effect is modeled by parameters ( $K1, K1w1, K1w2$ ).
- Improved history dependence of the body charges with two new parameters ( $F_{body}$ ,  $DLCB$ ).
- An instance parameter  $V_{bsur}$  is provided for users to set the transient initial condition of the body potential.
- The new charge-thickness capacitance model introduced in BSIM3v3.2,  $\text{capMod}=3$ , is included.

## Quadratic Temperature Compensation

SPICE temperature effects are the default, but MOSFET levels 18, 19 and 20 have a more advanced temperature compensation available. By specifying `TEMPMODEL=QUADRATIC` in the netlist, parameters can be interpolated quadratically between measured values extracted from data. See Section 2.1.14 for more details.

## MOSFET Equations

The following equations define an N-channel MOSFET. The P-channel devices use a reverse the sign for all voltages and currents. The equations use the following variables:

$V_{bs}$	=	intrinsic substrate-intrinsic source voltage
$V_{bd}$	=	intrinsic substrate-intrinsic drain voltage
$V_{ds}$	=	intrinsic drain-substrate source voltage
$V_{dsat}$	=	saturation voltage
$V_{gs}$	=	intrinsic gate-intrinsic source voltage
$V_{gd}$	=	intrinsic gate-intrinsic drain voltage
$V_t$	=	$kT/q$ (thermal voltage)
$V_{th}$	=	threshold voltage
$C_{ox}$	=	the gate oxide capacitance per unit area
$f$	=	noise frequency
$k$	=	Boltzmann's constant
$q$	=	electron charge
$L_{eff}$	=	effective channel length
$W_{eff}$	=	effective channel width
$T$	=	analysis temperature (K)
$T_0$	=	nominal temperature (set using TNOM option)

Other variables are listed in the BJT Equations section 2.3.14.

## All Levels

$$\begin{aligned}
 I_g &= \text{gate current} = 0 \\
 I_b &= \text{bulk current} = I_{bs} + I_{bd} \\
 \text{where} \\
 I_{bs} &= \text{bulk-source leakage current} = I_{ss} \left( e^{V_{bs}/(NV_t)} - 1 \right) \\
 I_{ds} &= \text{bulk-drain leakage current} = I_{ds} \left( e^{V_{bd}/(NV_t)} - 1 \right) \\
 \text{where} \\
 \text{if} \\
 \mathbf{JS} = 0, \text{ or } \mathbf{AS} = 0 \text{ or } \mathbf{AD} = 0 \\
 \text{then} \\
 I_{ss} &= \mathbf{IS} \\
 I_{ds} &= \mathbf{IS} \\
 \text{else} \\
 I_{ss} &= \mathbf{AS} \times \mathbf{JS} + \mathbf{PS} \times \mathbf{JSSW} \\
 I_{ds} &= \mathbf{AD} \times \mathbf{JS} + \mathbf{PD} \times \mathbf{JSSW} \\
 I_d &= \text{drain current} = I_{drain} - I_{bd} \\
 I_s &= \text{source current} = -I_{drain} - I_{bs}
 \end{aligned}$$

## Level 1: Idrain

**Normal Mode:**  $V_{ds} > 0$

### Case 1

For cutoff region:  $V_{gs} - V_{to} < 0$

$$I_{drain} = 0$$

### Case 2

For linear region:  $V_{ds} < V_{gs} - V_{to}$

$$I_{drain} = (W/L)(\mathbf{KN}/2)(1 + \mathbf{LAMBDA} \times V_{ds})V_{ds}(2(V_{gs} - V_{to}) - V_{ds})$$

### Case 3

For saturation region:  $0 \leq V_{gs} - V_{to} \leq V_{ds}$

$$I_{drain} = (W/L)(\mathbf{KN}/2)(1 + \mathbf{LAMBDA} \cdot V_{ds})(V_{gs} - V_{to})^2$$

where

$$V_{to} = \mathbf{VTO} + \mathbf{GAMMA} \cdot \left( (\mathbf{PHI} - V_{bs})^{1/2} \right)^{1/2}$$

**Inverted Mode:**  $V_{ds} < 0$

Here, simply switch the source and drain in the normal mode equations given above.

### Level 3: I<sub>drain</sub>

See Reference [16] below for detailed information.

## Capacitance

### Level 1 and 3

$C_{bs}$  = bulk-source capacitance = area cap. + sidewall cap. + transit time cap.

$C_{bd}$  = bulk-drain capacitance = area cap. + sidewall cap. + transit time cap.

where

*if*

$$\mathbf{CBS} = 0 \text{ and } \mathbf{CBD} = 0$$

*then*

$$\begin{aligned} C_{bs} &= \mathbf{AS} \cdot \mathbf{CJ} \cdot C_{bsj} + \mathbf{PS} \cdot \mathbf{CJSW} \cdot C_{bss} + \mathbf{TT} \cdot G_{bs} \\ C_{bd} &= \mathbf{AD} \cdot \mathbf{CJ} \cdot C_{bdj} + \mathbf{PD} \cdot \mathbf{CJSW} \cdot C_{bds} + \mathbf{TT} \cdot G_{ds} \end{aligned}$$

*else*

$$\begin{aligned} C_{bs} &= \mathbf{CBS} \cdot C_{bsj} + \mathbf{PS} \cdot \mathbf{CJSW} \cdot C_{bss} + \mathbf{TT} \cdot G_{bs} \\ C_{bd} &= \mathbf{CBD} \cdot C_{bdj} + \mathbf{PD} \cdot \mathbf{CJSW} \cdot C_{bds} + \mathbf{TT} \cdot G_{ds} \end{aligned}$$

*where*

$$\begin{aligned} G_{bs} &= \text{DC bulk-source conductance} = dI_{bs}/dV_{bs} \\ G_{bd} &= \text{DC bulk-drain conductance} = dI_{bd}/dV_{bd} \end{aligned}$$

*if*

$$V_{bs} \leq \mathbf{FC} \cdot \mathbf{PB}$$

*then*

$$\begin{aligned} C_{bsj} &= (1 - V_{bs}/\mathbf{PB})^{-\mathbf{MJ}} \\ C_{bss} &= (1 - V_{bs}/\mathbf{PBSW})^{-\mathbf{MJSW}} \end{aligned}$$

*if*

$$V_{bs} > \mathbf{FC} \cdot \mathbf{PB}$$

*then*

$$\begin{aligned} C_{bsj} &= (1 - \mathbf{FC})^{-(1+\mathbf{MJ})} (1 - \mathbf{FC}(1 + \mathbf{MJ}) + \mathbf{MJ} \cdot V_{bs}/\mathbf{PB}) \\ C_{bss} &= (1 - \mathbf{FC})^{-(1+\mathbf{MJSW})} (1 - \mathbf{FC}(1 + \mathbf{MJSW}) + \mathbf{MJSW} \cdot V_{bs}/\mathbf{PBSW}) \end{aligned}$$

*if*

$$V_{bd} \leq \mathbf{FC} \cdot \mathbf{PB}$$

*then*

$$\begin{aligned} C_{bdj} &= (1 - V_{bd}/\mathbf{PB})^{-\mathbf{MJ}} \\ C_{bds} &= (1 - V_{bd}/\mathbf{PBSW})^{-\mathbf{MJSW}} \end{aligned}$$

*if*

$$V_{bd} > \mathbf{FC} \cdot \mathbf{PB}$$

*then*

$$\begin{aligned} C_{bdj} &= (1 - \mathbf{FC})^{-(1+\mathbf{MJ})} (1 - \mathbf{FC}(1 + \mathbf{MJ}) + \mathbf{MJ} \cdot V_{bd}/\mathbf{PB}) \\ C_{bds} &= (1 - \mathbf{FC})^{-(1+\mathbf{MJSW})} (1 - \mathbf{FC}(1 + \mathbf{MJSW})) \end{aligned}$$

$$C_{gs} = \text{gate-source overlap capacitance} = \mathbf{CGSO} \cdot \mathbf{W}$$

$$C_{gd} = \text{gate-drain overlap capacitance} = \mathbf{CGDO} \cdot \mathbf{W}$$

$$C_{gb} = \text{gate-bulk overlap capacitance} = \mathbf{CGBO} \cdot \mathbf{L}$$

## Temperature Effects

### All Levels

$$\mathbf{IS}(T) = \mathbf{IS} \cdot \exp(E_g(T_0) \cdot T/T_0 - E_g(T)) / V_t$$

$$\mathbf{JS}(T) = \mathbf{JS} \cdot \exp(E_g(T_0) \cdot T/T_0 - E_g(T)) / V_t$$

$$\mathbf{JSSW}(T) = \mathbf{JSSW} \cdot \exp(E_g(T_0) \cdot T/T_0 - E_g(T)) / V_t$$

$$\mathbf{PB}(T) = \mathbf{PB} \cdot T/T_0 - 3V_t \ln(T/T_0) - E_g(T_0) \cdot T/T_0 + E_g T$$

$$\mathbf{PBSW}(T) = \mathbf{PBSW} \cdot T/T_0 - 3V_t \ln(T/T_0) - E_g(T_0) \cdot T/T_0 + E_g T$$

$$\mathbf{PHI}(T) = \mathbf{PHI} \cdot T/T_0 - 3V_t \ln(T/T_0) - E_g(T_0) \cdot T/T_0 + E_g T$$

where

$$E_g(T) = \text{silicon bandgap energy} = 1.16 - 0.000702T^2/(T + 1108)$$

$$\mathbf{CBD}(T) = \mathbf{CBD} \cdot (1 + \mathbf{MJ} \cdot (0.0004(T - T_0) + (1 - \mathbf{PB}(T)/\mathbf{PB})))$$

$$\mathbf{CBS}(T) = \mathbf{CBS} \cdot (1 + \mathbf{MJ} \cdot (0.0004(T - T_0) + (1 - \mathbf{PB}(T)/\mathbf{PB})))$$

$$\mathbf{CJ}(T) = \mathbf{CJ} \cdot (1 + \mathbf{MJ} \cdot (0.0004(T - T_0) + (1 - \mathbf{PB}(T)/\mathbf{PB})))$$

$$\mathbf{CJSW}(T) = \mathbf{CJSW} \cdot (1 + \mathbf{MJSW} \cdot (0.0004(T - T_0) + (1 - \mathbf{PB}(T)/\mathbf{PB})))$$

$$\mathbf{KP}(T) = \mathbf{KP} \cdot (T/T_0)^{-3/2}$$

$$\mathbf{UO}(T) = \mathbf{UO} \cdot (T/T_0)^{-3/2}$$

$$\mathbf{MUS}(T) = \mathbf{MUS} \cdot (T/T_0)^{-3/2}$$

$$\mathbf{MUZ}(T) = \mathbf{MUZ} \cdot (T/T_0)^{-3/2}$$

$$\mathbf{X3MS}(T) = \mathbf{X3MS} \cdot (T/T_0)^{-3/2}$$

## Level 1 MOSFET Tables

Table 2.54: MOSFET level 1 Device Instance Parameters

Parameter	Description	Units	Default
AD	Drain diffusion area	m <sup>2</sup>	0
AS	Source diffusion area	m <sup>2</sup>	0
IC1	Initial condition on Drain-Source voltage	V	0
IC2	Initial condition on Gate-Source voltage	V	0
IC3	Initial condition on Bulk-Source voltage	V	0
L	Channel length	m	0
M	Multiplier for M devices connected in parallel	–	1
NRD	Multiplier for RSH to yield parasitic resistance of drain	□	1
NRS	Multiplier for RSH to yield parasitic resistance of source	□	1
OFF	Initial condition of no voltage drops across device	logical (T/F)	false
PD	Drain diffusion perimeter	m	0
PS	Source diffusion perimeter	m	0
TEMP	Device temperature	°C	Ambient Temperature
W	Channel width	m	0

Table 2.55: MOSFET level 1 Device Model Parameters

Parameter	Description	Units	Default
AF	Flicker noise exponent	–	1
CBD	Zero-bias bulk-drain p-n capacitance	F	0
CBS	Zero-bias bulk-source p-n capacitance	F	0
CGB0	Gate-bulk overlap capacitance/channel length	F/m	0
CGD0	Gate-drain overlap capacitance/channel width	F/m	0
CGS0	Gate-source overlap capacitance/channel width	F/m	0
CJ	Bulk p-n zero-bias bottom capacitance/area	F/m <sup>2</sup>	0
CJSW	Bulk p-n zero-bias sidewall capacitance/area	F/m <sup>2</sup>	0
FC	Bulk p-n forward-bias capacitance coefficient	–	0.5
GAMMA	Bulk threshold parameter	V <sup>1/2</sup>	0
IS	Bulk p-n saturation current	A	1e-14
JS	Bulk p-n saturation current density	A/m <sup>2</sup>	0

Table 2.55: MOSFET level 1 Device Model Parameters

Parameter	Description	Units	Default
KF	Flicker noise coefficient	–	0
KP	Transconductance coefficient	$A/V^2$	2e-05
L	Default channel length	m	0.0001
LAMBDA	Channel-length modulation	$V^{-1}$	0
LD	Lateral diffusion length	m	0
MJ	Bulk p-n bottom grading coefficient	–	0.5
MJSW	Bulk p-n sidewall grading coefficient	–	0.5
NSS	Surface state density	$cm^{-2}$	0
NSUB	Substrate doping density	$cm^{-3}$	0
PB	Bulk p-n bottom potential	V	0.8
PHI	Surface potential	V	0.6
RD	Drain ohmic resistance	$\Omega$	0
RS	Source ohmic resistance	$\Omega$	0
RSH	Drain, source diffusion sheet resistance	$\Omega$	0
TEMPMODEL	Specifies the type of parameter interpolation over temperature	–	'NONE'
TNOM	Nominal device temperature	$^{\circ}C$	27
TOX	Gate oxide thickness	m	1e-07
TPG	Gate material type (-1 = same as substrate, 0 = aluminum, 1 = opposite of substrate)	–	0
U0	Surface mobility	$1/(Vcm^2s)$	600
U0	Surface mobility	$1/(Vcm^2s)$	600
VTO	Zero-bias threshold voltage	V	0
W	Default channel width	m	0.0001

## Level 2 MOSFET Tables (SPICE Level 2)

Table 2.56: MOSFET level 2 Device Instance Parameters

Parameter	Description	Units	Default
AD	Drain diffusion area	m <sup>2</sup>	0
AS	Source diffusion area	m <sup>2</sup>	0
IC1	Initial condition on Drain-Source voltage	V	0
IC2	Initial condition on Gate-Source voltage	V	0
IC3	Initial condition on Bulk-Source voltage	V	0
L	Channel length	m	0
M	Multiplier for M devices connected in parallel	–	1
NRD	Multiplier for RSH to yield parasitic resistance of drain	□	1
NRS	Multiplier for RSH to yield parasitic resistance of source	□	1
OFF	Initial condition of no voltage drops across device	logical (T/F)	false
PD	Drain diffusion perimeter	m	0
PS	Source diffusion perimeter	m	0
TEMP	Device temperature	°C	Ambient Temperature
W	Channel width	m	0

Table 2.57: MOSFET level 2 Device Model Parameters

Parameter	Description	Units	Default
AF	Flicker noise exponent	–	1
CBD	Zero-bias bulk-drain p-n capacitance	F	0
CBS	Zero-bias bulk-source p-n capacitance	F	0
CGBO	Gate-bulk overlap capacitance/channel length	F/m	0
CGDO	Gate-drain overlap capacitance/channel width	F/m	0
CGSO	Gate-source overlap capacitance/channel width	F/m	0
CJ	Bulk p-n zero-bias bottom capacitance/area	F/m <sup>2</sup>	0
CJSW	Bulk p-n zero-bias sidewall capacitance/area	F/m <sup>2</sup>	0
DELTA	Width effect on threshold	–	0
FC	Bulk p-n forward-bias capacitance coefficient	–	0.5
GAMMA	Bulk threshold parameter	V <sup>1/2</sup>	0
IS	Bulk p-n saturation current	A	1e-14

Table 2.57: MOSFET level 2 Device Model Parameters

Parameter	Description	Units	Default
JS	Bulk p-n saturation current density	A/m <sup>2</sup>	0
KF	Flicker noise coefficient	–	0
KP	Transconductance coefficient	A/V <sup>2</sup>	2e-05
L	Default channel length	m	0.0001
LAMBDA	Channel-length modulation	V <sup>-1</sup>	0
LD	Lateral diffusion length	m	0
MJ	Bulk p-n bottom grading coefficient	–	0.5
MJSW	Bulk p-n sidewall grading coefficient	–	0.5
NEFF	Total channel charge coeff.	–	1
NFS	Fast surface state density	–	0
NSS	Surface state density	cm <sup>-2</sup>	0
NSUB	Substrate doping density	cm <sup>-3</sup>	0
PB	Bulk p-n bottom potential	V	0.8
PHI	Surface potential	V	0.6
RD	Drain ohmic resistance	Ω	0
RS	Source ohmic resistance	Ω	0
RSH	Drain, source diffusion sheet resistance	Ω	0
TEMPMODEL	Specifies the type of parameter interpolation over temperature	–	'NONE'
TNOM	Nominal device temperature	°C	27
TOX	Gate oxide thickness	m	1e-07
TPG	Gate material type (-1 = same as substrate, 0 = aluminum, 1 = opposite of substrate)	–	0
U0	Surface mobility	1/(Vcm <sup>2</sup> s)	600
UCRIT	Crit. field for mob. degradation	–	10000
UEXP	Crit. field exp for mob. deg.	–	0
U0	Surface mobility	1/(Vcm <sup>2</sup> s)	600
VMAX	Maximum carrier drift velocity	–	0
VTO	Zero-bias threshold voltage	V	0
W	Default channel width	m	0.0001
XJ	Junction depth	–	0

## Level 3 MOSFET Tables

Table 2.58: MOSFET level 3 Device Instance Parameters

Parameter	Description	Units	Default
AD	Drain diffusion area	m <sup>2</sup>	0
AS	Source diffusion area	m <sup>2</sup>	0
IC1	Initial condition on Drain-Source voltage	V	0
IC2	Initial condition on Gate-Source voltage	V	0
IC3	Initial condition on Bulk-Source voltage	V	0
L	Channel length	m	0
M	Multiplier for M devices connected in parallel	–	1
NRD	Multiplier for RSH to yield parasitic resistance of drain	□	1
NRS	Multiplier for RSH to yield parasitic resistance of source	□	1
OFF	Initial condition of no voltage drops across device	logical (T/F)	false
PD	Drain diffusion perimeter	m	0
PS	Source diffusion perimeter	m	0
TEMP	Device temperature	°C	Ambient Temperature
W	Channel width	m	0

Table 2.59: MOSFET level 3 Device Model Parameters

Parameter	Description	Units	Default
AF	Flicker noise exponent	–	1
CBD	Zero-bias bulk-drain p-n capacitance	F	0
CBS	Zero-bias bulk-source p-n capacitance	F	0
CGB0	Gate-bulk overlap capacitance/channel length	F/m	0
CGD0	Gate-drain overlap capacitance/channel width	F/m	0
CGS0	Gate-source overlap capacitance/channel width	F/m	0
CJ	Bulk p-n zero-bias bottom capacitance/area	F/m <sup>2</sup>	0
CJSW	Bulk p-n zero-bias sidewall capacitance/area	F/m <sup>2</sup>	0
DELTA	Width effect on threshold	–	0
ETA	Static feedback	–	0
FC	Bulk p-n forward-bias capacitance coefficient	–	0.5
GAMMA	Bulk threshold parameter	V <sup>1/2</sup>	0

Table 2.59: MOSFET level 3 Device Model Parameters

Parameter	Description	Units	Default
IS	Bulk p-n saturation current	A	1e-14
JS	Bulk p-n saturation current density	A/m <sup>2</sup>	0
KAPPA	Saturation field factor	–	0.2
KF	Flicker noise coefficient	–	0
KP	Transconductance coefficient	A/V <sup>2</sup>	2e-05
L	Default channel length	m	0.0001
LD	Lateral diffusion length	m	0
MJ	Bulk p-n bottom grading coefficient	–	0.5
MJSW	Bulk p-n sidewall grading coefficient	–	0.33
NFS	Fast surface state density	cm <sup>-2</sup>	0
NSS	Surface state density	cm <sup>-2</sup>	0
NSUB	Substrate doping density	cm <sup>-3</sup>	0
PB	Bulk p-n bottom potential	V	0.8
PHI	Surface potential	V	0.6
RD	Drain ohmic resistance	Ω	0
RS	Source ohmic resistance	Ω	0
RSH	Drain, source diffusion sheet resistance	Ω	0
TEMPMODEL	Specifies the type of parameter interpolation over temperature	–	'NONE'
THETA	Mobility modulation	V <sup>-1</sup>	0
TNOM	Nominal device temperature	°C	27
TOX	Gate oxide thickness	m	1e-07
TPG	Gate material type (-1 = same as substrate, 0 = aluminum, 1 = opposite of substrate)	–	1
U0	Surface mobility	1/(Vcm <sup>2</sup> s)	600
U0	Surface mobility	1/(Vcm <sup>2</sup> s)	600
VMAX	Maximum drift velocity	m/s	0
VTO	Zero-bias threshold voltage	V	0
W	Default channel width	m	0.0001
XJ	Metallurgical junction depth	m	0

## Level 6 MOSFET Tables (SPICE Level 6)

Table 2.60: MOSFET level 6 Device Instance Parameters

Parameter	Description	Units	Default
AD	Drain diffusion area	m <sup>2</sup>	0
AS	Source diffusion area	m <sup>2</sup>	0
IC1	Initial condition on Drain-Source voltage	V	0
IC2	Initial condition on Gate-Source voltage	V	0
IC3	Initial condition on Bulk-Source voltage	V	0
L	Channel length	m	0
M	Multiplier for M devices connected in parallel	–	1
NRD	Multiplier for RSH to yield parasitic resistance of drain	□	1
NRS	Multiplier for RSH to yield parasitic resistance of source	□	1
OFF	Initial condition of no voltage drops across device	logical (T/F)	false
PD	Drain diffusion perimeter	m	0
PS	Source diffusion perimeter	m	0
TEMP	Device temperature	°C	Ambient Temperature
W	Channel width	m	0

Table 2.61: MOSFET level 6 Device Model Parameters

Parameter	Description	Units	Default
CBD	Zero-bias bulk-drain p-n capacitance	F	0
CBS	Zero-bias bulk-source p-n capacitance	F	0
CGBO	Gate-bulk overlap capacitance/channel length	F/m	0
CGDO	Gate-drain overlap capacitance/channel width	F/m	0
CGSO	Gate-source overlap capacitance/channel width	F/m	0
CJ	Bulk p-n zero-bias bottom capacitance/area	F/m <sup>2</sup>	0
CJSW	Bulk p-n zero-bias sidewall capacitance/area	F/m <sup>2</sup>	0
FC	Bulk p-n forward-bias capacitance coefficient	–	0.5
GAMMA	Bulk threshold parameter	–	0
GAMMA1	Bulk threshold parameter 1	–	0
IS	Bulk p-n saturation current	A	1e-14
JS	Bulk p-n saturation current density	A/m <sup>2</sup>	0

Table 2.61: MOSFET level 6 Device Model Parameters

Parameter	Description	Units	Default
KC	Saturation current factor	–	5e-05
KV	Saturation voltage factor	–	2
LAMBDA	Channel length modulation param.	–	0
LAMBDA0	Channel length modulation param. 0	–	0
LAMBDA1	Channel length modulation param. 1	–	0
LD	Lateral diffusion length	m	0
MJ	Bulk p-n bottom grading coefficient	–	0.5
MJSW	Bulk p-n sidewall grading coefficient	–	0.5
NC	Saturation current coeff.	–	1
NSS	Surface state density	cm <sup>-2</sup>	0
NSUB	Substrate doping density	cm <sup>-3</sup>	0
NV	Saturation voltage coeff.	–	0.5
NVTH	Threshold voltage coeff.	–	0.5
PB	Bulk p-n bottom potential	V	0.8
PHI	Surface potential	V	0.6
PS	Sat. current modification par.	–	0
RD	Drain ohmic resistance	Ω	0
RS	Source ohmic resistance	Ω	0
RSH	Drain, source diffusion sheet resistance	Ω	0
SIGMA	Static feedback effect par.	–	0
TEMPMODEL	Specifies the type of parameter interpolation over temperature	–	'NONE'
TNOM	Nominal device temperature	°C	27
TOX	Gate oxide thickness	m	1e-07
TPG	Gate material type (-1 = same as substrate, 0 = aluminum, 1 = opposite of substrate)	–	1
U0	Surface mobility	1/(Vcm <sup>2</sup> s)	600
U0	Surface mobility	1/(Vcm <sup>2</sup> s)	600
VTO	Zero-bias threshold voltage	V	0

## Level 9 MOSFET Tables (BSIM3)

For complete documentation of the BSIM3 model, see the users' manual for the BSIM3, available for download at <http://www-device.eecs.berkeley.edu/bsim/?page=BSIM3>. **Xyce** implements Version 3.2.2 of the BSIM3, you will have to get the documentation from the FTP archive on the Berkeley site.

In addition to the parameters shown in table 2.62, the BSIM3 supports a vector parameter for the initial conditions. IC1 through IC3 may therefore be specified compactly as IC=<ic1>,<ic2>,<ic3>.

**NOTE: Many BSIM3 parameters listed in tables 2.62 and 2.63 as having default values of zero are actually replaced with internally computed defaults if not given. Specifying zero in your model card will override this internal computation. It is recommended that you only set model parameters that you are actually changing from defaults and that you not generate model cards containing default values from the tables.**

Table 2.62: BSIM3 Device Instance Parameters

Parameter	Description	Units	Default
<b>Control Parameters</b>			
M	Multiplier for M devices connected in parallel	—	1
NQSMOD	Flag for NQS model	—	0
<b>Geometry Parameters</b>			
AD	Drain diffusion area	m <sup>2</sup>	0
AS	Source diffusion area	m <sup>2</sup>	0
L	Channel length	m	0
NRD	Multiplier for RSH to yield parasitic resistance of drain	□	1
NRS	Multiplier for RSH to yield parasitic resistance of source	□	1
PD	Drain diffusion perimeter	m	0
PS	Source diffusion perimeter	m	0
W	Channel width	m	0
<b>Temperature Parameters</b>			
TEMP	Device temperature	°C	Ambient Temperature
<b>Voltage Parameters</b>			
IC1	Initial condition on Vds	V	0
IC2	Initial condition on Vgs	V	0
IC3	Initial condition on Vbs	V	0
OFF	Initial condition of no voltage drops accross device	logical (T/F)	false

Table 2.63: BSIM3 Device Model Parameters

Parameter	Description	Units	Default
<b><i>Bin Parameters</i></b>			
LMAX	Maximum channel length	m	1
LMIN	Minimum channel length	m	0
WMAX	Maximum channel width	m	1
WMIN	Minimum channel width	m	0
<b><i>Capacitance Parameters</i></b>			
ACDE	Exponential coefficient for charge thickness in capmod = 3 for accumulation and depletion regions	m/V	1
CF	Firing field capacitance	F/m	0
CGBO	Gate-bulk overlap capacitance per unit channel length	F/m	0
CGDL	Light-doped drain-gate region overlap capacitance	F/m	0
CGDO	Non-LLD region drain-gate overlap capacitance per unit channel length	F/m	0
CGSL	Light-doped source-gate region overlap capacitance	F/m	0
CGSO	Non-LLD region source-gate overlap capacitance per unit channel length	F/m	0
CJ	Bulk p-n zero-bias bottom capacitance/area	F/m <sup>2</sup>	0.0005
CJSW	Bulk p-n zero-bias sidewall capacitance/area	F/m <sup>2</sup>	5e-10
CJSWG	Source/grain gate sidewall junction capacitance per unit width	F/m	0
CKAPPA	Coefficient for lightly doped region overlap capacitance firing field capacitance	F/m	0.6
CLC	Constant term for short-channel model	m	1e-07
CLE	Exponential term for the short-channel model	–	0.6
DLC	Length offset fitting parameter from C-V	m	0
DWC	Width offset fitting parameter from C-V	m	0
MJSWG	Source/grain gate sidewall junction capacitance grading coefficient	–	0
MOIN	Coefficient for the gate-bias dependent surface potential	–	15
NOFF	CV parameter in Vgsteff, CV for weak to strong inversion	–	1
PBSW	Source/drain side junction built-in potential	V	1
PBSWG	Source/drain gate sidewall junction built-in potential	V	0

Table 2.63: BSIM3 Device Model Parameters

Parameter	Description	Units	Default
VFBCV	Flat-band voltage parameter (for CAPMOD = 0 only)	V	-1
VOFFCV	CV parameter in Vgsteff, CV for weak to strong inversion	V	0
XPART	Charge partitioning rate flag	–	0
<b>Control Parameters</b>			
BINUNIT	Binning unit selector	–	1
CAPMOD	Flag for capacitance models	–	3
MOBMOD	Mobility model selector	–	1
NOIMOD	Flag for noise models	–	1
PARAMCHK	Parameter value check	–	0
VERSION	Version number	–	'3.2.2'
<b>DC Parameters</b>			
A0	Bulk charge effect coefficient for channel length	–	1
A1	First non-saturation effect parameter	$V^{-1}$	0
A2	Second non-saturation factor	–	1
AGS	Gate-bias coefficient of abulk	$V^{-1}$	0
ALPHA0	First parameter of impact-ionization current	m/V	0
ALPHA1	Isb parameter for length scaling	$V^{-1}$	0
B0	Bulk charge effect coefficient for channel width	m	0
B1	Bulk charge effect offset	m	0
BETA0	Second parameter of impact-ionization current	V	30
CDSC	Drain/source to channel coupling capacitance	F/m <sup>2</sup>	0.00024
CDSCB	Body-bias sensitivity of CDSC	F/(Vm <sup>2</sup> )	0
CDSCD	Drain-bias sensitivity of CDSC	F/(Vm <sup>2</sup> )	0
CIT	Interface trap capacitance	F/m <sup>2</sup>	0
DELTA	Effective Vds parameter	V	0.01
DROUT	L-depedance Coefficient of the DIBL correction parameter in Rout	–	0.56
DSUB	DIBL coefficient exponent in subthreshold region	–	0
DVT0	First coefficient of short-channel effect effect on threshold voltage	–	2.2
DVTOW	First coefficient of narrow-width effect effect on threshold voltage for small channel length	m <sup>-1</sup>	0
DVT1	Second coefficient of short-channel effect effect on threshold voltage	–	0.53

Table 2.63: BSIM3 Device Model Parameters

Parameter	Description	Units	Default
DVT1W	Second coefficient of narrow-width effect effect on threshold voltage for small channel length	$m^{-1}$	5.3e+06
DVT2	Body-bias coefficient of short-channel effect effect on threshold voltage	$V^{-1}$	-0.032
DVT2W	Body-bias coefficient of narrow-width effect effect on threshold voltage for small channel length	$V^{-1}$	-0.032
DWB	Coefficient of substrate body bias dependence of $W_{eff}$	$m/V^{1/2}$	0
DWG	Coefficient of gate dependence of $W_{eff}$	$m/V^{1/2}$	0
ETA0	DIBL coefficient in subthreshold region	—	0.08
ETAB	Body-bias coefficient for the subthreshold DIBL effect	$V^{-1}$	-0.07
IJTH	Diode limiting current	A	0.1
JSW	Sidewall saturation current per unit length	A/m	0
K1	First-order body effect coefficient	$V^{1/2}$	0
K2	second-order body effect coefficient	—	0
K3	Narrow width coefficient	—	80
K3B	Body effect coefficient of K3	$V^{-1}$	0
KETA	Body-bias coefficient of bulk charge effect	$V^{-1}$	-0.047
LINT	Length of offset fitting parameter from I-V without bias	m	0
NFACTOR	Subthreshold swing factor	—	1
NGATE	Poly gate doping concentration	$cm^{-3}$	0
NLX	Lateral non-uniform doping parameter	m	1.74e-07
PCLM	Channel length modulation parameter	—	1.3
PDIBLC1	First output resistance DIBL effect correction parameter	—	0.39
PDIBLC2	Second output resistance DIBL effect correction parameter	—	0.0086
PDIBLCB	Body effect coefficient of DIBL correction parameter	$V^{-1}$	0
PRWB	Body effect coefficient of RDSW	$V^{-1/2}$	0
PRWG	Gate-bias effect coefficient of RDSW	$V^{-1}$	0
PSCBE1	First substrate current body effect parameter	V/m	4.24e+08
PSCBE2	second substrate current body effect parameter	V/m	1e-05
PVAG	Gate dependence of early voltage	—	0
RDSW	Parasitic resistance per unit width	$\Omega \mu m$	0
UA	First-order mobility degradation coefficient	m/V	2.25e-09

Table 2.63: BSIM3 Device Model Parameters

Parameter	Description	Units	Default
UB	First-order mobility degradation coefficient	$\text{m}^2/\text{V}^2$	5.87e-19
UC	Body effect of mobility degradation coefficient	$\text{m}/\text{V}^2$	0
VBM	Maximum applied body-bias in threshold voltage calculation	V	-3
VFB	Flat-band voltage	V	0
VOFF	Offset voltage in the subthreshold region at large W and L	V	-0.08
VSAT	Saturation velocity at temp = TNOM	m/s	80000
VTH0	Threshold voltage at Vbs = 0 for large L	V	0
WO	Narrow-width parameter	m	2.5e-06
WINT	Width-offset fitting parameter from I-V without bias	m	0
WR	Width offset from Weff for Rds Calculation	—	1
<b>Dependency Parameters</b>			
LA0	Length dependence of A0	m	0
LA1	Length dependence of A1	m/V	0
LA2	Length dependence of A2	m	0
LACDE	Length dependence of ACDE	$\text{m}^2/\text{V}$	0
LAGS	Length dependence of AGS	m/V	0
LALPHA0	Length dependence of ALPHA0	$\text{m}^2/\text{V}$	0
LALPHA1	Length dependence of ALPHA1	m/V	0
LAT	Length dependence of AT	$\text{m}^2/\text{s}$	0
LB0	Length dependence of B0	$\text{m}^2$	0
LB1	Length dependence of B1	$\text{m}^2$	0
LBETA0	Length dependence of BETA0	Vm	0
LCDSC	Length dependence of CDSC	F/m	0
LCDS CB	Length dependence of CDSCB	F/(Vm)	0
LCDS CD	Length dependence of CDSCD	F/(Vm)	0
LCF	Length dependence of CF	F	0
LCGDL	Length dependence of CGDL	F	0
LCGSL	Length dependence of CGSL	F	0
LCIT	Length dependence of CIT	F/m	0
LCKAPPA	Length dependence of CKAPPA	F	0
LCLC	Length dependence of CLC	$\text{m}^2$	0
LCLE	Length dependence of CLE	m	0
LDELTA	Length dependence of DELTA	Vm	0

Table 2.63: BSIM3 Device Model Parameters

Parameter	Description	Units	Default
LDROUT	Length dependence of DROUT	m	0
LDSUB	Length dependence of DSUB	m	0
LDVT0	Length dependence of DVT0	m	0
LDVT0W	Length dependence of DVT0W	–	0
LDVT1	Length dependence of DVT1	m	0
LDVT1W	Length dependence of DVT1W	–	0
LDVT2	Length dependence of DVT2	m/V	0
LDVT2W	Length dependence of DVT2W	m/V	0
LDWB	Length dependence of DWB	$m^2/V^{1/2}$	0
LDWG	Length dependence of DWG	$m^2/V^{1/2}$	0
LELM	Length dependence of ELM	m	0
LETA0	Length dependence of ETA0	m	0
LETAB	Length dependence of ETAB	m/V	0
LGAMMA1	Length dependence of GAMMA1	$V^{1/2}m$	0
LGAMMA2	Length dependence of GAMMA2	$V^{1/2}m$	0
LK1	Length dependence of K1	$V^{1/2}m$	0
LK2	Length dependence of K2	m	0
LK3	Length dependence of K3	m	0
LK3B	Length dependence of K3B	m/V	0
LKETA	Length dependence of KETA	m/V	0
LKT1	Length dependence of KT1	Vm	0
LKT1L	Length dependence of KT1L	$Vm^2$	0
LKT2	Length dependence of KT2	m	0
LMOIN	Length dependence of MOIN	m	0
LNCH	Length dependence of NCH	$m/cm^3$	0
LNFACTOR	Length dependence of NFACTOR	m	0
LNGATE	Length dependence of NGATE	$m/cm^3$	0
LNLX	Length dependence of NLX	$m^2$	0
LNOFF	Length dependence of NOFF	m	0
LNSUB	Length dependence of NSUB	$m/cm^3$	0
LPCLM	Length dependence of PCLM	m	0
LPDIBLC1	Length dependence of PDIBLC1	m	0
LPDIBLC2	Length dependence of PDIBLC2	m	0
LPDIBLCB	Length dependence of PDIBLCB	m/V	0
LPRT	Length dependence of PRT	$\Omega \mu m m$	0

Table 2.63: BSIM3 Device Model Parameters

Parameter	Description	Units	Default
LPRWB	Length dependence of PRWB	$m/V^{1/2}$	0
LPRWG	Length dependence of PRWG	$m/V$	0
LPSCBE1	Length dependence of PSCBE1	V	0
LPSCBE2	Length dependence of PSCBE2	V	0
LPVAG	Length dependence of PVAG	m	0
LRDSW	Length dependence of RDSW	$\Omega \mu m m$	0
LU0	Length dependence of U0	$m/(Vcm^2s)$	0
LUA	Length dependence of UA	$m^2/V$	0
LUA1	Length dependence of UA1	$m^2/V$	0
LUB	Length dependence of UB	$m^3/V^2$	0
LUB1	Length dependence of UB1	$m^3/V^2$	0
LUC	Length dependence of UC	$m^2/V^2$	0
LUC1	Length dependence of UC1	$m^2/(^{\circ}CV^2)$	0
LUTE	Length dependence of UTE	m	0
LVBM	Length dependence of VBM	Vm	0
LVBX	Length dependence of VBX	Vm	0
LVFB	Length dependence of VFB	Vm	0
LVFBCV	Length dependence of VFBCV	Vm	0
LVOFF	Length dependence of VOFF	Vm	0
LVOFFCV	Length dependence of VOFFCV	Vm	0
LVSAT	Length dependence of VSAT	$m^2/s$	0
LVTH0	Length dependence of VTH0	Vm	0
LW0	Length dependence of W0	$m^2$	0
LWR	Length dependence of WR	m	0
LXJ	Length dependence of XJ	$m^2$	0
LXT	Length dependence of XT	$m^2$	0
PA0	Cross-term dependence of A0	m	0
PA1	Cross-term dependence of A1	$m/V$	0
PA2	Cross-term dependence of A2	m	0
PACDE	Cross-term dependence of ACDE	$m^2/V$	0
PAGS	Cross-term dependence of AGS	$m/V$	0
PALPHA0	Cross-term dependence of ALPHA0	$m^2/V$	0
PALPHA1	Cross-term dependence of ALPHA1	$m/V$	0
PAT	Cross-term dependence of AT	$m^2/s$	0
PB0	Cross-term dependence of B0	$m^2$	0

Table 2.63: BSIM3 Device Model Parameters

Parameter	Description	Units	Default
PB1	Cross-term dependence of B1	m <sup>2</sup>	0
PBETA0	Cross-term dependence of BETA0	Vm	0
PCDSC	Cross-term dependence of CDSC	F/m	0
PCDSCB	Cross-term dependence of CDSCB	F/(Vm)	0
PCDSCD	Cross-term dependence of CDSCD	F/(Vm)	0
PCF	Cross-term dependence of CF	F	0
PCGDL	Cross-term dependence of CGDL	F	0
PCGSL	Cross-term dependence of CGSL	F	0
PCIT	Cross-term dependence of CIT	F/m	0
PCKAPPA	Cross-term dependence of CKAPPA	F	0
PCLC	Cross-term dependence of CLC	m <sup>2</sup>	0
PCLE	Cross-term dependence of CLE	m	0
PDELTA	Cross-term dependence of DELTA	Vm	0
PDROUT	Cross-term dependence of DROUT	m	0
PDSUB	Cross-term dependence of DSUB	m	0
PDVT0	Cross-term dependence of DVT0	m	0
PDVT0W	Cross-term dependence of DVT0W	–	0
PDVT1	Cross-term dependence of DVT1	m	0
PDVT1W	Cross-term dependence of DVT1W	–	0
PDVT2	Cross-term dependence of DVT2	m/V	0
PDVT2W	Cross-term dependence of DVT2W	m/V	0
PDWB	Cross-term dependence of DWB	m <sup>2</sup> /V <sup>1/2</sup>	0
PDWG	Cross-term dependence of DWG	m <sup>2</sup> /V <sup>1/2</sup>	0
PELM	Cross-term dependence of ELM	m	0
PETA0	Cross-term dependence of ETA0	m	0
PETAB	Cross-term dependence of ETAB	m/V	0
PGAMMA1	Cross-term dependence of GAMMA1	V <sup>1/2</sup> m	0
PGAMMA2	Cross-term dependence of GAMMA2	V <sup>1/2</sup> m	0
PK1	Cross-term dependence of K1	V <sup>1/2</sup> m	0
PK2	Cross-term dependence of K2	m	0
PK3	Cross-term dependence of K3	m	0
PK3B	Cross-term dependence of K3B	m/V	0
PKETA	Cross-term dependence of KETA	m/V	0
PKT1	Cross-term dependence of KT1	Vm	0
PKT1L	Cross-term dependence of KT1L	Vm <sup>2</sup>	0

Table 2.63: BSIM3 Device Model Parameters

Parameter	Description	Units	Default
PKT2	Cross-term dependence of KT2	m	0
PMOIN	Cross-term dependence of MOIN	m	0
PNCH	Cross-term dependence of NCH	m/cm <sup>3</sup>	0
PNFACTOR	Cross-term dependence of NFACTOR	m	0
PNGATE	Cross-term dependence of NGATE	m/cm <sup>3</sup>	0
PNLX	Cross-term dependence of NLX	m <sup>2</sup>	0
PNOFF	Cross-term dependence of NOFF	m	0
PNSUB	Cross-term dependence of NSUB	m/cm <sup>3</sup>	0
PPCLM	Cross-term dependence of PCLM	m	0
PPDIBLC1	Cross-term dependence of PDIBLC1	m	0
PPDIBLC2	Cross-term dependence of PDIBLC2	m	0
PPDIBLCB	Cross-term dependence of PDIBLCB	m/V	0
PPRT	Cross-term dependence of PRT	$\Omega \mu\text{m m}$	0
PPRWB	Cross-term dependence of PRWB	m/V <sup>1/2</sup>	0
PPRWG	Cross-term dependence of PRWG	m/V	0
PPSCBE1	Cross-term dependence of PSCBE1	V	0
PPSCBE2	Cross-term dependence of PSCBE2	V	0
PPVAG	Cross-term dependence of PVAG	m	0
PRDSW	Cross-term dependence of RDSW	$\Omega \mu\text{m m}$	0
PU0	Cross-term dependence of U0	m/(Vcm <sup>2</sup> s)	0
PUA	Cross-term dependence of UA	m <sup>2</sup> /V	0
PUA1	Cross-term dependence of UA1	m <sup>2</sup> /V	0
PUB	Cross-term dependence of UB	m <sup>3</sup> /V <sup>2</sup>	0
PUB1	Cross-term dependence of UB1	m <sup>3</sup> /V <sup>2</sup>	0
PUC	Cross-term dependence of UC	m <sup>2</sup> /V <sup>2</sup>	0
PUC1	Cross-term dependence of UC1	m <sup>2</sup> /(°CV <sup>2</sup> )	0
PUTE	Cross-term dependence of UTE	m	0
PVBM	Cross-term dependence of VBM	Vm	0
PVBX	Cross-term dependence of VBX	Vm	0
PVFB	Cross-term dependence of VFB	Vm	0
PVFBCV	Cross-term dependence of VFBCV	Vm	0
PVOFF	Cross-term dependence of VOFF	Vm	0
PVOFFCV	Cross-term dependence of VOFFCV	Vm	0
PVSAT	Cross-term dependence of VSAT	m <sup>2</sup> /s	0
PVTH0	Cross-term dependence of VTH0	Vm	0

Table 2.63: BSIM3 Device Model Parameters

Parameter	Description	Units	Default
PW0	Cross-term dependence of W0	m <sup>2</sup>	0
PWR	Cross-term dependence of WR	m	0
PXJ	Cross-term dependence of XJ	m <sup>2</sup>	0
PXT	Cross-term dependence of XT	m <sup>2</sup>	0
WA0	Width dependence of A0	m	0
WA1	Width dependence of A1	m/V	0
WA2	Width dependence of A2	m	0
WACDE	Width dependence of ACDE	m <sup>2</sup> /V	0
WAGS	Width dependence of AGS	m/V	0
WALPHA0	Width dependence of ALPHA0	m <sup>2</sup> /V	0
WALPHA1	Width dependence of ALPHA1	m/V	0
WAT	Width dependence of AT	m <sup>2</sup> /s	0
WB0	Width dependence of B0	m <sup>2</sup>	0
WB1	Width dependence of B1	m <sup>2</sup>	0
WBETA0	Width dependence of BETA0	Vm	0
WCDSC	Width dependence of CDSC	F/m	0
WCDSCB	Width dependence of CDSCB	F/(Vm)	0
WCDSCD	Width dependence of CDSCD	F/(Vm)	0
WCF	Width dependence of CF	F	0
WCGDL	Width dependence of CGDL	F	0
WCGSL	Width dependence of CGSL	F	0
WCIT	Width dependence of CIT	F/m	0
WCKAPPA	Width dependence of CKAPPA	F	0
WCLC	Width dependence of CLC	m <sup>2</sup>	0
WCLE	Width dependence of CLE	m	0
WDELTA	Width dependence of DELTA	Vm	0
WDROUT	Width dependence of DROUT	m	0
WDSUB	Width dependence of DSUB	m	0
WDVT0	Width dependence of DVT0	m	0
WDVT0W	Width dependence of DVT0W	–	0
WDVT1	Width dependence of DVT1	m	0
WDVT1W	Width dependence of DVT1W	–	0
WDVT2	Width dependence of DVT2	m/V	0
WDVT2W	Width dependence of DVT2W	m/V	0
WDWB	Width dependence of DWB	m <sup>2</sup> /V <sup>1/2</sup>	0

Table 2.63: BSIM3 Device Model Parameters

Parameter	Description	Units	Default
WDWG	Width dependence of DWG	$m^2/V^{1/2}$	0
WELM	Width dependence of ELM	m	0
WETA0	Width dependence of ETA0	m	0
WETAB	Width dependence of ETAB	m/V	0
WGAMMA1	Width dependence of GAMMA1	$V^{1/2}m$	0
WGAMMA2	Width dependence of GAMMA2	$V^{1/2}m$	0
WK1	Width dependence of K1	$V^{1/2}m$	0
WK2	Width dependence of K2	m	0
WK3	Width dependence of K3	m	0
WK3B	Width dependence of K3B	m/V	0
WKETA	Width dependence of KETA	m/V	0
WKT1	Width dependence of KT1	Vm	0
WKT1L	Width dependence of KT1L	$Vm^2$	0
WKT2	Width dependence of KT2	m	0
WMOIN	Width dependence of MOIN	m	0
WNCH	Width dependence of NCH	$m/cm^3$	0
WNFACTOR	Width dependence of NFACTOR	m	0
WNGATE	Width dependence of NGATE	$m/cm^3$	0
WNLX	Width dependence of NLX	$m^2$	0
WNOFF	Width dependence of NOFF	m	0
WNSUB	Width dependence of NSUB	$m/cm^3$	0
WPCLM	Width dependence of PCLM	m	0
WPDIBLC1	Width dependence of PDIBLC1	m	0
WPDIBLC2	Width dependence of PDIBLC2	m	0
WPDIBLCB	Width dependence of PDIBLCB	m/V	0
WPRT	Width dependence of PRT	$\Omega \mu m m$	0
WPRWB	Width dependence of PRWB	$m/V^{1/2}$	0
WPRWG	Width dependence of PRWG	m/V	0
WPSCBE1	Width dependence of PSCBE1	V	0
WPSCBE2	Width dependence of PSCBE2	V	0
WPVAG	Width dependence of PVAG	m	0
WRDSW	Width dependence of RDSW	$\Omega \mu m m$	0
WU0	Width dependence of U0	$m/(Vcm^2s)$	0
WUA	Width dependence of UA	$m^2/V$	0
WUA1	Width dependence of UA1	$m^2/V$	0

Table 2.63: BSIM3 Device Model Parameters

Parameter	Description	Units	Default
WUB	Width dependence of UB	$\text{m}^3/\text{V}^2$	0
WUB1	Width dependence of UB1	$\text{m}^3/\text{V}^2$	0
WUC	Width dependence of UC	$\text{m}^2/\text{V}^2$	0
WUC1	Width dependence of UC1	$\text{m}^2/(\text{°CV}^2)$	0
WUTE	Width dependence of UTE	m	0
WVBM	Width dependence of VBM	Vm	0
WVBX	Width dependence of VBX	Vm	0
WVFB	Width dependence of VFB	Vm	0
WVBCV	Width dependence of VFBCV	Vm	0
WVOFF	Width dependence of VOFF	Vm	0
WVOFFCV	Width dependence of VOFFCV	Vm	0
WVSAT	Width dependence of VSAT	$\text{m}^2/\text{s}$	0
WVTHO	Width dependence of VTH0	Vm	0
WVO	Width dependence of W0	$\text{m}^2$	0
WWR	Width dependence of WR	m	0
WXJ	Width dependence of XJ	$\text{m}^2$	0
WXT	Width dependence of XT	$\text{m}^2$	0
<b><i>Doping Parameters</i></b>			
MJ	Bulk p-n bottom grading coefficient	–	0.5
MJSW	Bulk p-n sidewall grading coefficient	–	0.33
NSUB	Substrate doping density	$\text{cm}^{-3}$	$6\text{e}+16$
<b><i>Flicker Parameters</i></b>			
AF	Flicker noise exponent	–	1
EF	Flicker exponent	–	1
EM	Saturation field	V/m	$4.1\text{e}+07$
KF	Flicker noise coefficient	–	0
NOIA	Noise parameter a	–	0
NOIB	Noise parameter b	–	0
NOIC	Noise parameter c	–	0
<b><i>Geometry Parameters</i></b>			
L	Channel length	m	$5\text{e}-06$
LL	Coefficient of length dependence for length offset	$\text{m}^{LLN}$	0
LLC	Coefficient of length dependence for CV channel length offset	$\text{m}^{LLN}$	0
LLN	Power of length dependence for length offset	–	0

Table 2.63: BSIM3 Device Model Parameters

Parameter	Description	Units	Default
LW	Coefficient of width dependence for length offset	$m^{LWN}$	0
LWC	Coefficient of width dependence for channel length offset	$m^{LWN}$	0
LWL	Coefficient of length and width cross term for length offset	$m^{LLN+LWN}$	0
LWLC	Coefficient of length and width dependence for CV channel length offset	$m^{LLN+LWN}$	0
LWN	Power of width dependence for length offset	–	0
TOX	Gate oxide thickness	m	1.5e-08
W	Channel width	m	5e-06
WL	Coefficient of length dependence for width offset	$m^{WLN}$	0
WLC	Coefficient of length dependence for CV channel width offset	$m^{WLN}$	0
WLN	Power of length dependence of width offset	–	0
WW	Coefficient of width dependence for width offset	$m^{WWN}$	0
WWC	Coefficient of width dependence for CV channel width offset	$m^{WWN}$	0
WWL	Coefficient of length and width cross term for width offset	$m^{WLN+WWN}$	0
WWLC	Coefficient of length and width dependence for CV channel width offset	$m^{WLN+WWN}$	0
WWN	Power of width dependence of width offset	–	0
XJ	Junction depth	m	1.5e-07
<b><i>NQS Parameters</i></b>			
ELM	Elmore constant of the channel	–	5
<b><i>Resistance Parameters</i></b>			
RSH	Drain, source diffusion sheet resistance	$\Omega$	0
<b><i>Process Parameters</i></b>			
GAMMA1	Body effect coefficient near the surface	$V^{1/2}$	0
GAMMA2	Body effect coefficient in the bulk	$V^{1/2}$	0
JS	Bulk p-n saturation current density	$A/m^2$	0.0001
NCH	Channel doping concentration	$cm^{-3}$	1.7e+17
TOXM	Gate oxide thickness used in extraction	m	0
U0	Surface mobility	$1/(Vcm^2s)$	0
VBX	Vbs at which the depletion region = XT	V	0
XT	Doping depth	m	1.55e-07
<b><i>Temperature Parameters</i></b>			

Table 2.63: BSIM3 Device Model Parameters

Parameter	Description	Units	Default
AT	Temperature coefficient for saturation velocity	m/s	33000
KT1	Temperature coefficient for threshold voltage	V	-0.11
KT1L	Channel length dependence of the temperature coefficient for the threshold voltage	Vm	0
KT2	Body-bias coefficient for the threshold voltage temperature effect	—	0.022
NJ	Emission coefficient of junction	—	1
PRT	Temperature coefficient for RDSW	$\Omega \mu\text{m}$	0
TCJ	Temperature coefficient of $C_j$	$\text{K}^{-1}$	0
TCJSW	Temperature coefficient of $C_{swj}$	$\text{K}^{-1}$	0
TCJSWG	Temperature coefficient of $C_{jswg}$	$\text{K}^{-1}$	0
TNOM	Nominal device temperature	$^{\circ}\text{C}$	Ambient Temperature
TPB	Temperature coefficient of $P_b$	V/K	0
TPBSW	Temperature coefficient of $P_{bsw}$	V/K	0
TPBSWG	Temperature coefficient of $P_{bswg}$	V/K	0
UA1	Temperature coefficient for UA	m/V	4.31e-09
UB1	Temperature coefficient for UB	$\text{m}^2/\text{V}^2$	-7.61e-18
UC1	Temperature coefficient for UC	$\text{m}/(^{\circ}\text{C V}^2)$	0
UTE	Mobility temperature exponent	—	-1.5
XTI	Junction current temperature exponent coefficient	—	3
<b><i>Voltage Parameters</i></b>			
PB	Bulk p-n bottom potential	V	1

## Level 10 MOSFET Tables (BSIM SOI)

For complete documentation of the BSIMSOI model, see the users' manual for the BSIMSOI, available for download at <http://www-device.eecs.berkeley.edu/bsim/?page=BSIMSOI>. **Xyce** implements Version 3.2 of the BSIMSOI, you will have to get the documentation from the FTP archive on the Berkeley site.

In addition to the parameters shown in table 2.64, the BSIM3SOI supports a vector parameter for the initial conditions. IC1 through IC5 may therefore be specified compactly as IC=<ic1>,<ic2>,<ic3>,<ic4>,<ic5>.

**NOTE: Many BSIM SOI parameters listed in tables 2.64 and 2.65 as having default values of zero are actually replaced with internally computed defaults if not given. Specifying zero in your model card will override this internal computation. It is recommended that you only set model parameters that you are actually changing from defaults and that you not generate model cards containing default values from the tables.**

Table 2.64: BSIM3 SOI Device Instance Parameters

Parameter	Description	Units	Default
BJTOFF	BJT on/off flag	logical (T/F)	0
DEBUG	BJT on/off flag	logical (T/F)	0
TNODEOUT	Flag indicating external temp node	logical (T/F)	0
VLDEBUG		logical (T/F)	false
<b>Control Parameters</b>			
M	Multiplier for M devices connected in parallel	—	1
SOIMOD	SIO model selector, SOIMOD=0: BSIMPD, SOIMOD=1: undefined model for PD and FE, SOIMOD=2: ideal FD	—	0
<b>DC Parameters</b>			
VBSUSR	Vbs specified by user	V	0
<b>Geometry Parameters</b>			
AD	Drain diffusion area	m <sup>2</sup>	0
AEBCP	Substrate to body overlap area for bc parasitics	m <sup>2</sup>	0
AGBCP	Gate to body overlap area for bc parasitics	m <sup>2</sup>	0
AS	Source diffusion area	m <sup>2</sup>	0
FRBODY	Layout dependent body-resistance coefficient	—	1
L	Channel length	m	5e-06
NBC	Number of body contact isolation edge	—	0
NRB	Number of squares in body	—	1

Table 2.64: BSIM3 SOI Device Instance Parameters

Parameter	Description	Units	Default
NRD	Multiplier for RSH to yield parasitic resistance of drain	$\square$	1
NRS	Multiplier for RSH to yield parasitic resistance of source	$\square$	1
NSEG	Number segments for width partitioning	–	1
PD	Drain diffusion perimeter	m	0
PDBCP	Perimeter length for bc parasitics at drain side	m	0
PS	Source diffusion perimeter	m	0
PSBCP	Perimeter length for bc parasitics at source side	m	0
W	Channel width	m	5e-06
<b><i>RF Parameters</i></b>			
RGATEMOD	Gate resistance model selector	–	0
<b><i>Temperature Parameters</i></b>			
CTHO	Thermal capacitance	F	0
RTHO	normalized thermal resistance	$\Omega$	0
TEMP	Device temperature	$^{\circ}\text{C}$	27
<b><i>Voltage Parameters</i></b>			
IC1	Initial condition on Vds	V	0
IC2	Initial condition on Vgs	V	0
IC3	Initial condition on Vbs	V	0
IC4	Initial condition on Ves	V	0
IC5	Initial condition on Vps	V	0
OFF	Initial condition of no voltage drops accross device	logical (T/F)	false

Table 2.65: BSIM3 SOI Device Model Parameters

Parameter	Description	Units	Default
DELTAVOX	The smoothing parameter in the Vox smoothing function	–	0
DTOXCV	Delta oxide thickness in meters in CapMod3	m	0
FNOIMOD	Flicker noise model selector	–	1
IGBMOD	Flicker noise model selector	–	0
IGCMOD	Gate-channel tunneling current model selector	–	0
KB1	Scaling factor for backgate charge	–	1
NOIF	Floating body excess noise ideality factor	–	1

Table 2.65: BSIM3 SOI Device Model Parameters

Parameter	Description	Units	Default
NTNOI	Thermal noise parameter	–	1
POXEDGE	Factor for the gate edge $T_{ox}$	–	1
RNOIA	Thermal noise coefficient	–	0.577
RNOIB	Thermal noise coefficient	–	0.37
RSHG	Gate sheet resistance	–	0.1
TNOIA	Thermal noise parameter	–	1.5
TNOIB	Thermal noise parameter	–	3.5
TNOIMOD	Thermal noise model selector	–	0
VBSOFD	Lower bound of built-in potential lowering for FD operation	V	0.5
VBSOPD	Upper bound of built-in potential lowering for FD operation	–	0
VOXH	The limit of $V_{ox}$ in gate current calculation	–	0
VTHO	Threshold voltage	–	0
<b>Bin Parameters</b>			
LMAX	Maximum channel length	m	1
LMIN	Minimum channel length	m	0
WMAX	Maximum channel width	m	1
WMIN	Minimum channel width	m	0
<b>Capacitance Parameters</b>			
ACDE	Exponential coefficient for charge thickness in $capmod = 3$ for accumulation and depletion regions	m/V	1
ASD	Source/Drain bottom diffusion smoothing parameter	–	0.3
CF	Firing field capacitance	F/m	0
CGDL	Light-doped drain-gate region overlap capacitance	F/m	0
CGDO	Non-LLD region drain-gate overlap capacitance per unit channel length	F/m	0
CGEO	Gate substrate overlap capacitance per unit channel length	F/m	0
CGSL	Light-doped source-gate region overlap capacitance	F/m	0
CGSO	Non-LLD region source-gate overlap capacitance per unit channel length	F/m	0
CJSWG	Source/grain gate sidewall junction capacitance per unit width	F/m	1e-10

Table 2.65: BSIM3 SOI Device Model Parameters

Parameter	Description	Units	Default
CKAPPA	Coefficient for lightly doped region overlap capacitance firing field capacitance	F/m	0.6
CLC	Constant term for short-channel model	m	1e-08
CLE	Exponential term for the short-channel model	–	0
CSDSW	Source/Drain sidewall fringing capacitance per unit length	F/m	0
CSDMIN	Source/Drain bottom diffusion minimum capacitance	V	0
DELVT	Threshold voltage adjust for C-V	V	0
DLBG	Length offset fitting parameter for backgate charge	m	0
DLC	Length offset fitting parameter from C-V	m	0
DLCB	Length offset fitting parameter for body charge	m	0
DWC	Width offset fitting parameter from C-V	m	0
FBODY	Scaling factor for body charge	–	1
LDIFO	Channel length dependency coefficient of diffusion capacitance	–	1
MJSWG	Source/grain gate sidewall junction capacitance grading coefficient	–	0.5
MOIN	Coefficient for the gate-bias dependent surface potential	–	15
NDIF	Power coefficient of channel length dependency for diffusion capacitance	–	-1
NOFF	CV parameter in $V_{gsteff}$ , CV for weak to strong inversion	–	1
PBSWG	Source/drain gate sidewall junction built-in potential	V	0.7
TT	Diffusion capacitance transit time coefficient	s	1e-12
VSDFB	Source/Drain bottom diffusion capacitance flatband voltage	V	0
VSDTH	Source/Drain bottom diffusion capacitance threshold voltage	V	0
XPART	Charge partitioning rate flag	–	0
<b>Control Parameters</b>			
BINUNIT	Binning unit selector	–	1
CAPMOD	Flag for capacitance models	–	2
MOBMOD	Mobility model selector	–	1
PARAMCHK	Parameter value check	–	0

Table 2.65: BSIM3 SOI Device Model Parameters

Parameter	Description	Units	Default
SHMOD	Flag for self-heating, 0-no self-heating, 1-self-heating	–	0
TEMPMODEL	Specifies the type of parameter interpolation over temperature	–	'NONE'
VERSION	Version number	–	'3.2'
<b>Current Parameters</b>			
AIGC	Parameter for $I_{gc}$	$(F/g)^{1/2}s/mV$	0
AIGSD	Parameter for $I_{gs,d}$	$(F/g)^{1/2}s/mV$	0
BIGC	Parameter for $I_{gc}$	$(F/g)^{1/2}s/mV$	0
BIGSD	Parameter for $I_{gs,d}$	$(F/g)^{1/2}s/mV$	0
CIGC	Parameter for $I_{gc}$	$V^{-1}$	0
CIGSD	Parameter for $I_{gs,d}$	$V^{-1}$	0
DLICG	Delta L for $I_g$ model	$V^{-1}$	0
NIGC	Parameter for $I_{gc}$ slope	–	1
PIGCD	Parameter for $I_{gc}$ partition	–	1
<b>DC Parameters</b>			
A0	Bulk charge effect coefficient for channel length	–	1
A1	First non-saturation effect parameter	$V^{-1}$	0
A2	Second non-saturation factor	–	1
AELY	Channel length dependency of early voltage for bipolar current	V/m	0
AGIDL	GIDL constant	$\Omega^{-1}$	0
AGS	Gate-bias coefficient of $a_{bulk}$	$V^{-1}$	0
AHLI	High level injection parameter for bipolar current	–	0
ALPHA0	First parameter of impact-ionization current	m/V	0
B0	Bulk charge effect coefficient for channel width	m	0
B1	Bulk charge effect offset	m	0
BETA0	Second parameter of impact-ionization current	V	0
BETA1	Second $V_{ds}$ dependent parameter of impact ionization current	–	0
BETA2	Third $V_{ds}$ dependent parameter of impact ionization current	V	0.1
BGIDL	GIDL exponential coefficient	V/m	0
CDSC	Drain/source to channel coupling capacitance	F/m <sup>2</sup>	0.00024
CDSCB	Body-bias sensitivity of CDSC	F/(Vm <sup>2</sup> )	0
CDSCD	Drain-bias sensitivity of CDSC	F/(Vm <sup>2</sup> )	0

Table 2.65: BSIM3 SOI Device Model Parameters

Parameter	Description	Units	Default
CIT	Interface trap capacitance	F/m <sup>2</sup>	0
DELTA	Effective Vds parameter	V	0.01
DROUT	L-depedance Coefficient of the DIBL correction parameter in Rout	–	0.56
DSUB	DIBL coefficient exponent in subthreshold region	–	0
DVT0	First coefficient of short-channel effect effect on threshold voltage	–	2.2
DVT0W	First coefficient of narrow-width effect effect on threshold voltage for small channel length	m <sup>-1</sup>	0
DVT1	Second coefficient of short-channel effect effect on threshold voltage	–	0.53
DVT1W	Second coefficient of narrow-width effect effect on threshold voltage for small channel length	m <sup>-1</sup>	5.3e+06
DVT2	Body-bias coefficient of short-channel effect effect on threshold voltage	V <sup>-1</sup>	-0.032
DVT2W	Body-bias coefficient of narrow-width effect effect on threshold voltage for small channel length	V <sup>-1</sup>	-0.032
DWB	Coefficient of substrate body bias dependence of Weff	m/V <sup>1/2</sup>	0
DWBC	Width offset for body contact isolation edge	m	0
DWG	Coefficient of gate depedence of Weff	m/V <sup>1/2</sup>	0
ESATII	Saturation channel electric field for impact ionization current	V/m	1e+07
ETA0	DIBL coefficient in subthreshold region	–	0.08
ETAB	Body-bias coefficient for the subthreshold DIBL effect	V <sup>-1</sup>	-0.07
FBJTII	Fraction of bipolar current affecting the impact ionization	–	0
ISBJT	BJT injection saturation current	A/m <sup>2</sup>	1e-06
ISDIF	BOdy to source/drain injection saturation current	A/m <sup>2</sup>	0
ISREC	Recombinatin in depletion saturation current	A/m <sup>2</sup>	1e-05
ISTUN	Reverse tunneling saturation current	A/m <sup>2</sup>	0
K1	First-order body effect coefficient	V <sup>1/2</sup>	0.53
K1W1	First body effect width depenent parameter	m	0
K1W2	Second body effect width depenent parameter	m	0
K2	second-order body effect coefficient	–	-0.0186
K3	Narrow width coefficient	–	0
K3B	Body effect coefficient of K3	V <sup>-1</sup>	0

Table 2.65: BSIM3 SOI Device Model Parameters

Parameter	Description	Units	Default
KETA	Body-bias coefficient of bulk charge effect	$V^{-1}$	-0.6
KETAS	Surface potential adjustment for bulk charge effect	V	0
LBJTO	Reference channel length for bipolar current	m	2e-07
LII	Channel length dependent parameter at threshold for impact ionization current	–	0
LINT	Length of offset fitting parameter from I-V without bias	m	0
LN	Electron/hole diffusion length	m	2e-06
NBJT	Power coefficient of channel length	–	1
NDIODE	Diode non-ideality factor	–	1
NFACTOR	Subthreshold swing factor	–	1
NGATE	Poly gate doping concentration	$cm^{-3}$	0
NGIDL	GIDL Vds enhancement coefficient	V	1.2
NLX	Lateral non-uniform doping parameter	m	1.74e-07
NRECF0	Recombination non-ideality factor at forward bias	–	2
NRECR0	Recombination non-ideality factor at reverse bias	–	10
NTUN	Reverse tunneling non-ideality factor	–	10
PCLM	Channel length modulation parameter	–	1.3
PDIBLC1	First output resistance DIBL effect correction parameter	–	0.39
PDIBLC2	Second output resistance DIBL effect correction parameter	–	0.0086
PDIBLCB	Body effect coefficient of DIBL correction parameter	$V^{-1}$	0
PRWB	Body effect coefficient of RDSW	$V^{-1/2}$	0
PRWG	Gate-bias effect coefficient of RDSW	$V^{-1}$	0
PVAG	Gate dependence of early voltage	–	0
RBODY	Intrinsic body contact sheet resistance	$\Omega/\square$	0
RBSH	Intrinsic body contact sheet resistance	$\Omega/\square$	0
RDSW	Parasitic resistance per unit width	$\Omega \mu m$	100
RHALO	Body halo sheet resistance	$\Omega/m$	1e+15
SII0	First Vgs dependent parameter of impact ionization current	$V^{-1}$	0.5
SII1	Second Vgs dependent parameter of impact ionization current	$V^{-1}$	0.1
SII2	Third Vgs dependent parameter of impact ionization current	–	0

Table 2.65: BSIM3 SOI Device Model Parameters

Parameter	Description	Units	Default
SIID	Vds dependent parameter of drain saturation voltage for impact ionization current	$V^{-1}$	0
TII	Temperature dependent parameter for impact ionization current	–	0
UA	First-order mobility degradation coefficient	m/V	2.25e-09
UB	First-order mobility degradation coefficient	$m^2/V^2$	5.87e-19
UC	Body effect of mobility degradation coefficient	$m/V^2$	0
VABJT	Early voltage for bipolar current	V	10
VBM	Maximum applied body-bias in threshold voltage calculation	V	-3
VDSATII0	Normal drain saturation voltage at threshold for impact ionization current	V	0.9
VOFF	Offset voltage in the subthreshold region at large W and L	V	-0.08
VRECO	Voltage dependent parameter for recombination current	V	0
VSAT	Saturation velocity at temp = TNOM	m/s	80000
VTH0	Threshold voltage at Vbs = 0 for large L	V	0
VTUN0	Voltage dependent parameter for tunneling current	V	0
W0	Narrow-width parameter	m	2.5e-06
WINT	Width-offset fitting parameter from I-V without bias	m	0
WR	Width offset from Weff for Rds Calculation	–	1
<b>Dependency Parameters</b>			
LA0	Length dependence of A0	m	0
LA1	Length dependence of A1	m/V	0
LA2	Length dependence of A2	m	0
LACDE	Length dependence of ACDE	$m^2/V$	0
LAELY	Length dependence of AELY	V	0
LAGIDL	Length dependence of AGIDL	$m/\Omega$	0
LAGS	Length dependence of AGS	m/V	0
LAHLI	Length dependence of AHLI	m	0
LAIGC	Length dependence of AIGC	$(F/g)^{1/2} \text{smOnV}$	0
LAIGSD	Length dependence of AIGSD	$(F/g)^{1/2} \text{smOnV}$	0
LALPHA0	Length dependence of ALPHA0	$m^2/V$	0
LALPHAGB1	Length dependence of ALPHAGB1	m/V	0

Table 2.65: BSIM3 SOI Device Model Parameters

Parameter	Description	Units	Default
LALPHAGB2	Length dependence of ALPHAGB2	m/V	0
LAT	Length dependence of AT	m <sup>2</sup> /s	0
LB0	Length dependence of B0	m <sup>2</sup>	0
LB1	Length dependence of B1	m <sup>2</sup>	0
LBETA0	Length dependence of BETA0	Vm	0
LBETA1	Length dependence of BETA1	m	0
LBETA2	Length dependence of BETA2	Vm	0
LBETAGB1	Length dependence of BETAGB1	m/V <sup>2</sup>	0
LBETAGB2	Length dependence of BETAGB2	m/V <sup>2</sup>	0
LBGIDL	Length dependence of BGIDL	V	0
LBIGC	Length dependence of BIGC	(F/g) <sup>1/2</sup> sm0nV	0
LBIGSD	Length dependence of BIGSD	(F/g) <sup>1/2</sup> sm0nV	0
LCDSC	Length dependence of CDSC	F/m	0
LCDSGB	Length dependence of CDSCB	F/(Vm)	0
LCDSGD	Length dependence of CDSCD	F/(Vm)	0
LCGDL	Length dependence of CGDL	F	0
LCGSL	Length dependence of CGSL	F	0
LCIGC	Length dependence of CIGC	m/V	0
LCIGSD	Length dependence of CIGSD	m/V	0
LCIT	Length dependence of CIT	F/m	0
LCKAPPA	Length dependence of CKAPPA	F	0
LDELTA	Length dependence of DELTA	Vm	0
LDELVT	Length dependence of DELVT	Vm	0
LDROUT	Length dependence of DROUT	m	0
LDSUB	Length dependence of DSUB	m	0
LDVT0	Length dependence of DVT0	m	0
LDVT0W	Length dependence of DVT0W	–	0
LDVT1	Length dependence of DVT1	m	0
LDVT1W	Length dependence of DVT1W	–	0
LDVT2	Length dependence of DVT2	m/V	0
LDVT2W	Length dependence of DVT2W	m/V	0
LDWB	Length dependence of DWB	m <sup>2</sup> /V <sup>1/2</sup>	0
LDWG	Length dependence of DWG	m <sup>2</sup> /V <sup>1/2</sup>	0
LESATII	Length dependence of ESATII	V	0
LETA0	Length dependence of ETA0	m	0

Table 2.65: BSIM3 SOI Device Model Parameters

Parameter	Description	Units	Default
LETAB	Length dependence of ETAB	m/V	0
LFBJTII	Length dependence of FBJTII	m	0
LISBJT	Length dependence of ISBJT	A/m	0
LISDIF	Length dependence of ISDIF	A/m	0
LISREC	Length dependence of ISREC	A/m	0
LISTUN	Length dependence of ISTUN	A/m	0
LK1	Length dependence of K1	$V^{1/2}m$	0
LK1W1	Length dependence of K1W1	$m^2$	0
LK1W2	Length dependence of K1W2	$m^2$	0
LK2	Length dependence of K2	m	0
LK3	Length dependence of K3	m	0
LK3B	Length dependence of K3B	m/V	0
LKB1	Length dependence of KB1	m	0
LKETA	Length dependence of KETA	m/V	0
LKETAS	Length dependence of KETAS	Vm	0
LKT1	Length dependence of KT1	Vm	0
LKT1L	Length dependence of KT1L	$Vm^2$	0
LKT2	Length dependence of KT2	m	0
LLBJT0	Length dependence of LBJT0	$m^2$	0
LLII	Length dependence of LII	m	0
LMOIN	Length dependence of MOIN	m	0
LNBJT	Length dependence of NBJT	m	0
LNCH	Length dependence of NCH	$m/cm^3$	0
LNDIF	Length dependence of NDIF	m	0
LNDIODE	Length dependence of NDIODE	m	0
LNFACTOR	Length dependence of NFACTOR	m	0
LNGATE	Length dependence of NGATE	$m/cm^3$	0
LNGIDL	Length dependence of NGIDL	Vm	0
LNIGC	Length dependence of NIGC	m	0
LNLX	Length dependence of NLX	$m^2$	0
LNOFF	Length dependence of NOFF	m	0
LNRECF0	Length dependence of NRECF0	m	0
LNRECR0	Length dependence of NRECR0	m	0
LNSUB	Length dependence of NSUB	$m/cm^3$	0
LNTRECF	Length dependence of NTRECF	m	0

Table 2.65: BSIM3 SOI Device Model Parameters

Parameter	Description	Units	Default
LNTRECR	Length dependence of NTRECR	m	0
LNTUN	Length dependence of NTUN	m	0
LPCLM	Length dependence of PCLM	m	0
LPDIBLC1	Length dependence of PDIBLC1	m	0
LPDIBLC2	Length dependence of PDIBLC2	m	0
LPDIBLCB	Length dependence of PDIBLCB	m/V	0
LPIGCD	Length dependence of PIGCD	m	0
LPOXEDGE	Length dependence of POXEDGE	m	0
LPRT	Length dependence of PRT	$\Omega \mu\text{m m}$	0
LPRWB	Length dependence of PRWB	$\text{m}/\text{V}^{1/2}$	0
LPRWG	Length dependence of PRWG	m/V	0
LPVAG	Length dependence of PVAG	m	0
LRDSW	Length dependence of RDSW	$\Omega \mu\text{m m}$	0
LSII0	Length dependence of SII0	m/V	0
LSII1	Length dependence of SII1	m/V	0
LSII2	Length dependence of SII2	m	0
LSIID	Length dependence of SIID	m/V	0
LU0	Length dependence of U0	$\text{m}/(\text{Vcm}^2\text{s})$	0
LUA	Length dependence of UA	$\text{m}^2/\text{V}$	0
LUA1	Length dependence of UA1	$\text{m}^2/\text{V}$	0
LUB	Length dependence of UB	$\text{m}^3/\text{V}^2$	0
LUB1	Length dependence of UB1	$\text{m}^3/\text{V}^2$	0
LUC	Length dependence of UC	$\text{m}^2/\text{V}^2$	0
LUC1	Length dependence of UC1	$\text{m}^2/(\text{V}^2\text{C})$	0
LUTE	Length dependence of UTE	m	0
LVABJT	Length dependence of VABJT	Vm	0
LVDSATII0	Length dependence of VDSATII0	Vm	0
LVOFF	Length dependence of VOFF	Vm	0
LVREC0	Length dependence of VREC0	Vm	0
LVSAT	Length dependence of VSAT	$\text{m}^2/\text{s}$	0
LVSDFB	Length dependence of VSDFB	Vm	0
LVSDTH	Length dependence of VSDTH	Vm	0
LVTH0	Length dependence of VTH0	Vm	0
LVTUN0	Length dependence of VTUN0	Vm	0
LW0	Length dependence of W0	$\text{m}^2$	0

Table 2.65: BSIM3 SOI Device Model Parameters

Parameter	Description	Units	Default
LWR	Length dependence of WR	m	0
LXBJT	Length dependence of XBJT	m	0
LXDIF	Length dependence of XDIF	m	0
LXJ	Length dependence of XJ	m <sup>2</sup>	0
LXRCRG1	Length dependence of XRCRG1	m	0
LXRCRG2	Length dependence of XRCRG2	m	0
LXREC	Length dependence of XREC	m	0
LXTUN	Length dependence of XTUN	m	0
PA0	Cross-term dependence of A0	m	0
PA1	Cross-term dependence of A1	m/V	0
PA2	Cross-term dependence of A2	m	0
PACDE	Cross-term dependence of ACDE	m <sup>2</sup> /V	0
PAELY	Cross-term dependence of AELY	V	0
PAGIDL	Cross-term dependence of AGIDL	m/ $\Omega$	0
PAGS	Cross-term dependence of AGS	m/V	0
PAHLI	Cross-term dependence of AHLI	m	0
PAIGC	Cross-term dependence of AIGC	(F/g) <sup>1/2</sup> sm <sup>0</sup> nV	0
PAIGSD	Cross-term dependence of AIGSD	(F/g) <sup>1/2</sup> sm <sup>0</sup> nV	0
PALPHA0	Cross-term dependence of ALPHA0	m <sup>2</sup> /V	0
PALPHAGB1	Cross-term dependence of ALPHAGB1	m/V	0
PALPHAGB2	Cross-term dependence of ALPHAGB2	m/V	0
PAT	Cross-term dependence of AT	m <sup>2</sup> /s	0
PB0	Cross-term dependence of B0	m <sup>2</sup>	0
PB1	Cross-term dependence of B1	m <sup>2</sup>	0
PBETA0	Cross-term dependence of BETA0	Vm	0
PBETA1	Cross-term dependence of BETA1	m	0
PBETA2	Cross-term dependence of BETA2	Vm	0
PBETAGB1	Cross-term dependence of BETAGB1	m/V <sup>2</sup>	0
PBETAGB2	Cross-term dependence of BETAGB2	m/V <sup>2</sup>	0
PBGIDL	Cross-term dependence of BGIDL	V	0
PBIGC	Cross-term dependence of BIGC	(F/g) <sup>1/2</sup> sm <sup>0</sup> nV	0
PBIGSD	Cross-term dependence of BIGSD	(F/g) <sup>1/2</sup> sm <sup>0</sup> nV	0
PCDSC	Cross-term dependence of CDSC	F/m	0
PCDSCB	Cross-term dependence of CDSCB	F/(Vm)	0
PCDSCD	Cross-term dependence of CDSCD	F/(Vm)	0

Table 2.65: BSIM3 SOI Device Model Parameters

Parameter	Description	Units	Default
PCGDL	Cross-term dependence of CGDL	F	0
PCGSL	Cross-term dependence of CGSL	F	0
PCIGC	Cross-term dependence of CIGC	m/V	0
PCIGSD	Cross-term dependence of CIGSD	m/V	0
PCIT	Cross-term dependence of CIT	F/m	0
PCKAPPA	Cross-term dependence of CKAPPA	F	0
PDELTA	Cross-term dependence of DELTA	Vm	0
PDELVT	Cross-term dependence of DELVT	Vm	0
PDROUT	Cross-term dependence of DROUT	m	0
PDSUB	Cross-term dependence of DSUB	m	0
PDVT0	Cross-term dependence of DVT0	m	0
PDVT0W	Cross-term dependence of DVT0W	–	0
PDVT1	Cross-term dependence of DVT1	m	0
PDVT1W	Cross-term dependence of DVT1W	–	0
PDVT2	Cross-term dependence of DVT2	m/V	0
PDVT2W	Cross-term dependence of DVT2W	m/V	0
PDWB	Cross-term dependence of DWB	$m^2/V^{1/2}$	0
PDWG	Cross-term dependence of DWG	$m^2/V^{1/2}$	0
PESATII	Cross-term dependence of ESATII	V	0
PETA0	Cross-term dependence of ETA0	m	0
PETAB	Cross-term dependence of ETAB	m/V	0
PFBJTII	Cross-term dependence of FBJTII	m	0
PISBJT	Cross-term dependence of ISBJT	A/m	0
PISDIF	Cross-term dependence of ISDIF	A/m	0
PISREC	Cross-term dependence of ISREC	A/m	0
PISTUN	Cross-term dependence of ISTUN	A/m	0
PK1	Cross-term dependence of K1	$V^{1/2}m$	0
PK1W1	Cross-term dependence of K1W1	$m^2$	0
PK1W2	Cross-term dependence of K1W2	$m^2$	0
PK2	Cross-term dependence of K2	m	0
PK3	Cross-term dependence of K3	m	0
PK3B	Cross-term dependence of K3B	m/V	0
PKB1	Cross-term dependence of KB1	m	0
PKETA	Cross-term dependence of KETA	m/V	0
PKETAS	Cross-term dependence of KETAS	Vm	0

Table 2.65: BSIM3 SOI Device Model Parameters

Parameter	Description	Units	Default
PKT1	Cross-term dependence of KT1	Vm	0
PKT1L	Cross-term dependence of KT1L	Vm <sup>2</sup>	0
PKT2	Cross-term dependence of KT2	m	0
PLBJT0	Cross-term dependence of LBJT0	m <sup>2</sup>	0
PLII	Cross-term dependence of LII	m	0
PMOIN	Cross-term dependence of MOIN	m	0
PNBJT	Cross-term dependence of NBJT	m	0
PNCH	Cross-term dependence of NCH	m/cm <sup>3</sup>	0
PNDIF	Cross-term dependence of NDIF	m	0
PNDIODE	Cross-term dependence of NDIODE	m	0
PNFACTOR	Cross-term dependence of NFACTOR	m	0
PNGATE	Cross-term dependence of NGATE	m/cm <sup>3</sup>	0
PNGIDL	Cross-term dependence of NGIDL	Vm	0
PNIGC	Cross-term dependence of NIGC	m	0
PNLX	Cross-term dependence of NLX	m <sup>2</sup>	0
PNOFF	Cross-term dependence of NOFF	m	0
PNRECF0	Cross-term dependence of NRECF0	m	0
PNRECR0	Cross-term dependence of NRECR0	m	0
PNSUB	Cross-term dependence of NSUB	m/cm <sup>3</sup>	0
PNTRECF	Cross-term dependence of NTRECF	m	0
PNTRECR	Cross-term dependence of NTRECR	m	0
PNTUN	Cross-term dependence of NTUN	m	0
PPCLM	Cross-term dependence of PCLM	m	0
PPDIBLC1	Cross-term dependence of PDIBLC1	m	0
PPDIBLC2	Cross-term dependence of PDIBLC2	m	0
PPDIBLCB	Cross-term dependence of PDIBLCB	m/V	0
PPIGCD	Cross-term dependence of PIGCD	m	0
PPOXEDGE	Cross-term dependence of POXEDGE	m	0
PPRT	Cross-term dependence of PRT	$\Omega \mu\text{m m}$	0
PPRWB	Cross-term dependence of PRWB	m/V <sup>1/2</sup>	0
PPRWG	Cross-term dependence of PRWG	m/V	0
PPVAG	Cross-term dependence of PVAG	m	0
PRDSW	Cross-term dependence of RDSW	$\Omega \mu\text{m m}$	0
PSII0	Cross-term dependence of SII0	m/V	0
PSII1	Cross-term dependence of SII1	m/V	0

Table 2.65: BSIM3 SOI Device Model Parameters

Parameter	Description	Units	Default
PSII2	Cross-term dependence of SII2	m	0
PSIID	Cross-term dependence of SIID	m/V	0
PU0	Cross-term dependence of U0	m/(Vcm <sup>2</sup> s)0	0
PUA	Cross-term dependence of UA	m <sup>2</sup> /V	0
PUA1	Cross-term dependence of UA1	m <sup>2</sup> /V	0
PUB	Cross-term dependence of UB	m <sup>3</sup> /V <sup>2</sup>	0
PUB1	Cross-term dependence of UB1	m <sup>3</sup> /V <sup>2</sup>	0
PUC	Cross-term dependence of UC	m <sup>2</sup> /V <sup>2</sup>	0
PUC1	Cross-term dependence of UC1	m <sup>2</sup> /(°CV <sup>2</sup> )	0
PUTE	Cross-term dependence of UTE	m	0
PVABJT	Cross-term dependence of VABJT	Vm	0
PVDSATII0	Cross-term dependence of VDSATII0	Vm	0
PVOFF	Cross-term dependence of VOFF	Vm	0
PVREC0	Cross-term dependence of VREC0	Vm	0
PVSAT	Cross-term dependence of VSAT	m <sup>2</sup> /s	0
PVSDFB	Cross-term dependence of VSDFB	Vm	0
PVSDTH	Cross-term dependence of VSDTH	Vm	0
PVTH0	Cross-term dependence of VTH0	Vm	0
PVTUN0	Cross-term dependence of VTUN0	Vm	0
PW0	Cross-term dependence of W0	m <sup>2</sup>	0
PWR	Cross-term dependence of WR	m	0
PXBJT	Cross-term dependence of XBJT	m	0
PXDIF	Cross-term dependence of XDIF	m	0
PXJ	Cross-term dependence of XJ	m <sup>2</sup>	0
PXRCRG1	Cross-term dependence of XRCRG1	m	0
PXRCRG2	Cross-term dependence of XRCRG2	m	0
PXREC	Cross-term dependence of XREC	m	0
PXTUN	Cross-term dependence of XTUN	m	0
WA0	Width dependence of A0	m	0
WA1	Width dependence of A1	m/V	0
WA2	Width dependence of A2	m	0
WACDE	Width dependence of ACDE	m <sup>2</sup> /V	0
WAELY	Width dependence of AELY	V	0
WAGIDL	Width dependence of AGIDL	m/Ω	0
WAGS	Width dependence of AGS	m/V	0

Table 2.65: BSIM3 SOI Device Model Parameters

Parameter	Description	Units	Default
WAHLI	Width dependence of AHLI	m	0
WAIGC	Width dependence of AIGC	$(F/g)^{1/2} \text{sm}^2/\text{V}$	0
WAIGSD	Width dependence of AIGSD	$(F/g)^{1/2} \text{sm}^2/\text{V}$	0
WALPHA0	Width dependence of ALPHA0	$\text{m}^2/\text{V}$	0
WALPHAGB1	Width dependence of ALPHAGB1	m/V	0
WALPHAGB2	Width dependence of ALPHAGB2	m/V	0
WAT	Width dependence of AT	$\text{m}^2/\text{s}$	0
WBO	Width dependence of B0	$\text{m}^2$	0
WB1	Width dependence of B1	$\text{m}^2$	0
WBETA0	Width dependence of BETA0	Vm	0
WBETA1	Width dependence of BETA1	m	0
WBETA2	Width dependence of BETA2	Vm	0
WBETAGB1	Width dependence of BETAGB1	$\text{m}/\text{V}^2$	0
WBETAGB2	Width dependence of BETAGB2	$\text{m}/\text{V}^2$	0
WBGIDL	Width dependence of BGIDL	V	0
WBIGC	Width dependence of BIGC	$(F/g)^{1/2} \text{sm}^2/\text{V}$	0
WBIGSD	Width dependence of BIGSD	$(F/g)^{1/2} \text{sm}^2/\text{V}$	0
WCDSC	Width dependence of CDSC	F/m	0
WCDSCB	Width dependence of CDSCB	$\text{F}/(\text{Vm})$	0
WCDSCD	Width dependence of CDSCD	$\text{F}/(\text{Vm})$	0
WCGDL	Width dependence of CGDL	F	0
WCGSL	Width dependence of CGSL	F	0
WCIGC	Width dependence of CIGC	m/V	0
WCIGSD	Width dependence of CIGSD	m/V	0
WCIT	Width dependence of CIT	F/m	0
WCKAPPA	Width dependence of CKAPPA	F	0
WDELTA	Width dependence of DELTA	Vm	0
WDELVT	Width dependence of DELVT	Vm	0
WDROUT	Width dependence of DROUT	m	0
WDSUB	Width dependence of DSUB	m	0
WDVT0	Width dependence of DVT0	m	0
WDVT0W	Width dependence of DVT0W	–	0
WDVT1	Width dependence of DVT1	m	0
WDVT1W	Width dependence of DVT1W	–	0
WDVT2	Width dependence of DVT2	m/V	0

Table 2.65: BSIM3 SOI Device Model Parameters

Parameter	Description	Units	Default
WDVT2W	Width dependence of DVT2W	m/V	0
WDWB	Width dependence of DWB	$m^2/V^{1/2}$	0
WDWG	Width dependence of DWG	$m^2/V^{1/2}$	0
WESATII	Width dependence of ESATII	V	0
WETA0	Width dependence of ETA0	m	0
WETAB	Width dependence of ETAB	m/V	0
WFBJTII	Width dependence of FBJTII	m	0
WISBJT	Width dependence of ISBJT	A/m	0
WISDIF	Width dependence of ISDIF	A/m	0
WISREC	Width dependence of ISREC	A/m	0
WISTUN	Width dependence of ISTUN	A/m	0
WK1	Width dependence of K1	$V^{1/2}m$	0
WK1W1	Width dependence of K1W1	$m^2$	0
WK1W2	Width dependence of K1W2	$m^2$	0
WK2	Width dependence of K2	m	0
WK3	Width dependence of K3	m	0
WK3B	Width dependence of K3B	m/V	0
WKB1	Width dependence of KB1	m	0
WKETA	Width dependence of KETA	m/V	0
WKETAS	Width dependence of KETAS	Vm	0
WKT1	Width dependence of KT1	Vm	0
WKT1L	Width dependence of KT1L	$Vm^2$	0
WKT2	Width dependence of KT2	m	0
WLBJT0	Width dependence of LBJT0	$m^2$	0
WLII	Width dependence of LII	m	0
WMOIN	Width dependence of MOIN	m	0
WNBJT	Width dependence of NBJT	m	0
WNCH	Width dependence of NCH	$m/cm^3$	0
WNDIF	Width dependence of NDIF	m	0
WNDIODE	Width dependence of NDIODE	m	0
WNFACTOR	Width dependence of NFACTOR	m	0
WNGATE	Width dependence of NGATE	$m/cm^3$	0
WNGIDL	Width dependence of NGIDL	Vm	0
WNIGC	Width dependence of NIGC	m	0
WNLX	Width dependence of NLX	$m^2$	0

Table 2.65: BSIM3 SOI Device Model Parameters

Parameter	Description	Units	Default
WNOFF	Width dependence of NOFF	m	0
WNRECF0	Width dependence of NRECF0	m	0
WNRECR0	Width dependence of NRECR0	m	0
WNSUB	Width dependence of NSUB	m/cm <sup>3</sup>	0
WNTRECF	Width dependence of NTRECF	m	0
WNTRECR	Width dependence of NTRECR	m	0
WNTUN	Width dependence of NTUN	m	0
WPCLM	Width dependence of PCLM	m	0
WPDIBLC1	Width dependence of PDIBLC1	m	0
WPDIBLC2	Width dependence of PDIBLC2	m	0
WPDIBLCB	Width dependence of PDIBLCB	m/V	0
WPIGCD	Width dependence of PIGCD	m	0
WPOXEDGE	Width dependence of POXEDGE	m	0
WPRT	Width dependence of PRT	$\Omega \mu\text{m m}$	0
WPRWB	Width dependence of PRWB	$\text{m}/\text{V}^{1/2}$	0
WPRWG	Width dependence of PRWG	m/V	0
WPVAG	Width dependence of PVAG	m	0
WRDSW	Width dependence of RDSW	$\Omega \mu\text{m m}$	0
WSII0	Width dependence of SII0	m/V	0
WSII1	Width dependence of SII1	m/V	0
WSII2	Width dependence of SII2	m	0
WSIID	Width dependence of SIID	m/V	0
WU0	Width dependence of U0	$\text{m}/(\text{Vcm}^2\text{s})$	0
WUA	Width dependence of UA	$\text{m}^2/\text{V}$	0
WUA1	Width dependence of UA1	$\text{m}^2/\text{V}$	0
WUB	Width dependence of UB	$\text{m}^3/\text{V}^2$	0
WUB1	Width dependence of UB1	$\text{m}^3/\text{V}^2$	0
WUC	Width dependence of UC	$\text{m}^2/\text{V}^2$	0
WUC1	Width dependence of UC1	$\text{m}^2/({}^\circ\text{CV}^2)$	0
WUTE	Width dependence of UTE	m	0
WVABJT	Width dependence of VABJT	Vm	0
WVDSATII0	Width dependence of VDSATII0	Vm	0
WVOFF	Width dependence of VOFF	Vm	0
WVREC0	Width dependence of VREC0	Vm	0
WVSAT	Width dependence of VSAT	$\text{m}^2/\text{s}$	0

Table 2.65: BSIM3 SOI Device Model Parameters

Parameter	Description	Units	Default
WVSDFB	Width dependence of VSDFB	Vm	0
WVSDTH	Width dependence of VSDTH	Vm	0
WVTH0	Width dependence of VTH0	Vm	0
WVTUN0	Width dependence of VTUN0	Vm	0
WV0	Width dependence of W0	m <sup>2</sup>	0
WWR	Width dependence of WR	m	0
WXBJT	Width dependence of XBJT	m	0
WXDIF	Width dependence of XDIF	m	0
WXJ	Width dependence of XJ	m <sup>2</sup>	0
WXRCRG1	Width dependence of XRCRG1	m	0
WXRCRG2	Width dependence of XRCRG2	m	0
WXREC	Width dependence of XREC	m	0
WXTUN	Width dependence of XTUN	m	0
<b><i>Doping Parameters</i></b>			
NSUB	Substrate doping density	cm <sup>-3</sup>	6e+16
<b><i>Flicker Parameters</i></b>			
AF	Flicker noise exponent	–	1
EF	Flicker exponent	–	1
EM	Saturation field	V/m	4.1e+07
KF	Flicker noise coefficient	–	0
NOIA	Noise parameter a	–	0
NOIB	Noise parameter b	–	0
NOIC	Noise parameter c	–	8.75e+09
<b><i>Geometry Parameters</i></b>			
L	Channel length	m	5e-06
LL	Coefficient of length dependence for length offset	m <sup>LLN</sup>	0
LLC	Coefficient of length dependence for CV channel length offset	m <sup>LLN</sup>	0
LLN	Power of length dependence for length offset	–	1
LW	Coefficient of width dependence for length offset	m <sup>LWN</sup>	0
LWC	Coefficient of width dependence for channel length offset	m <sup>LWN</sup>	0
LWL	Coefficient of length and width cross term for length offset	m <sup>LLN+LWN</sup>	0
LWLC	Coefficient of length and width dependence for CV channel length offset	m <sup>LLN+LWN</sup>	0

Table 2.65: BSIM3 SOI Device Model Parameters

Parameter	Description	Units	Default
LWN	Power of width dependence for length offset	–	1
TOX	Gate oxide thickness	m	1e-08
W	Channel width	m	5e-06
WL	Coefficient of length dependence for width offset	$m^{WLN}$	0
WLC	Coefficient of length dependence for CV channel width offset	$m^{WLN}$	0
WLN	Power of length dependence of width offset	–	1
WW	Coefficient of width dependence for width offset	$m^{WWN}$	0
WWC	Coefficient of width dependence for CV channel width offset	$m^{WWN}$	0
WWL	Coefficient of length and width cross term for width offset	$m^{WLN+WWN}$	0
WWLC	Coefficient of length and width dependence for CV channel width offset	$m^{WLN+WWN}$	0
WWN	Power of width dependence of width offset	–	1
XJ	Junction depth	m	0
<b>Resistance Parameters</b>			
RSH	Drain, source diffusion sheet resistance	$\Omega$	0
<b>Process Parameters</b>			
GAMMA1	Body effect coefficient near the surface	$V^{1/2}$	0
GAMMA2	Body effect coefficient in the bulk	$V^{1/2}$	0
NCH	Channel doping concentration	$cm^{-3}$	1.7e+17
TBOX	Buried oxide thickness	m	3e-07
TOXM	Gate oxide thickness used in extraction	m	0
TSI	Silicon film thickness	m	1e-07
U0	Surface mobility	$1/(Vcm^2s)$	0
VBX	Vbs at which the depletion region = XT	V	0
XT	Doping depth	m	1.55e-07
<b>RF Parameters</b>			
BUG1830FIX	Voltage limiter fix for bug 1830	–	0
NGCON	Number of gate contacts	–	1
RGATEMOD	Gate resistance model selector	–	0
XGL	Offset of the gate length due to variations in patterning	m	0
XGW	Distance from the gate contact to the channel edge	m	0

Table 2.65: BSIM3 SOI Device Model Parameters

Parameter	Description	Units	Default
XRCRG1	Parameter for distributed channel resistance effect for intrinsic input resistance	—	12
XRCRG2	Parameter to account for the excess channel diffusion resistance for intrinsic input resistance	—	1
<b>Temperature Parameters</b>			
AT	Temperature coefficient for saturation velocity	m/s	33000
CTHO	Thermal capacitance per unit width	F/m	1e-05
KT1	Temperature coefficient for threshold voltage	V	-0.11
KT1L	Channel length dependence of the temperature coefficient for the threshold voltage	Vm	0
KT2	Body-bias coefficient to the threshold voltage temperature effect	—	0.022
NTRECF	Temperature coefficient for NRECF	—	0
NTRECR	Temperature coefficient for NRECR	—	0
PRT	Temperature coefficient for RDSW	$\Omega \mu\text{m}$	0
RTHO	Thermal resistance per unit width	$\Omega/\text{m}$	0
TCJSWG	Temperature coefficient of Cjswg	$\text{K}^{-1}$	0
TNOM	Nominal device temperature	$^{\circ}\text{C}$	Ambient Temperature
TPBSWG	Temperature coefficient of Pbswg	V/K	0
UA1	Temperature coefficient for UA	m/V	4.31e-09
UB1	Temperature coefficient for UB	$\text{m}^2/\text{V}^2$	-7.61e-18
UC1	Temperature coefficient for UC	$\text{m}/(^{\circ}\text{C V}^2)$	0
UTE	Mobility temperature exponent	—	-1.5
WTHO	Minimum width for thermal resistance calculation	m	0
XBJT	Power dependence of JBJT on temperature	—	1
XDIF	Power dependence of JDIF on temperature	—	0
XREC	Power dependence of JREC on temperature	—	1
XTUN	Power dependence of JTUN on temperature	—	0
<b>Tunnelling Parameters</b>			
ALPHAGB1	First Vox dependent parameter for gate current in inversion	$\text{V}^{-1}$	0.35
ALPHAGB2	First Vox dependent parameter for gate current in accumulation	$\text{V}^{-1}$	0.43
BETAGB1	Second Vox dependent parameter for gate current in inversion	$\text{V}^{-2}$	0.03

Table 2.65: BSIM3 SOI Device Model Parameters

Parameter	Description	Units	Default
BETAGB2	First Vox dependent parameter for gate current in accumulation	$V^{-2}$	0.05
EBG	Effective bandgap in gate current calculation	V	1.2
IGMOD	Gate current model selector	–	0
NTOX	Power term of gate current	–	1
TOXQM	Oxide thickness for Igb calculation	m	0
TOXREF	Target oxide thickness	m	2.5e-09
VECB	Vaux parameter for conduction band electron tunneling	–	0.026
VEVB	Vaux parameter for valence band electron tunneling	–	0.075
VGB1	Third Vox dependent parameter for gate current in inversion	V	300
VGB2	Third Vox dependent parameter for gate current in accumulation	V	17
<b><i>Built-in Potential Lowering Parameters</i></b>			
DK2B	Third backgate body effect parameter for short channel effect	–	0
DVBD0	First short channel effect parameter in FD module	–	0
DVBD1	Second short channel effect parameter in FD module	–	0
K1B	First backgate body effect parameter	–	1
K2B	Second backgate body effect parameter for short channel effect	–	0
MOINFD	Gate bias dependance coefficient of surface potential in FD module	–	1000
NOFFFD	Smoothing parameter in FD module	–	1
SOIMOD	SIO model selector, SOIMOD=0: BSIMPD, SOIMOD=1: undefined model for PD and FE, SOIMOD=2: ideal FD	–	0
VBSA	Offset voltage due to non-idealities	V	0
VOFFFD	Smoothing parameter in FD module	V	0

## Level 14 MOSFET Tables (BSIM4)

For complete documentation of the BSIM4 model, see the users' manual for the BSIM4, available for download at <http://www-device.eecs.berkeley.edu/bsim/?page=BSIM4>. **Xyce** implements Version 4.6.1 of the BSIM4, you will have to get the documentation from the FTP archive on the Berkeley site.

The level 14 MOSFET device in **Xyce** is based on the Berkeley BSIM4 model version 4.6.1. Its parameters are given in the following tables. Note that the table is not yet in its final form and parameters have not all been properly categorized with units in place. For correct units, see the BSIM4 documentation available at the BSIM group's web site, <http://www-device.eecs.berkeley.edu/bsim/>.

Table 2.66: BSIM4 Device Instance Parameters

Parameter	Description	Units	Default
AD	Drain area	—	0
AS	Source area	—	0
IC2		—	0
IC3		—	0
L	Length	—	5e-06
M	Number of parallel copies	—	1
MIN	Minimize either D or S	—	0
NF	Number of fingers	—	1
NGCON	Number of gate contacts	—	0
OFF	Device is initially off	—	false
PD	Drain perimeter	—	0
PS	Source perimeter	—	0
RBDB	Body resistance	—	0
RBPB	Body resistance	—	0
RBPD	Body resistance	—	0
RBPS	Body resistance	—	0
RBSB	Body resistance	—	0
SA	distance between OD edge to poly of one side	—	0
SB	distance between OD edge to poly of the other side	—	0
SC	Distance to a single well edge	—	0
SCA	Integral of the first distribution function for scattered well dopant	—	0
SCB	Integral of the second distribution function for scattered well dopant	—	0

Table 2.66: BSIM4 Device Instance Parameters

Parameter	Description	Units	Default
SCC	Integral of the third distribution function for scattered well dopant	–	0
SD	distance between neighbour fingers	–	0
W	Width	–	5e-06
XGW	Distance from gate contact center to device edge	–	0
<b>Basic Parameters</b>			
DELVT0	Zero bias threshold voltage variation	V	0
<b>Control Parameters</b>			
ACNQSMOD	AC NQS model selector	–	0
GEOMOD	Geometry dependent parasitics model selector	–	0
RBODYMOD	Distributed body R model selector	–	0
RGATEMOD	Gate resistance model selector	–	0
RGEOMOD	S/D resistance and contact model selector	–	0
TRNQSMOD	Transient NQS model selector	–	0
<b>Temperature Parameters</b>			
TEMP	Device temperature	°C	Ambient Temperature
<b>Voltage Parameters</b>			
IC1	Vector of initial values: Vds, Vgs, Vbs	V	0
<b>Asymmetric and Bias-Dependent <math>R_{ds}</math> Parameters</b>			
NRD	Number of squares in drain	–	1
NRS	Number of squares in source	–	1

Table 2.67: BSIM4 Device Model Parameters

Parameter	Description	Units	Default
AF	Flicker noise exponent	–	1
AIGSD	Parameter for $I_{gs,d}$	–	0.0136
AT	Temperature coefficient of $v_{sat}$	–	33000
BIGSD	Parameter for $I_{gs,d}$	–	0.00171
BVD	Drain diode breakdown voltage	–	10
BVS	Source diode breakdown voltage	–	10
CIGSD	Parameter for $I_{gs,d}$	–	0.075
CJD	Drain bottom junction capacitance per unit area	–	0.0005
CJS	Source bottom junction capacitance per unit area	–	0.0005

Table 2.67: BSIM4 Device Model Parameters

Parameter	Description	Units	Default
CJSWD	Drain sidewall junction capacitance per unit periphery	—	5e-10
CJSWGD	Drain (gate side) sidewall junction capacitance per unit width	—	0
CJSWGS	Source (gate side) sidewall junction capacitance per unit width	—	0
CJSWS	Source sidewall junction capacitance per unit periphery	—	5e-10
DLGIG	Delta L for Ig model	—	0
DMCG	Distance of Mid-Contact to Gate edge	—	0
DMCGT	Distance of Mid-Contact to Gate edge in Test structures	—	0
DMCI	Distance of Mid-Contact to Isolation	—	0
DMDG	Distance of Mid-Diffusion to Gate edge	—	0
DWJ	Delta W for S/D junctions	—	0
EF	Flicker noise frequency exponent	—	1
EM	Flicker noise parameter	—	4.1e+07
EPSRGATE	Dielectric constant of gate relative to vacuum	—	11.7
GBMIN	Minimum body conductance	$\Omega^{-1}$	1e-12
IJTHDFWD	Forward drain diode forward limiting current	—	0.1
IJTHDREV	Reverse drain diode forward limiting current	—	0.1
IJTHSFWD	Forward source diode forward limiting current	—	0.1
IJTHSREV	Reverse source diode forward limiting current	—	0.1
JSD	Bottom drain junction reverse saturation current density	—	0.0001
JSS	Bottom source junction reverse saturation current density	—	0.0001
JSWD	Isolation edge sidewall drain junction reverse saturation current density	—	0
JSWGD	Gate edge drain junction reverse saturation current density	—	0
JSWGS	Gate edge source junction reverse saturation current density	—	0
JSWS	Isolation edge sidewall source junction reverse saturation current density	—	0
JTSD	Drain bottom trap-assisted saturation current density	—	0
JTSS	Source bottom trap-assisted saturation current density	—	0

Table 2.67: BSIM4 Device Model Parameters

Parameter	Description	Units	Default
JTSSWD	Drain STI sidewall trap-assisted saturation current density	—	0
JTSSWGD	Drain gate-edge sidewall trap-assisted saturation current density	—	0
JTSSWGS	Source gate-edge sidewall trap-assisted saturation current density	—	0
JTSSWS	Source STI sidewall trap-assisted saturation current density	—	0
K2WE	K2 shift factor for well proximity effect	—	0
K3B	Body effect coefficient of k3	—	0
KF	Flicker noise coefficient	—	0
KT1	Temperature coefficient of Vth	—	-0.11
KT1L	Temperature coefficient of Vth	—	0
KT2	Body-coefficient of kt1	—	0.022
KU0	Mobility degradation/enhancement coefficient for LOD	—	0
KUOWE	Mobility degradation factor for well proximity effect	—	0
KVSAT	Saturation velocity degradation/enhancement parameter for LOD	—	0
KVTH0	Threshold degradation/enhancement parameter for LOD	—	0
KVTHWE	Threshold shift factor for well proximity effect	—	0
LA0	Length dependence of a0	—	0
LA1	Length dependence of a1	—	0
LA2	Length dependence of a2	—	0
LACDE	Length dependence of acde	—	0
LAGIDL	Length dependence of agidl	—	0
LAGISL	Length dependence of agisl	—	0
LAGS	Length dependence of ags	—	0
LAIGBACC	Length dependence of aigbacc	—	0
LAIGBINV	Length dependence of aigbinv	—	0
LAIGC	Length dependence of aigc	—	0
LAIGD	Length dependence of aigd	—	0
LAIGS	Length dependence of aigs	—	0
LAIGSD	Length dependence of aigsd	—	0
LALPHA0	Length dependence of alpha0	—	0

Table 2.67: BSIM4 Device Model Parameters

Parameter	Description	Units	Default
LALPHA1	Length dependence of alpha1	—	0
LAT	Length dependence of at	—	0
LB0	Length dependence of b0	—	0
LB1	Length dependence of b1	—	0
LBETA0	Length dependence of beta0	—	0
LBGIDL	Length dependence of bgidl	—	0
LBGISL	Length dependence of bgisl	—	0
LBIGBACC	Length dependence of bigbacc	—	0
LBIGBINV	Length dependence of bigbinv	—	0
LBIGC	Length dependence of bigc	—	0
LBIGD	Length dependence of bigd	—	0
LBIGS	Length dependence of bigs	—	0
LBIGSD	Length dependence of bigsd	—	0
LCDSC	Length dependence of cdsc	—	0
LCDSCB	Length dependence of cdsch	—	0
LCDSCD	Length dependence of cdsd	—	0
LCF	Length dependence of cf	—	0
LCGDL	Length dependence of cgdl	—	0
LCGIDL	Length dependence of cgidl	—	0
LCGISL	Length dependence of cgisl	—	0
LCGSL	Length dependence of cgsl	—	0
LCIGBACC	Length dependence of cigbacc	—	0
LCIGBINV	Length dependence of cigbinv	—	0
LCIGC	Length dependence of cigc	—	0
LCIGD	Length dependence of cigd	—	0
LCIGS	Length dependence of cigs	—	0
LCIGSD	Length dependence of cigsd	—	0
LCIT	Length dependence of cit	—	0
LCKAPPAD	Length dependence of ckappad	—	0
LCKAPPAS	Length dependence of ckappas	—	0
LCLC	Length dependence of clc	—	0
LCLE	Length dependence of cle	—	0
LDELTA	Length dependence of delta	—	0
LDROUT	Length dependence of drout	—	0
LDSUB	Length dependence of dsub	—	0

Table 2.67: BSIM4 Device Model Parameters

Parameter	Description	Units	Default
LDVT0	Length dependence of dvt0	—	0
LDVT0W	Length dependence of dvt0w	—	0
LDVT1	Length dependence of dvt1	—	0
LDVT1W	Length dependence of dvt1w	—	0
LDVT2	Length dependence of dvt2	—	0
LDVT2W	Length dependence of dvt2w	—	0
LDVTP0	Length dependence of dvtp0	—	0
LDVTP1	Length dependence of dvtp1	—	0
LDWB	Length dependence of dwb	—	0
LDWG	Length dependence of dwg	—	0
LEGIDL	Length dependence of egidl	—	0
LEGISL	Length dependence of egisl	—	0
LEIGBINV	Length dependence for eigbinv	—	0
LETA0	Length dependence of eta0	—	0
LETAB	Length dependence of etab	—	0
LEU	Length dependence of eu	—	0
LFPROUT	Length dependence of pdiblcb	—	0
LGAMMA1	Length dependence of gamma1	—	0
LGAMMA2	Length dependence of gamma2	—	0
LINTNOI	lint offset for noise calculation	—	0
LK1	Length dependence of k1	—	0
LK2	Length dependence of k2	—	0
LK2WE	Length dependence of k2we	—	0
LK3	Length dependence of k3	—	0
LK3B	Length dependence of k3b	—	0
LKETA	Length dependence of keta	—	0
LKT1	Length dependence of kt1	—	0
LKT1L	Length dependence of kt1l	—	0
LKT2	Length dependence of kt2	—	0
LKU0	Length dependence of ku0	—	0
LKU0WE	Length dependence of ku0we	—	0
LKVTH0	Length dependence of kvth0	—	0
LKVTH0WE	Length dependence of kvth0we	—	0
LL	Length reduction parameter	—	0
LLAMBDA	Length dependence of lambda	—	0

Table 2.67: BSIM4 Device Model Parameters

Parameter	Description	Units	Default
LLC	Length reduction parameter for CV	—	0
LLN	Length reduction parameter	—	1
LLODKU0	Length parameter for u0 LOD effect	—	0
LLODVTH	Length parameter for vth LOD effect	—	0
LLP	Length dependence of lp	—	0
LLPE0	Length dependence of lpe0	—	0
LLPEB	Length dependence of lpeb	—	0
LMAX	Maximum length for the model	—	1
LMIN	Minimum length for the model	—	0
LMINV	Length dependence of minv	—	0
LMINVCV	Length dependence of minvcv	—	0
LMOIN	Length dependence of moin	—	0
LNDEP	Length dependence of ndep	—	0
LNFACTOR	Length dependence of nfactor	—	0
LNGATE	Length dependence of ngate	—	0
LNIGBACC	Length dependence of nighbacc	—	0
LNIGBINV	Length dependence of nighbinv	—	0
LNIGC	Length dependence of nignc	—	0
LNOFF	Length dependence of noff	—	0
LNSD	Length dependence of nsd	—	0
LNSUB	Length dependence of nsub	—	0
LNTOX	Length dependence of ntox	—	0
LODETA0	eta0 shift modification factor for stress effect	—	1
LODK2	K2 shift modification factor for stress effect	—	1
LPCLM	Length dependence of pclm	—	0
LPDIBLC1	Length dependence of pdiblc1	—	0
LPDIBLC2	Length dependence of pdiblc2	—	0
LPDIBLCB	Length dependence of pdiblcb	—	0
LPDITS	Length dependence of pdits	—	0
LPDITSD	Length dependence of pditsd	—	0
LPHIN	Length dependence of phin	—	0
LPIGCD	Length dependence for pigcd	—	0
LPOXEDGE	Length dependence for poxedge	—	0
LPRT	Length dependence of prt	—	0
LPRWB	Length dependence of prwb	—	0

Table 2.67: BSIM4 Device Model Parameters

Parameter	Description	Units	Default
LPRWG	Length dependence of prwg	—	0
LPSCBE1	Length dependence of pscbe1	—	0
LPSCBE2	Length dependence of pscbe2	—	0
LPVAG	Length dependence of pvag	—	0
LRDSW	Length dependence of rdsw	—	0
LRDW	Length dependence of rdw	—	0
LRSW	Length dependence of rsw	—	0
LTVFBSDOFF	Length dependence of tvfbsdoff	—	0
LTVOFF	Length dependence of tvoff	—	0
LU0	Length dependence of u0	—	0
LUA	Length dependence of ua	—	0
LUA1	Length dependence of ua1	—	0
LUB	Length dependence of ub	—	0
LUB1	Length dependence of ub1	—	0
LUC	Length dependence of uc	—	0
LUC1	Length dependence of uc1	—	0
LUD	Length dependence of ud	—	0
LUD1	Length dependence of ud1	—	0
LUP	Length dependence of up	—	0
LUTE	Length dependence of ute	—	0
LVBM	Length dependence of vbm	—	0
LVBX	Length dependence of vbx	—	0
LVFB	Length dependence of vfb	—	0
LVFBCV	Length dependence of vfbcv	—	0
LVFBSDOFF	Length dependence of vfbsdoff	—	0
LVOFF	Length dependence of voff	—	0
LVOFFCV	Length dependence of voffcv	—	0
LVSAT	Length dependence of vsat	—	0
LVTHO		—	0
LVTL	Length dependence of vtl	—	0
LW	Length reduction parameter	—	0
LW0	Length dependence of w0	—	0
LWC	Length reduction parameter for CV	—	0
LWL	Length reduction parameter	—	0
LWLC	Length reduction parameter for CV	—	0

Table 2.67: BSIM4 Device Model Parameters

Parameter	Description	Units	Default
LWN	Length reduction parameter	—	1
LWR	Length dependence of $w_r$	—	0
LXJ	Length dependence of $x_j$	—	0
LXN	Length dependence of $x_n$	—	0
LXRCRG1	Length dependence of $xrcrg1$	—	0
LXRCRG2	Length dependence of $xrcrg2$	—	0
LXT	Length dependence of $x_t$	—	0
MJD	Drain bottom junction capacitance grading coefficient	—	0.5
MJS	Source bottom junction capacitance grading coefficient	—	0.5
MJSWD	Drain sidewall junction capacitance grading coefficient	—	0.33
MJSWGD	Drain (gate side) sidewall junction capacitance grading coefficient	—	0.33
MJSWGS	Source (gate side) sidewall junction capacitance grading coefficient	—	0.33
MJSWS	Source sidewall junction capacitance grading coefficient	—	0.33
NGCON	Number of gate contacts	—	1
NJD	Drain junction emission coefficient	—	1
NJS	Source junction emission coefficient	—	1
NJTS	Non-ideality factor for bottom junction	—	20
NJTSD	Non-ideality factor for bottom junction drain side	—	20
NJTSSW	Non-ideality factor for STI sidewall junction	—	20
NJTSSWD	Non-ideality factor for STI sidewall junction drain side	—	20
NJTSSWG	Non-ideality factor for gate-edge sidewall junction	—	20
NJTSSWGD	Non-ideality factor for gate-edge sidewall junction drain side	—	20
NTNOI	Thermal noise parameter	—	1
PA0	Cross-term dependence of $a_0$	—	0
PA1	Cross-term dependence of $a_1$	—	0
PA2	Cross-term dependence of $a_2$	—	0
PACDE	Cross-term dependence of $acde$	—	0
PAGIDL	Cross-term dependence of $agidl$	—	0
PAGISL	Cross-term dependence of $agisl$	—	0

Table 2.67: BSIM4 Device Model Parameters

Parameter	Description	Units	Default
PAGS	Cross-term dependence of ags	—	0
PAIGBACC	Cross-term dependence of aigbacc	—	0
PAIGBINV	Cross-term dependence of aigbinv	—	0
PAIGC	Cross-term dependence of aigc	—	0
PAIGD	Cross-term dependence of aigd	—	0
PAIGS	Cross-term dependence of aigs	—	0
PAIGSD	Cross-term dependence of aigsd	—	0
PALPHA0	Cross-term dependence of alpha0	—	0
PALPHA1	Cross-term dependence of alpha1	—	0
PAT	Cross-term dependence of at	—	0
PB0	Cross-term dependence of b0	—	0
PB1	Cross-term dependence of b1	—	0
PBD	Drain junction built-in potential	—	1
PBETA0	Cross-term dependence of beta0	—	0
PBGIDL	Cross-term dependence of bgidl	—	0
PBGISL	Cross-term dependence of bgisl	—	0
PBIGBACC	Cross-term dependence of bigbacc	—	0
PBIGBINV	Cross-term dependence of bigbinv	—	0
PBIGC	Cross-term dependence of bigc	—	0
PBIGD	Cross-term dependence of bigd	—	0
PBIGS	Cross-term dependence of bigs	—	0
PBIGSD	Cross-term dependence of bigsd	—	0
PBS	Source junction built-in potential	—	1
PBSWD	Drain sidewall junction capacitance built in potential	—	1
PBSWGD	Drain (gate side) sidewall junction capacitance built in potential	—	0
PBSWGS	Source (gate side) sidewall junction capacitance built in potential	—	0
PBSWS	Source sidewall junction capacitance built in potential	—	1
PCDSC	Cross-term dependence of cdsc	—	0
PCDSCB	Cross-term dependence of cdsch	—	0
PCDSCD	Cross-term dependence of cdsd	—	0
PCF	Cross-term dependence of cf	—	0
PCGDL	Cross-term dependence of cgdl	—	0

Table 2.67: BSIM4 Device Model Parameters

Parameter	Description	Units	Default
PCGIDL	Cross-term dependence of cgidl	—	0
PCGISL	Cross-term dependence of cgisl	—	0
PCGSL	Cross-term dependence of cgsl	—	0
PCIGBACC	Cross-term dependence of cigbacc	—	0
PCIGBINV	Cross-term dependence of cigbinv	—	0
PCIGC	Cross-term dependence of cigc	—	0
PCIGD	Cross-term dependence of cigd	—	0
PCIGS	Cross-term dependence of cigs	—	0
PCIGSD	Cross-term dependence of cigsd	—	0
PCIT	Cross-term dependence of cit	—	0
PCKAPPAD	Cross-term dependence of ckappad	—	0
PCKAPPAS	Cross-term dependence of ckappas	—	0
PCLC	Cross-term dependence of clc	—	0
PCLE	Cross-term dependence of cle	—	0
PDELTA	Cross-term dependence of delta	—	0
PDROUT	Cross-term dependence of drout	—	0
PDSUB	Cross-term dependence of dsub	—	0
PDVT0	Cross-term dependence of dvt0	—	0
PDVT0W	Cross-term dependence of dvt0w	—	0
PDVT1	Cross-term dependence of dvt1	—	0
PDVT1W	Cross-term dependence of dvt1w	—	0
PDVT2	Cross-term dependence of dvt2	—	0
PDVT2W	Cross-term dependence of dvt2w	—	0
PDVTP0	Cross-term dependence of dvtp0	—	0
PDVTP1	Cross-term dependence of dvtp1	—	0
PDWB	Cross-term dependence of dwb	—	0
PDWG	Cross-term dependence of dwg	—	0
PEGIDL	Cross-term dependence of egidl	—	0
PEGISL	Cross-term dependence of egisl	—	0
PEIGBINV	Cross-term dependence for eigbinv	—	0
PETA0	Cross-term dependence of eta0	—	0
PETAB	Cross-term dependence of etab	—	0
PEU	Cross-term dependence of eu	—	0
PFPROUT	Cross-term dependence of pdiblcb	—	0
PGAMMA1	Cross-term dependence of gamma1	—	0

Table 2.67: BSIM4 Device Model Parameters

Parameter	Description	Units	Default
PGAMMA2	Cross-term dependence of gamma2	—	0
PHIG	Work Function of gate	—	4.05
PK1	Cross-term dependence of k1	—	0
PK2	Cross-term dependence of k2	—	0
PK2WE	Cross-term dependence of k2we	—	0
PK3	Cross-term dependence of k3	—	0
PK3B	Cross-term dependence of k3b	—	0
PKETA	Cross-term dependence of keta	—	0
PKT1	Cross-term dependence of kt1	—	0
PKT1L	Cross-term dependence of kt1l	—	0
PKT2	Cross-term dependence of kt2	—	0
PKU0	Cross-term dependence of ku0	—	0
PKU0WE	Cross-term dependence of ku0we	—	0
PKVTH0	Cross-term dependence of kvth0	—	0
PKVTH0WE	Cross-term dependence of kvth0we	—	0
PLAMBDA	Cross-term dependence of lambda	—	0
PLP	Cross-term dependence of lp	—	0
PLPE0	Cross-term dependence of lpe0	—	0
PLPEB	Cross-term dependence of lpeb	—	0
PMINV	Cross-term dependence of minv	—	0
PMINVCV	Cross-term dependence of minvcv	—	0
PMOIN	Cross-term dependence of moin	—	0
PNDEP	Cross-term dependence of ndep	—	0
PNFACTOR	Cross-term dependence of nfactor	—	0
PNGATE	Cross-term dependence of ngate	—	0
PNIGBACC	Cross-term dependence of nigbacc	—	0
PNIGBINV	Cross-term dependence of nigbinv	—	0
PNIGC	Cross-term dependence of nignc	—	0
PNOFF	Cross-term dependence of noff	—	0
PNSD	Cross-term dependence of nsd	—	0
PNSUB	Cross-term dependence of nsub	—	0
PNTOX	Cross-term dependence of ntox	—	0
PPCLM	Cross-term dependence of pclm	—	0
PPDIBLC1	Cross-term dependence of pdiblc1	—	0
PPDIBLC2	Cross-term dependence of pdiblc2	—	0

Table 2.67: BSIM4 Device Model Parameters

Parameter	Description	Units	Default
PPDIBLCB	Cross-term dependence of pdiblcb	—	0
PPDITS	Cross-term dependence of pdits	—	0
PPDITSD	Cross-term dependence of pditsd	—	0
PPHIN	Cross-term dependence of phin	—	0
PPIGCD	Cross-term dependence for pigcd	—	0
PPOXEDGE	Cross-term dependence for poxedg	—	0
PPRT	Cross-term dependence of prt	—	0
PPRWB	Cross-term dependence of prwb	—	0
PPRWG	Cross-term dependence of prwg	—	0
PPSCBE1	Cross-term dependence of pscbe1	—	0
PPSCBE2	Cross-term dependence of pscbe2	—	0
PPVAG	Cross-term dependence of pvag	—	0
PRDSW	Cross-term dependence of rdsw	—	0
PRDW	Cross-term dependence of rdw	—	0
PRSW	Cross-term dependence of rsw	—	0
PRT	Temperature coefficient of parasitic resistance	—	0
PTVFBSDOFF	Cross-term dependence of tvfbsdoff	—	0
PTVOFF	Cross-term dependence of tvoff	—	0
PU0	Cross-term dependence of u0	—	0
PUA	Cross-term dependence of ua	—	0
PUA1	Cross-term dependence of ua1	—	0
PUB	Cross-term dependence of ub	—	0
PUB1	Cross-term dependence of ub1	—	0
PUC	Cross-term dependence of uc	—	0
PUC1	Cross-term dependence of uc1	—	0
PUD	Cross-term dependence of ud	—	0
PUD1	Cross-term dependence of ud1	—	0
PUP	Cross-term dependence of up	—	0
PUTE	Cross-term dependence of ute	—	0
PVAG	Gate dependence of output resistance parameter	—	0
PVBM	Cross-term dependence of vbm	—	0
PVBX	Cross-term dependence of vbv	—	0
PVFB	Cross-term dependence of vfb	—	0
PVFBCV	Cross-term dependence of vfbcv	—	0
PVFBSDOFF	Cross-term dependence of vfbsdoff	—	0

Table 2.67: BSIM4 Device Model Parameters

Parameter	Description	Units	Default
PVOFF	Cross-term dependence of voff	—	0
PVOFFCV	Cross-term dependence of voffcv	—	0
PVSAT	Cross-term dependence of vsat	—	0
PVTH0		—	0
PVTL	Cross-term dependence of vtl	—	0
PW0	Cross-term dependence of w0	—	0
PWR	Cross-term dependence of wr	—	0
PXJ	Cross-term dependence of xj	—	0
PXN	Cross-term dependence of xn	—	0
PXRCRG1	Cross-term dependence of xrcrg1	—	0
PXRCRG2	Cross-term dependence of xrcrg2	—	0
PXT	Cross-term dependence of xt	—	0
RBDB	Resistance between bNode and dbNode	$\Omega$	50
RDBX0	Body resistance RDBX scaling	—	100
RDBY0	Body resistance RDBY scaling	—	100
RBPB	Resistance between bNodePrime and bNode	$\Omega$	50
RBPBX0	Body resistance RBPBX scaling	—	100
RBPBXL	Body resistance RBPBX L scaling	—	0
RBPBXNF	Body resistance RBPBX NF scaling	—	0
RBPBXW	Body resistance RBPBX W scaling	—	0
RBPBY0	Body resistance RBPBY scaling	—	100
RBPBYL	Body resistance RBPBY L scaling	—	0
RBPBYNF	Body resistance RBPBY NF scaling	—	0
RBPBYW	Body resistance RBPBY W scaling	—	0
RBPDP	Resistance between bNodePrime and bNode	$\Omega$	50
RBPDO	Body resistance RBPDP scaling	—	50
RBPDL	Body resistance RBPDP L scaling	—	0
RBPDPNF	Body resistance RBPDP NF scaling	—	0
RBPDPW	Body resistance RBPDP W scaling	—	0
RBPS	Resistance between bNodePrime and sbNode	$\Omega$	50
RBPS0	Body resistance RBPS scaling	—	50
RBPSL	Body resistance RBPS L scaling	—	0
RBPSNF	Body resistance RBPS NF scaling	—	0
RBPSW	Body resistance RBPS W scaling	—	0
RBSB	Resistance between bNode and sbNode	$\Omega$	50

Table 2.67: BSIM4 Device Model Parameters

Parameter	Description	Units	Default
RBSBX0	Body resistance RBSBX scaling	—	100
RBSBY0	Body resistance RBSBY scaling	—	100
RBSDBXL	Body resistance RBSDBX L scaling	—	0
RBSDBXNF	Body resistance RBSDBX NF scaling	—	0
RBSDBXW	Body resistance RBSDBX W scaling	—	0
RBSDBYL	Body resistance RBSDBY L scaling	—	0
RBSDBYNF	Body resistance RBSDBY NF scaling	—	0
RBSDBYW	Body resistance RBSDBY W scaling	—	0
RNOIA	Thermal noise coefficient	—	0.577
RNOIB	Thermal noise coefficient	—	0.5164
SAREF	Reference distance between OD edge to poly of one side	—	1e-06
SBREF	Reference distance between OD edge to poly of the other side	—	1e-06
SCREF	Reference distance to calculate SCA, SCB and SCC	—	1e-06
STETA0	eta0 shift factor related to stress effect on vth	—	0
STK2	K2 shift factor related to stress effect on vth	—	0
TCJ	Temperature coefficient of cj	—	0
TCJSW	Temperature coefficient of cjsw	—	0
TCJSWG	Temperature coefficient of cjswg	—	0
TKU0	Temperature coefficient of KU0	—	0
TNJTS	Temperature coefficient for NJTS	—	0
TNJTSD	Temperature coefficient for NJTSD	—	0
TNJTSSW	Temperature coefficient for NJTSSW	—	0
TNJTSSWD	Temperature coefficient for NJTSSWD	—	0
TNJTSSWG	Temperature coefficient for NJTSSWG	—	0
TNJTSSWGD	Temperature coefficient for NJTSSWGD	—	0
TNOIA	Thermal noise parameter	—	1.5
TNOIB	Thermal noise parameter	—	3.5
TNOM	Parameter measurement temperature	—	Ambient Temperature
TPB	Temperature coefficient of pb	—	0
TPBSW	Temperature coefficient of pbsw	—	0
TPBSWG	Temperature coefficient of pbswg	—	0

Table 2.67: BSIM4 Device Model Parameters

Parameter	Description	Units	Default
TVFBSDOFF	Temperature parameter for vfbsdoff	—	0
TVOFF	Temperature parameter for voff	—	0
UA1	Temperature coefficient of ua	—	1e-09
UB1	Temperature coefficient of ub	—	-1e-18
UC1	Temperature coefficient of uc	—	0
UD1	Temperature coefficient of ud	—	0
UTE	Temperature coefficient of mobility	—	-1.5
VTSD	Drain bottom trap-assisted voltage dependent parameter	—	10
VTSS	Source bottom trap-assisted voltage dependent parameter	—	10
VTSSWD	Drain STI sidewall trap-assisted voltage dependent parameter	—	10
VTSSWGD	Drain gate-edge sidewall trap-assisted voltage dependent parameter	—	10
VTSSWGS	Source gate-edge sidewall trap-assisted voltage dependent parameter	—	10
VTSSWS	Source STI sidewall trap-assisted voltage dependent parameter	—	10
WA0	Width dependence of a0	—	0
WA1	Width dependence of a1	—	0
WA2	Width dependence of a2	—	0
WACDE	Width dependence of acde	—	0
WAGIDL	Width dependence of agidl	—	0
WAGISL	Width dependence of agisl	—	0
WAGS	Width dependence of ags	—	0
WAIGBACC	Width dependence of aigbacc	—	0
WAIGBINV	Width dependence of aigbinv	—	0
WAIGC	Width dependence of aigc	—	0
WAIGD	Width dependence of aigd	—	0
WAIGS	Width dependence of aigs	—	0
WAIGSD	Width dependence of aigsd	—	0
WALPHA0	Width dependence of alpha0	—	0
WALPHA1	Width dependence of alpha1	—	0
WAT	Width dependence of at	—	0
WB0	Width dependence of b0	—	0
WB1	Width dependence of b1	—	0

Table 2.67: BSIM4 Device Model Parameters

Parameter	Description	Units	Default
WBETA0	Width dependence of beta0	—	0
WBGIDL	Width dependence of bgidl	—	0
WBGISL	Width dependence of bgisl	—	0
WBIGBACC	Width dependence of bigbacc	—	0
WBIGBINV	Width dependence of bigbinv	—	0
WBIGC	Width dependence of bigc	—	0
WBIGD	Width dependence of bigd	—	0
WBIGS	Width dependence of bigs	—	0
WBIGSD	Width dependence of bigsd	—	0
WCDSC	Width dependence of cdsc	—	0
WCDSCB	Width dependence of cdscb	—	0
WCDSCD	Width dependence of cdscd	—	0
WCF	Width dependence of cf	—	0
WCGDL	Width dependence of cgdl	—	0
WCGIDL	Width dependence of cgidl	—	0
WCGISL	Width dependence of cgisl	—	0
WCGSL	Width dependence of cgsl	—	0
WCIGBACC	Width dependence of cigbacc	—	0
WCIGBINV	Width dependence of cigbinv	—	0
WCIGC	Width dependence of cigc	—	0
WCIGD	Width dependence of cigd	—	0
WCIGS	Width dependence of cigs	—	0
WCIGSD	Width dependence of cigsd	—	0
WCIT	Width dependence of cit	—	0
WCKAPPAD	Width dependence of ckappad	—	0
WCKAPPAS	Width dependence of ckappas	—	0
WCLC	Width dependence of clc	—	0
WCLE	Width dependence of cle	—	0
WDELTA	Width dependence of delta	—	0
WDROUT	Width dependence of drout	—	0
WDSUB	Width dependence of dsub	—	0
WDVT0	Width dependence of dvt0	—	0
WDVT0W	Width dependence of dvt0w	—	0
WDVT1	Width dependence of dvt1	—	0
WDVT1W	Width dependence of dvt1w	—	0

Table 2.67: BSIM4 Device Model Parameters

Parameter	Description	Units	Default
WDVT2	Width dependence of dvt2	—	0
WDVT2W	Width dependence of dvt2w	—	0
WDVTP0	Width dependence of dvtp0	—	0
WDVTP1	Width dependence of dvtp1	—	0
WDWB	Width dependence of dwb	—	0
WDWG	Width dependence of dwg	—	0
WEB	Coefficient for SCB	—	0
WEC	Coefficient for SCC	—	0
WEGIDL	Width dependence of egidl	—	0
WEGISL	Width dependence of egisl	—	0
WEIGBINV	Width dependence for eigbinv	—	0
WETA0	Width dependence of eta0	—	0
WETAB	Width dependence of etab	—	0
WEU	Width dependence of eu	—	0
WFPROUT	Width dependence of pdiblcb	—	0
WGAMMA1	Width dependence of gamma1	—	0
WGAMMA2	Width dependence of gamma2	—	0
WK1	Width dependence of k1	—	0
WK2	Width dependence of k2	—	0
WK2WE	Width dependence of k2we	—	0
WK3	Width dependence of k3	—	0
WK3B	Width dependence of k3b	—	0
WKETA	Width dependence of keta	—	0
WKT1	Width dependence of kt1	—	0
WKT1L	Width dependence of kt1l	—	0
WKT2	Width dependence of kt2	—	0
WKU0	Width dependence of ku0	—	0
WKUOWE	Width dependence of ku0we	—	0
WKVTH0	Width dependence of kvth0	—	0
WKVTHOWE	Width dependence of kvth0we	—	0
WL	Width reduction parameter	—	0
WLAMBDA	Width dependence of lambda	—	0
WLC	Width reduction parameter for CV	—	0
WLN	Width reduction parameter	—	1
WLOD	Width parameter for stress effect	—	0

Table 2.67: BSIM4 Device Model Parameters

Parameter	Description	Units	Default
WLODKU0	Width parameter for u0 LOD effect	—	0
WLODVTH	Width parameter for vth LOD effect	—	0
WLP	Width dependence of $I_p$	—	0
WLPE0	Width dependence of $I_{pe0}$	—	0
WLPEB	Width dependence of $I_{peb}$	—	0
WMAX	Maximum width for the model	—	1
WMIN	Minimum width for the model	—	0
WMINV	Width dependence of $\min v$	—	0
WMINVCV	Width dependence of $\min v_{cv}$	—	0
WMOIN	Width dependence of $m_{oin}$	—	0
WNDEP	Width dependence of $n_{dep}$	—	0
WNFACTOR	Width dependence of $n_{factor}$	—	0
WNGATE	Width dependence of $n_{gate}$	—	0
WNIGBACC	Width dependence of $n_{igbacc}$	—	0
WNIGBINV	Width dependence of $n_{igbinv}$	—	0
WNIGC	Width dependence of $n_{igc}$	—	0
WNOFF	Width dependence of $n_{off}$	—	0
WNSD	Width dependence of $n_{sd}$	—	0
WNSUB	Width dependence of $n_{sub}$	—	0
WNTOX	Width dependence of $n_{tox}$	—	0
WPCLM	Width dependence of $p_{clm}$	—	0
WPDIBLC1	Width dependence of $p_{diblc1}$	—	0
WPDIBLC2	Width dependence of $p_{diblc2}$	—	0
WPDIBLCB	Width dependence of $p_{diblcb}$	—	0
WPDITS	Width dependence of $p_{dits}$	—	0
WPDITSD	Width dependence of $p_{ditsd}$	—	0
WPEMOD	Flag for WPE model (WPEMOD=1 to activate this model)	—	0
WPHIN	Width dependence of $\phi_{in}$	—	0
WPIGCD	Width dependence for $p_{igcd}$	—	0
WPOXEDGE	Width dependence for $p_{oxedge}$	—	0
WPRT	Width dependence of $p_{rt}$	—	0
WPRWB	Width dependence of $p_{rwb}$	—	0
WPRWG	Width dependence of $p_{rwg}$	—	0
WPSCBE1	Width dependence of $p_{scbe1}$	—	0

Table 2.67: BSIM4 Device Model Parameters

Parameter	Description	Units	Default
WPSCBE2	Width dependence of pscbe2	—	0
WPVAG	Width dependence of pvag	—	0
WRDSW	Width dependence of rds	—	0
WRDW	Width dependence of rdw	—	0
WRSW	Width dependence of rsw	—	0
WTVFBSDOFF	Width dependence of tvfbsdoff	—	0
WTVOFF	Width dependence of tvoff	—	0
WU0	Width dependence of u0	—	0
WUA	Width dependence of ua	—	0
WUA1	Width dependence of ua1	—	0
WUB	Width dependence of ub	—	0
WUB1	Width dependence of ub1	—	0
WUC	Width dependence of uc	—	0
WUC1	Width dependence of uc1	—	0
WUD	Width dependence of ud	—	0
WUD1	Width dependence of ud1	—	0
WUP	Width dependence of up	—	0
WUTE	Width dependence of ute	—	0
WVBM	Width dependence of vbm	—	0
WVBX	Width dependence of vb	—	0
WVFB	Width dependence of vfb	—	0
WVFBCV	Width dependence of vfbcv	—	0
WVFBSDOFF	Width dependence of vfbsdoff	—	0
WVOFF	Width dependence of voff	—	0
WVOFFCV	Width dependence of voffcv	—	0
WVSAT	Width dependence of vsat	—	0
WVTHO		—	0
WVTL	Width dependence of vtl	—	0
WW	Width reduction parameter	—	0
WU0	Width dependence of w0	—	0
WWC	Width reduction parameter for CV	—	0
WWL	Width reduction parameter	—	0
WWLC	Width reduction parameter for CV	—	0
WWN	Width reduction parameter	—	1
WWR	Width dependence of wr	—	0

Table 2.67: BSIM4 Device Model Parameters

Parameter	Description	Units	Default
WXJ	Width dependence of $x_j$	—	0
WXN	Width dependence of $x_n$	—	0
WXRCRG1	Width dependence of $xrcrg1$	—	0
WXRCRG2	Width dependence of $xrcrg2$	—	0
WXT	Width dependence of $x_t$	—	0
XGL	Variation in $L_{drawn}$	—	0
XGW	Distance from gate contact center to device edge	—	0
XJBVD	Fitting parameter for drain diode breakdown current	—	1
XJBVS	Fitting parameter for source diode breakdown current	—	1
XL	L offset for channel length due to mask/etch effect	—	0
XRCRG1	First fitting parameter the bias-dependent $R_g$	—	12
XRCRG2	Second fitting parameter the bias-dependent $R_g$	—	1
XTID	Drain junction current temperature exponent	—	3
XTIS	Source junction current temperature exponent	—	3
XTSD	Power dependence of $J_{TSD}$ on temperature	—	0.02
XTSS	Power dependence of $J_{TSS}$ on temperature	—	0.02
XTSSWD	Power dependence of $J_{TSSWD}$ on temperature	—	0.02
XTSSWGD	Power dependence of $J_{TSSWGD}$ on temperature	—	0.02
XTSSWGS	Power dependence of $J_{TSSWGS}$ on temperature	—	0.02
XTSSWS	Power dependence of $J_{TSSWS}$ on temperature	—	0.02
XW	W offset for channel width due to mask/etch effect	—	0
<b>Basic Parameters</b>			
A0	Non-uniform depletion width effect coefficient.	—	1
A1	Non-saturation effect coefficient	$V^{-1}$	0
A2	Non-saturation effect coefficient	—	1
ADOS	Charge centroid parameter	—	1
AGS	Gate bias coefficient of $A_{bulk}$ .	$V^{-1}$	0
B0	$A_{bulk}$ narrow width parameter	m	0
B1	$A_{bulk}$ narrow width parameter	m	0
BDOS	Charge centroid parameter	—	1
BGOSUB	Band-gap of substrate at $T=0K$	eV	1.16

Table 2.67: BSIM4 Device Model Parameters

Parameter	Description	Units	Default
CDSC	Drain/Source and channel coupling capacitance	F/m <sup>2</sup>	0.00024
CDSCB	Body-bias dependence of cdsc	F/(Vm <sup>2</sup> )	0
CDSCD	Drain-bias dependence of cdsc	F/(Vm <sup>2</sup> )	0
CIT	Interface state capacitance	F/m <sup>2</sup>	0
DELTA	Effective Vds parameter	V	0.01
DROUT	DIBL coefficient of output resistance	–	0.56
DSUB	DIBL coefficient in the subthreshold region	–	0
DVTO	Short channel effect coeff. 0	–	2.2
DVTOW	Narrow Width coeff. 0	–	0
DVT1	Short channel effect coeff. 1	–	0.53
DVT1W	Narrow Width effect coeff. 1	m <sup>-1</sup>	5.3e+06
DVT2	Short channel effect coeff. 2	V <sup>-1</sup>	-0.032
DVT2W	Narrow Width effect coeff. 2	V <sup>-1</sup>	-0.032
DVTP0	First parameter for Vth shift due to pocket	m	0
DVTP1	Second parameter for Vth shift due to pocket	V <sup>-1</sup>	0
DWB	Width reduction parameter	m/V <sup>1/2</sup>	0
DWG	Width reduction parameter	m/V	0
EASUB	Electron affinity of substrate	V	4.05
EPSRSUB	Dielectric constant of substrate relative to vacuum	–	11.7
ETA0	Subthreshold region DIBL coefficient	–	0.08
ETAB	Subthreshold region DIBL coefficient	V <sup>-1</sup>	-0.07
EU	Mobility exponent	–	0
FPROUT	Rout degradation coefficient for pocket devices	V/m <sup>1/2</sup>	0
K1	Bulk effect coefficient 1	V <sup>-1/2</sup>	0
K2	Bulk effect coefficient 2	–	0
K3	Narrow width effect coefficient	–	80
KETA	Body-bias coefficient of non-uniform depletion width effect.	V <sup>-1</sup>	-0.047
LAMBDA	Velocity overshoot parameter	–	0
LC	back scattering parameter	m	5e-09
LINT	Length reduction parameter	m	0
LP	Channel length exponential factor of mobility	m	1e-08
LPE0	Equivalent length of pocket region at zero bias	m	1.74e-07
LPEB	Equivalent length of pocket region accounting for body bias	m	0

Table 2.67: BSIM4 Device Model Parameters

Parameter	Description	Units	Default
MINV	Fitting parameter for moderate inversion in $V_{gsteff}$	—	0
NFACTOR	Subthreshold swing Coefficient	—	1
NIOSUB	Intrinsic carrier concentration of substrate at 300.15K	$cm^{-3}$	1.45e+10
PCLM	Channel length modulation Coefficient	—	1.3
PDIBLC1	Drain-induced barrier lowering coefficient	—	0.39
PDIBLC2	Drain-induced barrier lowering coefficient	—	0.0086
PDIBLCB	Body-effect on drain-induced barrier lowering	$V^{-1}$	0
PDITS	Coefficient for drain-induced $V_{th}$ shifts	$V^{-1}$	0
PDITSD	$V_{ds}$ dependence of drain-induced $V_{th}$ shifts	$V^{-1}$	0
PDITSL	Length dependence of drain-induced $V_{th}$ shifts	$m^{-1}$	0
PHIN	Adjusting parameter for surface potential due to non-uniform vertical doping	V	0
PSCBE1	Substrate current body-effect coefficient	V/m	4.24e+08
PSCBE2	Substrate current body-effect coefficient	m/V	1e-05
TBGASUB	First parameter of band-gap change due to temperature	eV/K	0.000702
TBGBSUB	Second parameter of band-gap change due to temperature	K	1108
U0	Low-field mobility at $T_{nom}$	$m^2/(Vs)$	0
UA	Linear gate dependence of mobility	m/V	0
UB	Quadratic gate dependence of mobility	$m^2/V^2$	1e-19
UC	Body-bias dependence of mobility	$V^{-1}$	0
UD	Coulomb scattering factor of mobility	$m^{-2}$	0
UP	Channel length linear factor of mobility	$m^{-2}$	0
VBM	Maximum body voltage	V	-3
VDDEOT	Voltage for extraction of equivalent gate oxide thickness	V	1.5
VFB	Flat Band Voltage	V	-1
VOFF	Threshold voltage offset	V	-0.08
VOFFL	Length dependence parameter for $V_{th}$ offset	V	0
VSAT	Saturation velocity at $t_{nom}$	m/s	80000
VTH0		V	0
VTL	thermal velocity	m/s	200000
W0	Narrow width effect parameter	m	2.5e-06
WINT	Width reduction parameter	m	0

Table 2.67: BSIM4 Device Model Parameters

Parameter	Description	Units	Default
XN	back scattering parameter	–	3
<b>Capacitance Parameters</b>			
ACDE	Exponential coefficient for finite charge thickness	m/V	1
CF	Fringe capacitance parameter	F/m	0
CGB0	Gate-bulk overlap capacitance per length	–	0
CGDL	New C-V model parameter	F/m	0
CGD0	Gate-drain overlap capacitance per width	F/m	0
CGSL	New C-V model parameter	F/m	0
CGS0	Gate-source overlap capacitance per width	F/m	0
CKAPPAD	D/G overlap C-V parameter	V	0.6
CKAPPAS	S/G overlap C-V parameter	V	0.6
CLC	Vdsat parameter for C-V model	m	1e-07
CLE	Vdsat parameter for C-V model	–	0.6
DLC	Delta L for C-V model	m	0
DWC	Delta W for C-V model	m	0
MINVCV	Fitting parameter for moderate inversion in Vgsteffcv	–	0
MOIN	Coefficient for gate-bias dependent surface potential	–	15
NOFF	C-V turn-on/off parameter	–	1
VFBCV	Flat Band Voltage parameter for capmod=0 only	V	-1
VOFFCV	C-V lateral-shift parameter	V	0
VOFFCVL	Length dependence parameter for Vth offset in CV	–	0
XPART	Channel charge partitioning	F/m	0
<b>Control Parameters</b>			
ACNQSMOD	AC NQS model selector	–	0
BINUNIT	Bin unit selector	–	1
CAPMOD	Capacitance model selector	–	2
CVCHARGEMOD	Capacitance charge model selector	–	0
DIOMOD	Diode IV model selector	–	1
FNOIMOD	Flicker noise model selector	–	1
GEOMOD	Geometry dependent parasitics model selector	–	0
IGBMOD	Gate-to-body Ig model selector	–	0
IGCMOD	Gate-to-channel Ig model selector	–	0
MOBMOD	Mobility model selector	–	0

Table 2.67: BSIM4 Device Model Parameters

Parameter	Description	Units	Default
MTRLMOD	parameter for nonm-silicon substrate or metal gate selector	–	0
PARAMCHK	Model parameter checking selector	–	1
PERMOD	Pd and Ps model selector	–	1
RBODYMOD	Distributed body R model selector	–	0
RDSMOD	Bias-dependent S/D resistance model selector	–	0
RGATEMOD	Gate R model selector	–	0
TEMPMOD	Temperature model selector	–	0
TNOIMOD	Thermal noise model selector	–	0
TRNQSMOD	Transient NQS model selector	–	0
VERSION	parameter for model version	–	'4.6.1'
<b><i>Flicker Parameters</i></b>			
NOIA	Flicker Noise parameter a	–	0
NOIB	Flicker Noise parameter b	–	0
NOIC	Flicker Noise parameter c	–	0
<b><i>Process Parameters</i></b>			
DTOX	Defined as (toxe - toxp)	m	0
EOT	Equivalent gate oxide thickness in meters	m	1.5e-09
EPSROX	Dielectric constant of the gate oxide relative to vacuum	–	3.9
GAMMA1	Vth body coefficient	$V^{1/2}$	0
GAMMA2	Vth body coefficient	$V^{1/2}$	0
NDEP	Channel doping concentration at the depletion edge	$cm^{-3}$	1.7e+17
NGATE	Poly-gate doping concentration	$cm^{-3}$	0
NSD	S/D doping concentration	$cm^{-3}$	1e+20
NSUB	Substrate doping concentration	$cm^{-3}$	6e+16
RSH	Source-drain sheet resistance	$\Omega/\square$	0
RSHG	Gate sheet resistance	$\Omega/\square$	0.1
TOXE	Electrical gate oxide thickness in meters	m	3e-09
TOXM	Gate oxide thickness at which parameters are extracted	m	3e-09
TOXP	Physical gate oxide thickness in meters	m	3e-09
VBX	Vth transition body Voltage	V	0
XJ	Junction depth in meters	m	1.5e-07
XT	Doping depth	m	1.55e-07

Table 2.67: BSIM4 Device Model Parameters

Parameter	Description	Units	Default
<b><i>Tunnelling Parameters</i></b>			
AIGBACC	Parameter for Igb	$(Fs^2/g)^{1/2}/n$	0.0136
AIGBINV	Parameter for Igb	$(Fs^2/g)^{1/2}/n$	0.0111
AIGC	Parameter for Igc	$(Fs^2/g)^{1/2}/n$	0.0136
AIGD	Parameter for Igd	$(Fs^2/g)^{1/2}/n$	0.0136
AIGS	Parameter for Igs	$(Fs^2/g)^{1/2}/n$	0.0136
BIGBACC	Parameter for Igb	$(Fs^2/g)^{1/2}/n$	0.00171
BIGBINV	Parameter for Igb	$(Fs^2/g)^{1/2}/n$	0.000949
BIGC	Parameter for Igc	$(Fs^2/g)^{1/2}/n$	0.00171
BIGD	Parameter for Igd	$(Fs^2/g)^{1/2}/n$	0.00171
BIGS	Parameter for Igs	$(Fs^2/g)^{1/2}/n$	0.00171
CIGBACC	Parameter for Igb	$V^{-1}$	0.075
CIGBINV	Parameter for Igb	$V^{-1}$	0.006
CIGC	Parameter for Igc	$V^{-1}$	0.075
CIGD	Parameter for Igd	$V^{-1}$	0.075
CIGS	Parameter for Igs	$V^{-1}$	0.075
DLICGD	Delta L for Ig model drain side	m	0
EIGBINV	Parameter for the Si bandgap for Igbinv	V	1.1
NIGBACC	Parameter for Igbacc slope	—	1
NIGBINV	Parameter for Igbinv slope	—	3
NIGC	Parameter for Igc slope	—	1
NTOX	Exponent for Tox ratio	—	1
PIGCD	Parameter for Igc partition	—	1
POXEDGE	Factor for the gate edge Tox	—	1
TOXREF	Target tox value	m	3e-09
VFBSDOFF	S/D flatband voltage offset	V	0
<b><i>Asymmetric and Bias-Dependent <math>R_{ds}</math> Parameters</i></b>			
PRWB	Body-effect on parasitic resistance	$V^{-1}$	0
PRWG	Gate-bias effect on parasitic resistance	$V^{-1}$	1
RDSW	Source-drain resistance per width	$\Omega \mu m$	200
RDSWMIN	Source-drain resistance per width at high Vg	$\Omega \mu m$	0
RDW	Drain resistance per width	$\Omega \mu m$	100
RDWMIN	Drain resistance per width at high Vg	$\Omega \mu m$	0
RSW	Source resistance per width	$\Omega \mu m$	100
RSWMIN	Source resistance per width at high Vg	$\Omega \mu m$	0

Table 2.67: BSIM4 Device Model Parameters

Parameter	Description	Units	Default
WR	Width dependence of $r_{ds}$	—	1
<b><i>Impact Ionization Current Parameters</i></b>			
ALPHA0	substrate current model parameter	m/V	0
ALPHA1	substrate current model parameter	$V^{-1}$	0
BETA0	substrate current model parameter	$V^{-1}$	0
<b><i>Gate-induced Drain Leakage Model Parameters</i></b>			
AGIDL	Pre-exponential constant for GIDL	$\Omega^{-1}$	0
AGISL	Pre-exponential constant for GISL	$\Omega^{-1}$	0
BGIDL	Exponential constant for GIDL	V/m	2.3e+09
BGISL	Exponential constant for GISL	V/m	2.3e-09
CGIDL	Parameter for body-bias dependence of GIDL	$V^3$	0.5
CGISL	Parameter for body-bias dependence of GISL	$V^3$	0.5
EGIDL	Fitting parameter for Bandbending	V	0.8
EGISL	Fitting parameter for Bandbending	V	0.8

## Level 18 MOSFET Tables (VDMOS)

The vertical double-diffused power MOSFET model is based on the uniform charge control model (UCCM) developed at Rensselaer Polytechnic Institute [10]. The VDMOS current-voltage characteristics are described by a single, continuous analytical expression for all regimes of operation. The physics-based model includes effects such as velocity saturation in the channel, drain induced barrier lowering, finite output conductance in saturation, the quasi-saturation effect through a bias dependent drain parasitic resistance, effects of bulk charge, and bias dependent low-field mobility. An important feature of the implementation is the utilization of a single continuous expression for the drain current, which is valid below and above threshold, effectively removing discontinuities and improving convergence properties.

The following tables give parameters for the level 18 MOSFET.

Table 2.68: Power MOSFET Device Instance Parameters

Parameter	Description	Units	Default
AD	Drain diffusion area	m <sup>2</sup>	0
AS	Source diffusion area	m <sup>2</sup>	0
L	Channel length	m	0
M	Multiplier for M devices connected in parallel	–	1
NRD	Multiplier for RSH to yield parasitic resistance of drain	□	1
NRS	Multiplier for RSH to yield parasitic resistance of source	□	1
PD	Drain diffusion perimeter	m	0
PS	Source diffusion perimeter	m	0
TEMP	Device temperature	°C	Ambient Temperature
W	Channel width	m	0

Table 2.69: Power MOSFET Device Model Parameters

Parameter	Description	Units	Default
AI		–	2e+09
ALPHA	Parameter accounting for the threshold dependence on the channel potential	–	0
ARTD		–	0
BI		–	8e+08
BRTD		–	0.035
CBD	Zero-bias bulk-drain p-n capacitance	F	0
CBS	Zero-bias bulk-source p-n capacitance	F	0

Table 2.69: Power MOSFET Device Model Parameters

Parameter	Description	Units	Default
CGB0	Gate-bulk overlap capacitance/channel length	F/m	0
CGD0	Gate-drain overlap capacitance/channel width	F/m	0
CGS0	Gate-source overlap capacitance/channel width	F/m	0
CJ	Bulk p-n zero-bias bottom capacitance/area	F/m <sup>2</sup>	0
CJSW	Bulk p-n zero-bias sidewall capacitance/area	F/m <sup>2</sup>	0
CRTD		–	0.1472
CV	Charge model storage selector	–	1
CVE	Meyer-like capacitor model selector	–	1
D1AF	Drain-source diode flicker noise exponent	–	1
D1BV	Drain-source diode reverse breakdown voltage	V	1e+99
D1CJ0	Drain-source diode junction capacitance	F	0
D1EG	Drain-source diode activation energy	eV	1.11
D1FC	Drain-source diode forward bias depletion capacitance	–	0.5
D1IBV	Drain-source diode current at breakdown voltage	A	0.001
D1IKF	Drain-source diode high injection knee current	A	0
D1IS	Drain-Source diode saturation current	A	1e-14
D1ISR	Drain-source diode recombination saturation current	A	0
D1KF	Drain-source diode flicker noise coefficient	–	0
D1M	Drain-source diode grading coefficient	–	0.5
D1N	Drain-source diode emission coefficient	–	1
D1NR	Drain-source diode recombination emission coefficient	–	2
D1RS	Drain-source diode ohmic resistance	Ω	0
D1TNOM	Drain-source diode nominal temperature	°C	300.15
D1TT	Drain-source diode transit time	s	0
D1VJ	Drain-source diode junction potential	V	1
D1XTI	Drain-source diode sat. current temperature exponent	–	3
DELMAX		–	0.9
DELTA	Transition width parameter	–	5
DRIFTPARAMA	Drift region resistance intercept parameter	Ω	0.08
DRIFTPARAMB	Drift region resistance slope parameter	Ω V <sup>-1</sup>	0.013
DRTD		–	0.0052
ETA	Subthreshold ideality factor	–	1.32

Table 2.69: Power MOSFET Device Model Parameters

Parameter	Description	Units	Default
FC	Coefficient for forward-bias depletion capacitance formula	—	0.5
FPE	Charge partitioning scheme selector	—	1
GAMMALO	Body effect constant in front of linear term	—	0
GAMMASO	Body effect constant in front of square root term	$V^{-1/2}$	0.5
IS	Bulk p-n saturation current	A	1e-14
ISUBMOD		—	0
JS	Bulk p-n saturation current density	A/m <sup>2</sup>	0
K		—	0
KVS		—	0
KVT		—	0
LO	Gate length of nominal device	m	0
LAMBDA	Output conductance parameter	$V^{-1}$	0.048
LD	Lateral diffusion length	m	0
LGAMMAL	Sensitivity of gL on device length	—	0
LGAMMAS	Sensitivity of gS on device length	$V^{-1/2}$	0
LS		—	3.5e-08
M	Knee shape parameter	—	4
MC		—	3
MCV	Transition width parameter used by the charge partitioning scheme	—	10
MD		—	2
MDTEMP		—	0
MJ	Bulk p-n bottom grading coefficient	—	0.5
MJSW	Bulk p-n sidewall grading coefficient	—	0.5
MTH		—	0
N2		—	1
NRTD		—	0.115
NSS	Surface state density	cm <sup>-2</sup>	0
NSUB	Substrate doping density	cm <sup>-3</sup>	0
PB	Bulk p-n bottom potential	V	0.8
PHI	Surface potential	V	0.6
RD	Drain ohmic resistance	$\Omega$	0
RDSSHUNT	Drain-source shunt resistance	$\Omega$	0
RG	Gate ohmic resistance	$\Omega$	0

Table 2.69: Power MOSFET Device Model Parameters

Parameter	Description	Units	Default
RS	Source ohmic resistance	$\Omega$	0
RSH	Drain, source diffusion sheet resistance	$\Omega$	0
RSUB		–	0
SIGMA0	DIBL parameter	–	0.048
TEMPMODEL	Specifies the type of parameter interpolation over temperature	–	'NONE'
THETA	Mobility degradation parameter	m/V	0
TNOM	Nominal device temperature	$^{\circ}\text{C}$	Ambient Temperature
TOX	Gate oxide thickness	m	1e-07
TPG	Gate material type (-1 = same as substrate, 0 = aluminum, 1 = opposite of substrate)	–	1
TS		–	0
TVS		–	0
U0	Surface mobility	$1/(\text{Vcm}^2\text{s})$	280
U0	Surface mobility	$1/(\text{Vcm}^2\text{s})$	280
VFB	Flat band voltage	V	0
VMAX	Maximum drift velocity for carriers	m/s	40000
VP		–	0
VSIGMA	DIBL parameter	V	0.2
VSIGMAT	DIBL parameter	V	1.7
VTO	Zero-bias threshold voltage	V	0
W0	Gate width of nominal device	m	0
WGAMMAL	Sensitivity of gL on device width	–	0
WGAMMAS	Sensitivity of gS on device width	$\text{V}^{-1/2}$	0
XJ	Metallurgical junction depth	m	0
XQC	Charge partitioning factor	–	0.6

## Level 103 MOSFET Tables (PSP version 103.1)

**Xyce** includes the PSP MOSFET model, version 103.1 [24]. Full documentation for the PSP model is available on its web site, [http://pspmodel.asu.edu/psp\\_documentation.htm](http://pspmodel.asu.edu/psp_documentation.htm). Instance and model parameters for the PSP model are given in tables 2.70 and 2.71.

Table 2.70: PSP103VA MOSFET Device Instance Parameters

Parameter	Description	Units	Default
ABDRAIN	Bottom area of drain junction	–	1e-12
ABSOURCE	Bottom area of source junction	–	1e-12
AD	Bottom area of drain junction	–	1e-12
AS	Bottom area of source junction	–	1e-12
DELVTO	Threshold voltage shift parameter	V	0
FACTU0	Zero-field mobility pre-factor	–	1
JW	Gate-edge length of source/drain junction	m	1e-06
L	Design length	m	1e-05
LGDRAIN	Gate-edge length of drain junction	m	1e-06
LGSOURCE	Gate-edge length of source junction	m	1e-06
LSDRAIN	STI-edge length of drain junction	m	1e-06
LSSOURCE	STI-edge length of source junction	m	1e-06
MULT	Number of devices in parallel	–	1
NF	Number of fingers	–	1
NGCON	Number of gate contacts	–	1
NRD	Number of squares of drain diffusion	–	0
NRS	Number of squares of source diffusion	–	0
PD	Perimeter of drain junction	m	1e-06
PS	Perimeter of source junction	m	1e-06
SA	Distance between OD-edge and poly from one side	m	0
SB	Distance between OD-edge and poly from other side	m	0
SC	Distance between OD-edge and nearest well edge	m	0
SCA	Integral of the first distribution function for scattered well dopants	–	0
SCB	Integral of the second distribution function for scattered well dopants	–	0
SCC	Integral of the third distribution function for scattered well dopants	–	0
SD	Distance between neighbouring fingers	m	0

Table 2.70: PSP103VA MOSFET Device Instance Parameters

Parameter	Description	Units	Default
W	Design width	m	1e-05
XGW	Distance from the gate contact to the channel edge	m	1e-07

Table 2.71: PSP103VA MOSFET Device Model Parameters

Parameter	Description	Units	Default
A1	Impact-ionization pre-factor	—	1
A1L	Length dependence of A1	—	0
A10	Geometry independent impact-ionization pre-factor	—	1
A1W	Width dependence of A1	—	0
A2	Impact-ionization exponent at TR	V	10
A20	Impact-ionization exponent at TR	V	10
A3	Saturation-voltage dependence of impact-ionization	—	1
A3L	Length dependence of A3	—	0
A30	Geometry independent saturation-voltage dependence of II	—	1
A3W	Width dependence of A3	—	0
A4	Back-bias dependence of impact-ionization	$V^{-1/2}$	0
A4L	Length dependence of A4	—	0
A40	Geometry independent back-bias dependence of II	$V^{-1/2}$	0
A4W	Width dependence of A4	—	0
AGIDL	GIDL pre-factor	$A/V^3$	0
AGIDLD	GIDL pre-factor for drain side	$A/V^3$	0
AGIDLDW	Width dependence of GIDL pre-factor for drain side	$A/V^3$	0
AGIDLW	Width dependence of GIDL pre-factor	$A/V^3$	0
ALP	CLM pre-factor	—	0.01
ALP1	CLM enhancement factor above threshold	V	0
ALP1L1	Length dependence of CLM enhancement factor above threshold	V	0
ALP1L2	Second order length dependence of ALP1	—	0
ALP1LEXP	Exponent for length dependence of ALP1	—	0.5
ALP1W	Width dependence of ALP1	—	0

Table 2.71: PSP103VA MOSFET Device Model Parameters

Parameter	Description	Units	Default
ALP2	CLM enhancement factor below threshold	$V^{-1}$	0
ALP2L1	Length dependence of CLM enhancement factor below threshold	$V^{-1}$	0
ALP2L2	Second order length dependence of ALP2	–	0
ALP2LEXP	Exponent for length dependence of ALP2	–	0.5
ALP2W	Width dependence of ALP2	–	0
ALPL	Length dependence of ALP	–	0.0005
ALPLEXP	Exponent for length dependence of ALP	–	1
ALPNOI	Exponent for length offset for flicker noise	–	2
ALPW	Width dependence of ALP	–	0
AX	Linear/saturation transition factor	–	3
AXL	Length dependence of AX	–	0.4
AX0	Geometry independent linear/saturation transition factor	–	18
BETN	Channel aspect ratio times zero-field mobility	$m^2/(Vs)$	0.07
BETW1	First higher-order width scaling coefficient of BETN	–	0
BETW2	Second higher-order width scaling coefficient of BETN	–	0
BGIDL	GIDL probability factor at TR	V	41
BGIDLD	GIDL probability factor at TR for drain side	V	41
BGIDLDO	GIDL probability factor at TR for drain side	V	41
BGIDL0	GIDL probability factor at TR	V	41
CBBTBOT	Band-to-band tunneling prefactor of bottom component for source-bulk junction	–	1e-12
CBBTBOTD	Band-to-band tunneling prefactor of bottom component for drain-bulk junction	–	1e-12
CBBTGAT	Band-to-band tunneling prefactor of gate-edge component for source-bulk junction	–	1e-18
CBBTGATD	Band-to-band tunneling prefactor of gate-edge component for drain-bulk junction	–	1e-18
CBBTSTI	Band-to-band tunneling prefactor of STI-edge component for source-bulk junction	–	1e-18
CBBTSTID	Band-to-band tunneling prefactor of STI-edge component for drain-bulk junction	–	1e-18
CF	DIBL-parameter	–	0
CFB	Back bias dependence of CF	$V^{-1}$	0
CFB0	Back-bias dependence of CF	$V^{-1}$	0

Table 2.71: PSP103VA MOSFET Device Model Parameters

Parameter	Description	Units	Default
CFL	Length dependence of DIBL-parameter	–	0
CFLEXP	Exponent for length dependence of CF	–	2
CFR	Outer fringe capacitance	F	0
CFRD	Outer fringe capacitance for drain side	F	0
CFRDW	Outer fringe capacitance for 1 um wide channel for drain side	F	0
CFRW	Outer fringe capacitance for 1 um wide channel	F	0
CFW	Width dependence of CF	–	0
CGBOV	Oxide capacitance for gate-bulk overlap	F	0
CGBOVL	Oxide capacitance for gate-bulk overlap for 1 um long channel	F	0
CGIDL	Back-bias dependence of GIDL	–	0
CGIDLD	Back-bias dependence of GIDL for drain side	–	0
CGIDLDO	Back-bias dependence of GIDL for drain side	–	0
CGIDL0	Back-bias dependence of GIDL	–	0
CGOV	Oxide capacitance for gate-drain/source overlap	F	1e-15
CGOVD	Oxide capacitance for gate-drain overlap	F	1e-15
CHIB	Tunneling barrier height	V	3.1
CHIB0	Tunneling barrier height	V	3.1
CJORBOT	Zero-bias capacitance per unit-of-area of bottom component for source-bulk junction	–	0.001
CJORBOTD	Zero-bias capacitance per unit-of-area of bottom component for drain-bulk junction	–	0.001
CJORGAT	Zero-bias capacitance per unit-of-length of gate-edge component for source-bulk junction	–	1e-09
CJORGATD	Zero-bias capacitance per unit-of-length of gate-edge component for drain-bulk junction	–	1e-09
CJORSTI	Zero-bias capacitance per unit-of-length of STI-edge component for source-bulk junction	–	1e-09
CJORSTID	Zero-bias capacitance per unit-of-length of STI-edge component for drain-bulk junction	–	1e-09
COX	Oxide capacitance for intrinsic channel	F	1e-14
CS	Coulomb scattering parameter at TR	–	0
CSL	Length dependence of CS	–	0
CSLEXP	Exponent for length dependence of CS	–	1
CSLW	Area dependence of CS	–	0
CS0	Geometry independent coulomb scattering parameter at TR	–	0

Table 2.71: PSP103VA MOSFET Device Model Parameters

Parameter	Description	Units	Default
CSRHBOT	Shockley-Read-Hall prefactor of bottom component for source-bulk junction	–	100
CSRHBOTD	Shockley-Read-Hall prefactor of bottom component for drain-bulk junction	–	100
CSRHGAT	Shockley-Read-Hall prefactor of gate-edge component for source-bulk junction	–	0.0001
CSRHGATD	Shockley-Read-Hall prefactor of gate-edge component for drain-bulk junction	–	0.0001
CSRHSTI	Shockley-Read-Hall prefactor of STI-edge component for source-bulk junction	–	0.0001
CSRHSTID	Shockley-Read-Hall prefactor of STI-edge component for drain-bulk junction	–	0.0001
CSW	Width dependence of CS	–	0
CT	Interface states factor	–	0
CTATBOT	Trap-assisted tunneling prefactor of bottom component for source-bulk junction	–	100
CTATBOTD	Trap-assisted tunneling prefactor of bottom component for drain-bulk junction	–	100
CTATGAT	Trap-assisted tunneling prefactor of gate-edge component for source-bulk junction	–	0.0001
CTATGATD	Trap-assisted tunneling prefactor of gate-edge component for drain-bulk junction	–	0.0001
CTATSTI	Trap-assisted tunneling prefactor of STI-edge component for source-bulk junction	–	0.0001
CTATSTID	Trap-assisted tunneling prefactor of STI-edge component for drain-bulk junction	–	0.0001
CTL	Length dependence of interface states factor	–	0
CTLEXP	Exponent for length dependence of interface states factor	–	1
CTLW	Area dependence of interface states factor	–	0
CTO	Geometry-independent interface states factor	–	0
CTW	Width dependence of interface states factor	–	0
DELVTAC	Offset parameter for PHIB in separate charge calculation	V	0
DELVTACL	Length dependence of DELVTAC	V	0
DELVTACLEXP	Exponent for length dependence of offset of DELVTAC	–	1
DELVTACLW	Area dependence of DELVTAC	V	0
DELVTACO	Geom. independent offset parameter for PHIB in separate charge calculation	V	0

Table 2.71: PSP103VA MOSFET Device Model Parameters

Parameter	Description	Units	Default
DELVTACW	Width dependence of DELVTAC	V	0
DLQ	Effective channel length reduction for CV	m	0
DLSIL	Silicide extension over the physical gate length	m	0
DNSUB	Effective doping bias-dependence parameter	$V^{-1}$	0
DNSUBO	Effective doping bias-dependence parameter	$V^{-1}$	0
DPHIB	Offset parameter for PHIB	V	0
DPHIBL	Length dependence offset of PHIB	V	0
DPHIBLEXP	Exponent for length dependence of offset of PHIB	–	1
DPHIBLW	Area dependence of offset of PHIB	V	0
DPHIBO	Geometry independent offset of PHIB	V	0
DPHIBW	Width dependence of offset of PHIB	V	0
DTA	Temperature offset w.r.t. ambient temperature	K	0
DVSBNUD	Vsb-range for NUD-effect	V	1
DVSBNUDO	Vsb range for NUD-effect	V	1
DWQ	Effective channel width reduction for CV	m	0
EF	Flicker noise frequency exponent	–	1
EFO	Flicker noise frequency exponent	–	1
EPSROX	Relative permittivity of gate dielectric	–	3.9
EPSROXO	Relative permittivity of gate dielectric	–	3.9
FACNEFFAC	Pre-factor for effective substrate doping in separate charge calculation	–	1
FACNEFFACL	Length dependence of FACNEFFAC	–	0
FACNEFFACLW	Area dependence of FACNEFFAC	–	0
FACNEFFACO	Geom. independent pre-factor for effective substrate doping in separate charge calculation	–	1
FACNEFFACW	Width dependence of FACNEFFAC	–	0
FBTTRBOT	Normalization field at the reference temperature for band-to-band tunneling of bottom component for source-bulk junction	–	1e+09
FBTTRBOTD	Normalization field at the reference temperature for band-to-band tunneling of bottom component for drain-bulk junction	–	1e+09
FBTTRGAT	Normalization field at the reference temperature for band-to-band tunneling of gate-edge component for source-bulk junction	–	1e+09

Table 2.71: PSP103VA MOSFET Device Model Parameters

Parameter	Description	Units	Default
FBBTRGATD	Normalization field at the reference temperature for band-to-band tunneling of gate-edge component for drain-bulk junction	–	1e+09
FBBTRSTI	Normalization field at the reference temperature for band-to-band tunneling of STI-edge component for source-bulk junction	–	1e+09
FBBTRSTID	Normalization field at the reference temperature for band-to-band tunneling of STI-edge component for drain-bulk junction	–	1e+09
FBET1	Relative mobility decrease due to first lateral profile	–	0
FBET1W	Width dependence of relative mobility decrease due to first lateral profile	–	0
FBET2	Relative mobility decrease due to second lateral profile	–	0
FETA	Effective field parameter	–	1
FETA0	Effective field parameter	–	1
FJUNQ	Fraction below which source-bulk junction capacitance components are considered negligible	–	0.03
FJUNQD	Fraction below which drain-bulk junction capacitance components are considered negligible	–	0.03
FNT	Thermal noise coefficient	–	1
FNT0	Thermal noise coefficient	–	1
FOL1	First length dependence coefficient for short channel body effect	–	0
FOL2	Second length dependence coefficient for short channel body effect	–	0
GC2	Gate current slope factor	–	0.375
GC20	Gate current slope factor	–	0.375
GC3	Gate current curvature factor	–	0.063
GC30	Gate current curvature factor	–	0.063
GCO	Gate tunneling energy adjustment	–	0
GCO0	Gate tunneling energy adjustment	–	0
GFACNUD	Bodyfactor change due to NUD-effect	–	1
GFACNUDL	Length dependence of GFACNUD	–	0
GFACNUDLEXP	Exponent for length dependence of GFACNUD	–	1
GFACNUDLW	Area dependence of GFACNUD	–	0

Table 2.71: PSP103VA MOSFET Device Model Parameters

Parameter	Description	Units	Default
GFACNUDO	Geom. independent bodyfactor change due to NUD-effect	–	1
GFACNUDW	Width dependence of GFACNUD	–	0
IDSATRBOT	Saturation current density at the reference temperature of bottom component for source-bulk junction	–	1e-12
IDSATRBOTD	Saturation current density at the reference temperature of bottom component for drain-bulk junction	–	1e-12
IDSATRGAT	Saturation current density at the reference temperature of gate-edge component for source-bulk junction	–	1e-18
IDSATRGATD	Saturation current density at the reference temperature of gate-edge component for drain-bulk junction	–	1e-18
IDSATRSTI	Saturation current density at the reference temperature of STI-edge component for source-bulk junction	–	1e-18
IDSATRSTID	Saturation current density at the reference temperature of STI-edge component for drain-bulk junction	–	1e-18
IGINV	Gate channel current pre-factor	A	0
IGINVLW	Gate channel current pre-factor for 1 $\mu\text{m}^2$ channel area	A	0
IGOV	Gate overlap current pre-factor	A	0
IGOVD	Gate overlap current pre-factor for drain side	A	0
IGOVDW	Gate overlap current pre-factor for 1 $\mu\text{m}$ wide channel for drain side	A	0
IGOVW	Gate overlap current pre-factor for 1 $\mu\text{m}$ wide channel	A	0
IMAX	Maximum current up to which forward current behaves exponentially	–	1000
KUO	Mobility degradation/enhancement coefficient	m	0
KUOWEL	Length dependent mobility degradation factor	–	0
KUOWELW	Area dependent mobility degradation factor	–	0
KUOWEO	Geometrical independent mobility degradation factor	–	0
KUOWEW	Width dependent mobility degradation factor	–	0
KVSAT	Saturation velocity degradation/enhancement coefficient	m	0
KVTHO	Threshold shift parameter	Vm	0

Table 2.71: PSP103VA MOSFET Device Model Parameters

Parameter	Description	Units	Default
KVTHOWEL	Length dependent threshold shift parameter	–	0
KVTHOWELW	Area dependent threshold shift parameter	–	0
KVTHOWEO	Geometrical independent threshold shift parameter	–	0
KVTHOWEW	Width dependent threshold shift parameter	–	0
LAP	Effective channel length reduction per side	m	0
LEVEL	Model level	–	103
LINTNOI	Length offset for flicker noise	m	0
LKUO	Length dependence of KUO	–	0
LKVTHO	Length dependence of KVTHO	–	0
LLODKUO	Length parameter for UO stress effect	–	0
LLODVTH	Length parameter for VTH-stress effect	–	0
LMAX	Dummy parameter to label binning set	m	1
LMIN	Dummy parameter to label binning set	m	0
LODETA0	eta0 shift modification factor for stress effect	–	1
LOV	Overlap length for gate/drain and gate/source overlap capacitance	m	0
LOVD	Overlap length for gate/drain overlap capacitance	m	0
LP1	Mobility-related characteristic length of first lateral profile	m	1e-08
LP1W	Width dependence of mobility-related characteristic length of first lateral profile	–	0
LP2	Mobility-related characteristic length of second lateral profile	m	1e-08
LPCK	Char. length of lateral doping profile	m	1e-08
LPCKW	Width dependence of char. length of lateral doping profile	–	0
LVARL	Length dependence of LVAR	–	0
LVARO	Geom. independent difference between actual and programmed gate length	m	0
LVARW	Width dependence of LVAR	–	0
MEFFTATBOT	Effective mass (in units of m0) for trap-assisted tunneling of bottom component for source-bulk junction	–	0.25
MEFFTATBOTD	Effective mass (in units of m0) for trap-assisted tunneling of bottom component for drain-bulk junction	–	0.25

Table 2.71: PSP103VA MOSFET Device Model Parameters

Parameter	Description	Units	Default
MEFFTATGAT	Effective mass (in units of $m_0$ ) for trap-assisted tunneling of gate-edge component for source-bulk junction	–	0.25
MEFFTATGATD	Effective mass (in units of $m_0$ ) for trap-assisted tunneling of gate-edge component for drain-bulk junction	–	0.25
MEFFTATSTI	Effective mass (in units of $m_0$ ) for trap-assisted tunneling of STI-edge component for source-bulk junction	–	0.25
MEFFTATSTID	Effective mass (in units of $m_0$ ) for trap-assisted tunneling of STI-edge component for drain-bulk junction	–	0.25
MUE	Mobility reduction coefficient at TR	m/V	0.5
MUEO	Geometry independent mobility reduction coefficient at TR	m/V	0.5
MUEW	Width dependence of mobility reduction coefficient at TR	–	0
NEFF	Effective substrate doping	$m^{-3}$	5e+23
NFA	First coefficient of flicker noise	–	8e+22
NFALW	First coefficient of flicker noise for 1 $\mu m^2$ channel area	–	8e+22
NFB	Second coefficient of flicker noise	–	3e+07
NFBLW	Second coefficient of flicker noise for 1 $\mu m^2$ channel area	–	3e+07
NFC	Third coefficient of flicker noise	$V^{-1}$	0
NFCLW	Third coefficient of flicker noise for 1 $\mu m^2$ channel area	$V^{-1}$	0
NOV	Effective doping of overlap region	$m^{-3}$	5e+25
NOVD	Effective doping of overlap region for drain side	$m^{-3}$	5e+25
NOVDO	Effective doping of overlap region for drain side	$m^{-3}$	5e+25
NOVO	Effective doping of overlap region	$m^{-3}$	5e+25
NP	Gate poly-silicon doping	$m^{-3}$	1e+26
NPCK	Pocket doping level	$m^{-3}$	1e+24
NPCKW	Width dependence of pocket doping NPCK due to segregation	–	0
NPL	Length dependence of gate poly-silicon doping	–	0
NPO	Geometry-independent gate poly-silicon doping	$m^{-3}$	1e+26
NSLP	Effective doping bias-dependence parameter	V	0.05
NSLPO	Effective doping bias-dependence parameter	V	0.05

Table 2.71: PSP103VA MOSFET Device Model Parameters

Parameter	Description	Units	Default
NSUBO	Geometry independent substrate doping	$\text{m}^{-3}$	3e+23
NSUBW	Width dependence of background doping NSUBO due to segregation	—	0
PBOT	Grading coefficient of bottom component for source-bulk junction	—	0.5
PBOTD	Grading coefficient of bottom component for drain-bulk junction	—	0.5
PBRBOT	Breakdown onset tuning parameter of bottom component for source-bulk junction	—	4
PBRBOTD	Breakdown onset tuning parameter of bottom component for drain-bulk junction	—	4
PBRGAT	Breakdown onset tuning parameter of gate-edge component for source-bulk junction	—	4
PBRGATD	Breakdown onset tuning parameter of gate-edge component for drain-bulk junction	—	4
PBRSTI	Breakdown onset tuning parameter of STI-edge component for source-bulk junction	—	4
PBRSTID	Breakdown onset tuning parameter of STI-edge component for drain-bulk junction	—	4
PGAT	Grading coefficient of gate-edge component for source-bulk junction	—	0.5
PGATD	Grading coefficient of gate-edge component for drain-bulk junction	—	0.5
PHIGBOT	Zero-temperature bandgap voltage of bottom component for source-bulk junction	—	1.16
PHIGBOTD	Zero-temperature bandgap voltage of bottom component for drain-bulk junction	—	1.16
PHIGGAT	Zero-temperature bandgap voltage of gate-edge component for source-bulk junction	—	1.16
PHIGGATD	Zero-temperature bandgap voltage of gate-edge component for drain-bulk junction	—	1.16
PHIGSTI	Zero-temperature bandgap voltage of STI-edge component for source-bulk junction	—	1.16
PHIGSTID	Zero-temperature bandgap voltage of STI-edge component for drain-bulk junction	—	1.16
PKUO	Cross-term dependence of KUO	—	0
PKVTHO	Cross-term dependence of KVTHO	—	0
PLA1	Coefficient for the length dependence of A1	—	0
PLA3	Coefficient for the length dependence of A3	—	0
PLA4	Coefficient for the length dependence of A4	—	0

Table 2.71: PSP103VA MOSFET Device Model Parameters

Parameter	Description	Units	Default
PLAGIDL	Coefficient for the length dependence of AGIDL	$A/V^3$	0
PLAGIDLD	Coefficient for the length dependence of AGIDL for drain side	$A/V^3$	0
PLALP	Coefficient for the length dependence of ALP	–	0
PLALP1	Coefficient for the length dependence of ALP1	V	0
PLALP2	Coefficient for the length dependence of ALP2	–	0
PLAX	Coefficient for the length dependence of AX	–	0
PLBETN	Coefficient for the length dependence of BETN	–	0
PLCF	Coefficient for the length dependence of CF	–	0
PLCFR	Coefficient for the length dependence of CFR	F	0
PLCFRD	Coefficient for the length dependence of CFR for drain side	F	0
PLCGBOV	Coefficient for the length dependence of CGBOV	F	0
PLCGOV	Coefficient for the length dependence of CGOV	F	0
PLCGOVD	Coefficient for the length dependence of CGOV for drain side	F	0
PLCOX	Coefficient for the length dependence of COX	F	0
PLCS	Coefficient for the length dependence of CS	–	0
PLCT	Coefficient for the length dependence of CT	–	0
PLDELVTAC	Coefficient for the length dependence of DELVTAC	V	0
PLDPHIB	Coefficient for the length dependence of DPHIB	V	0
PLFACNEFFAC	Coefficient for the length dependence of FACNEFFAC	–	0
PLGFACNUD	Coefficient for the length dependence of GFACNUD	–	0
PLIGINV	Coefficient for the length dependence of IGINV	A	0
PLIGOV	Coefficient for the length dependence of IGOV	A	0
PLIGOVD	Coefficient for the length dependence of IGOV for drain side	A	0
PLKUOWE	Coefficient for the length dependence part of KUOWE	–	0
PLKVTHOWE	Coefficient for the length dependence part of KVTHOWE	–	0
PLMUE	Coefficient for the length dependence of MUE	–	0
PLNEFF	Coefficient for the length dependence of NEFF	$m^{-3}$	0
PLNFA	Coefficient for the length dependence of NFA	–	0
PLNFB	Coefficient for the length dependence of NFB	–	0

Table 2.71: PSP103VA MOSFET Device Model Parameters

Parameter	Description	Units	Default
PLNFC	Coefficient for the length dependence of NFC	–	0
PLNOV	Coefficient for the length dependence of NOV	$m^{-3}$	0
PLNOVD	Coefficient for the length dependence of NOV for drain side	$m^{-3}$	0
PLNP	Coefficient for the length dependence of NP	$m^{-3}$	0
PLRS	Coefficient for the length dependence of RS	–	0
PLSTBET	Coefficient for the length dependence of STBET	–	0
PLSTTHESAT	Coefficient for the length dependence of STTHESAT	–	0
PLSTVFB	Coefficient for the length dependence of STVFB	V/K	0
PLTHESAT	Coefficient for the length dependence of THESAT	–	0
PLTHESATB	Coefficient for the length dependence of THESATB	–	0
PLTHESATG	Coefficient for the length dependence of THESATG	–	0
PLVFB	Coefficient for the length dependence of VFB	V	0
PLWA1	Coefficient for the length times width dependence of A1	–	0
PLWA3	Coefficient for the length times width dependence of A3	–	0
PLWA4	Coefficient for the length times width dependence of A4	–	0
PLWAGIDL	Coefficient for the length times width dependence of AGIDL	$A/V^3$	0
PLWAGIDLD	Coefficient for the length times width dependence of AGIDL for drain side	$A/V^3$	0
PLWALP	Coefficient for the length times width dependence of ALP	–	0
PLWALP1	Coefficient for the length times width dependence of ALP1	V	0
PLWALP2	Coefficient for the length times width dependence of ALP2	–	0
PLWAX	Coefficient for the length times width dependence of AX	–	0
PLWBETN	Coefficient for the length times width dependence of BETN	–	0
PLWCF	Coefficient for the length times width dependence of CF	–	0
PLWCFR	Coefficient for the length times width dependence of CFR	F	0

Table 2.71: PSP103VA MOSFET Device Model Parameters

Parameter	Description	Units	Default
PLWCFRD	Coefficient for the length times width dependence of CFR for drain side	F	0
PLWCGBOV	Coefficient for the length times width dependence of CGBOV	F	0
PLWCGOV	Coefficient for the length times width dependence of CGOV	F	0
PLWCGOVD	Coefficient for the length times width dependence of CGOV for drain side	F	0
PLWCOX	Coefficient for the length times width dependence of COX	F	0
PLWCS	Coefficient for the length times width dependence of CS	–	0
PLWCT	Coefficient for the length times width dependence of CT	–	0
PLWDELVTAC	Coefficient for the length times width dependence of DELVTAC	V	0
PLWDPHIB	Coefficient for the length times width dependence of DPHIB	V	0
PLWFACNEFFAC	Coefficient for the length times width dependence of FACNEFFAC	–	0
PLWGFACNUD	Coefficient for the length times width dependence of GFACNUD	–	0
PLWIGINV	Coefficient for the length times width dependence of IGINV	A	0
PLWIGOV	Coefficient for the length times width dependence of IGOV	A	0
PLWIGOVD	Coefficient for the length times width dependence of IGOV for drain side	A	0
PLWKUOWE	Coefficient for the length times width dependence part of KUOWE	–	0
PLWKVTHOWE	Coefficient for the length times width dependence part of KVTHOWE	–	0
PLWMUE	Coefficient for the length times width dependence of MUE	–	0
PLWNEFF	Coefficient for the length times width dependence of NEFF	m <sup>-3</sup>	0
PLWNFA	Coefficient for the length times width dependence of NFA	–	0
PLWNFB	Coefficient for the length times width dependence of NFB	–	0
PLWNFC	Coefficient for the length times width dependence of NFC	–	0

Table 2.71: PSP103VA MOSFET Device Model Parameters

Parameter	Description	Units	Default
PLWNOV	Coefficient for the length times width dependence of NOV	$m^{-3}$	0
PLWNOVD	Coefficient for the length times width dependence of NOV for drain side	$m^{-3}$	0
PLWNP	Coefficient for the length times width dependence of NP	$m^{-3}$	0
PLWRS	Coefficient for the length times width dependence of RS	–	0
PLWSTBET	Coefficient for the length times width dependence of STBET	–	0
PLWSTTHESAT	Coefficient for the length times width dependence of STTHESAT	–	0
PLWSTVFB	Coefficient for the length times width dependence of STVFB	V/K	0
PLWTHESAT	Coefficient for the length times width dependence of THESAT	–	0
PLWTHESATB	Coefficient for the length times width dependence of THESATB	–	0
PLWTHESATG	Coefficient for the length times width dependence of THESATG	–	0
PLWVFB	Coefficient for the length times width dependence of VFB	V	0
PLWXCOR	Coefficient for the length times width dependence of XCOR	–	0
PLXCOR	Coefficient for the length dependence of XCOR	–	0
POA1	Coefficient for the geometry independent part of A1	–	1
POA2	Coefficient for the geometry independent part of A2	V	10
POA3	Coefficient for the geometry independent part of A3	–	1
POA4	Coefficient for the geometry independent part of A4	–	0
POAGIDL	Coefficient for the geometry independent part of AGIDL	$A/V^3$	0
POAGIDLD	Coefficient for the geometry independent part of AGIDL for drain side	$A/V^3$	0
POALP	Coefficient for the geometry independent part of ALP	–	0.01
POALP1	Coefficient for the geometry independent part of ALP1	V	0

Table 2.71: PSP103VA MOSFET Device Model Parameters

Parameter	Description	Units	Default
POALP2	Coefficient for the geometry independent part of ALP2	–	0
POAX	Coefficient for the geometry independent part of AX	–	3
POBETN	Coefficient for the geometry independent part of BETN	–	0.07
POBGIDL	Coefficient for the geometry independent part of BGIDL	V	41
POBGIDLD	Coefficient for the geometry independent part of BGIDL for drain side	V	41
POCF	Coefficient for the geometry independent part of CF	–	0
POCFB	Coefficient for the geometry independent part of CFB	–	0
POCFR	Coefficient for the geometry independent part of CFR	F	0
POCFRD	Coefficient for the geometry independent part of CFR for drain side	F	0
POCGBOV	Coefficient for the geometry independent part of CGBOV	F	0
POCGIDL	Coefficient for the geometry independent part of CGIDL	–	0
POCGIDLD	Coefficient for the geometry independent part of CGIDL for drain side	–	0
POCGOV	Coefficient for the geometry independent part of CGOV	F	1e-15
POCGOVD	Coefficient for the geometry independent part of CGOV for drain side	F	1e-15
POCHIB	Coefficient for the geometry independent part of CHIB	V	3.1
POCOX	Coefficient for the geometry independent part of COX	F	1e-14
POCS	Coefficient for the geometry independent part of CS	–	0
POCT	Coefficient for the geometry independent part of CT	–	0
PODELVTAC	Coefficient for the geometry independent part of DELVTAC	V	0
PODNSUB	Coefficient for the geometry independent part of DNSUB	–	0
PODPHIB	Coefficient for the geometry independent part of DPHIB	V	0

Table 2.71: PSP103VA MOSFET Device Model Parameters

Parameter	Description	Units	Default
PODVSBNUD	Coefficient for the geometry independent part of DVSBNUD	V	1
POEF	Coefficient for the flicker noise frequency exponent	–	1
POEPSROX	Coefficient for the geometry independent part of EPSOX	–	3.9
POFACNEFFAC	Coefficient for the geometry independent part of FACNEFFAC	–	1
POFETA	Coefficient for the geometry independent part of FETA	–	1
POFNT	Coefficient for the geometry independent part of FNT	–	1
POGC2	Coefficient for the geometry independent part of GC2	–	0.375
POGC3	Coefficient for the geometry independent part of GC3	–	0.063
POGCO	Coefficient for the geometry independent part of GCO	–	0
POGFACNUD	Coefficient for the geometry independent part of GFACNUD	–	1
POIGINV	Coefficient for the geometry independent part of IGINV	A	0
POIGOV	Coefficient for the geometry independent part of IGOV	A	0
POIGOVD	Coefficient for the geometry independent part of IGOV for drain side	A	0
POKUOWE	Coefficient for the geometry independent part of KUOWE	–	0
POKVTHOWE	Coefficient for the geometry independent part of KVTHOWE	–	0
POMUE	Coefficient for the geometry independent part of MUE	–	0.5
PONEFF	Coefficient for the geometry independent part of NEFF	m <sup>-3</sup>	5e+23
PONFA	Coefficient for the geometry independent part of NFA	–	8e+22
PONFB	Coefficient for the geometry independent part of NFB	–	3e+07
PONFC	Coefficient for the geometry independent part of NFC	–	0
PONOV	Coefficient for the geometry independent part of NOV	m <sup>-3</sup>	5e+25

Table 2.71: PSP103VA MOSFET Device Model Parameters

Parameter	Description	Units	Default
PONOV	Coefficient for the geometry independent part of NOV for drain side	$m^{-3}$	5e+25
PONP	Coefficient for the geometry independent part of NP	$m^{-3}$	1e+26
PONSLP	Coefficient for the geometry independent part of NSLP	V	0.05
PORS	Coefficient for the geometry independent part of RS	–	30
PORSB	Coefficient for the geometry independent part of RSB	–	0
PORSG	Coefficient for the geometry independent part of RSG	–	0
POSTA2	Coefficient for the geometry independent part of STA2	V	0
POSTBET	Coefficient for the geometry independent part of STBET	–	1
POSTBGIDL	Coefficient for the geometry independent part of STBGIDL	V/K	0
POSTBGIDLD	Coefficient for the geometry independent part of STBGIDL for drain side	V/K	0
POSTCS	Coefficient for the geometry independent part of STCS	–	0
POSTIG	Coefficient for the geometry independent part of STIG	–	2
POSTMUE	Coefficient for the geometry independent part of STMUE	–	0
POSTRS	Coefficient for the geometry independent part of STRS	–	1
POSTTHEMU	Coefficient for the geometry independent part of STTHEMU	–	1.5
POSTTHESAT	Coefficient for the geometry independent part of STTHESAT	–	1
POSTVFB	Coefficient for the geometry independent part of STVFB	V/K	0.0005
POSTXCOR	Coefficient for the geometry independent part of STXCOR	–	0
POTHEMU	Coefficient for the geometry independent part of THEMU	–	1.5
POTHEMAT	Coefficient for the geometry independent part of THESAT	–	1
POTHEMATB	Coefficient for the geometry independent part of THESATB	–	0

Table 2.71: PSP103VA MOSFET Device Model Parameters

Parameter	Description	Units	Default
POTHSATG	Coefficient for the geometry independent part of THESATG	–	0
POTOX	Coefficient for the geometry independent part of TOX	m	2e-09
POTOXOV	Coefficient for the geometry independent part of TOXOV	m	2e-09
POTOXOVD	Coefficient for the geometry independent part of TOXOV for drain side	m	2e-09
POVFB	Coefficient for the geometry independent part of VFB	V	-1
POVNSUB	Coefficient for the geometry independent part of VNSUB	V	0
POVP	Coefficient for the geometry independent part of VP	V	0.05
POVSBNUD	Coefficient for the geometry independent part of VSBNUD	V	0
POXCOR	Coefficient for the geometry independent part of XCOR	–	0
PSTI	Grading coefficient of STI-edge component for source-bulk junction	–	0.5
PSTID	Grading coefficient of STI-edge component for drain-bulk junction	–	0.5
PWA1	Coefficient for the width dependence of A1	–	0
PWA3	Coefficient for the width dependence of A3	–	0
PWA4	Coefficient for the width dependence of A4	–	0
PWAGIDL	Coefficient for the width dependence of AGIDL	A/V <sup>3</sup>	0
PWAGIDLD	Coefficient for the width dependence of AGIDL for drain side	A/V <sup>3</sup>	0
PWALP	Coefficient for the width dependence of ALP	–	0
PWALP1	Coefficient for the width dependence of ALP1	V	0
PWALP2	Coefficient for the width dependence of ALP2	–	0
PWAX	Coefficient for the width dependence of AX	–	0
PWBETN	Coefficient for the width dependence of BETN	–	0
PWCF	Coefficient for the width dependence of CF	–	0
PWCFR	Coefficient for the width dependence of CFR	F	0
PWCFRD	Coefficient for the width dependence of CFR for drain side	F	0
PWCGBOV	Coefficient for the width dependence of CGBOV	F	0
PWCGOV	Coefficient for the width dependence of CGOV	F	0

Table 2.71: PSP103VA MOSFET Device Model Parameters

Parameter	Description	Units	Default
PWCGOVD	Coefficient for the width dependence of CGOV for drain side	F	0
PWCOX	Coefficient for the width dependence of COX	F	0
PWCS	Coefficient for the width dependence of CS	–	0
PWCT	Coefficient for the width dependence of CT	–	0
PWDELVTAC	Coefficient for the width dependence of DELVTAC	V	0
PWDPHIB	Coefficient for the width dependence of DPHIB	V	0
PWFACNEFFAC	Coefficient for the width dependence of FACNEFFAC	–	0
PWGFACTUD	Coefficient for the width dependence of GFACNUD	–	0
PWIGINV	Coefficient for the width dependence of IGINV	A	0
PWIGOV	Coefficient for the width dependence of IGOV	A	0
PWIGOVD	Coefficient for the width dependence of IGOV for drain side	A	0
PWKUOWE	Coefficient for the width dependence part of KUOWE	–	0
PWKVTHOWE	Coefficient for the width dependence part of KVTHOWE	–	0
PWMUE	Coefficient for the width dependence of MUE	–	0
PWNEFF	Coefficient for the width dependence of NEFF	$m^{-3}$	0
PWNFA	Coefficient for the width dependence of NFA	–	0
PWNFB	Coefficient for the width dependence of NFB	–	0
PWNFC	Coefficient for the width dependence of NFC	–	0
PWNOV	Coefficient for the width dependence of NOV	$m^{-3}$	0
PWNOVD	Coefficient for the width dependence of NOV for drain side	$m^{-3}$	0
PWNP	Coefficient for the width dependence of NP	$m^{-3}$	0
PWRS	Coefficient for the width dependence of RS	–	0
PWSTBET	Coefficient for the width dependence of STBET	–	0
PWSTTHESAT	Coefficient for the width dependence of STTHESAT	–	0
PWSTVFB	Coefficient for the width dependence of STVFB	V/K	0
PWTHESAT	Coefficient for the width dependence of THESAT	–	0
PWTHESATB	Coefficient for the width dependence of THESATB	–	0

Table 2.71: PSP103VA MOSFET Device Model Parameters

Parameter	Description	Units	Default
PWTHESATG	Coefficient for the width dependence of THESATG	—	0
PWVFB	Coefficient for the width dependence of VFB	V	0
PWXCOR	Coefficient for the width dependence of XCOR	—	0
QMC	Quantum-mechanical correction factor	—	1
RBULK	Bulk resistance between node BP and BI	$\Omega$	0
RBULK0	Bulk resistance between node BP and BI	$\Omega$	0
RDE	External drain resistance	$\Omega$	0
RG	Gate resistance	$\Omega$	0
RGO	Gate resistance	$\Omega$	0
RINT	Contact resistance between silicide and ploy	$\Omega \text{ m}^2$	0
RJUND	Drain-side bulk resistance between node BI and BD	$\Omega$	0
RJUNDO	Drain-side bulk resistance between node BI and BD	$\Omega$	0
RJUNS	Source-side bulk resistance between node BI and BS	$\Omega$	0
RJUNSO	Source-side bulk resistance between node BI and BS	$\Omega$	0
RS	Series resistance at TR	$\Omega$	30
RSB	Back-bias dependence of series resistance	$V^{-1}$	0
RSBO	Back-bias dependence of series resistance	$V^{-1}$	0
RSE	External source resistance	$\Omega$	0
RSG	Gate-bias dependence of series resistance	$V^{-1}$	0
RSGO	Gate-bias dependence of series resistance	$V^{-1}$	0
RSH	Sheet resistance of source diffusion	$\Omega/\square$	0
RSHD	Sheet resistance of drain diffusion	$\Omega/\square$	0
RSHG	Gate electrode diffusion sheet resistance	$\Omega/\square$	0
RSW1	Source/drain series resistance for 1 $\mu\text{m}$ wide channel at TR	$\Omega$	50
RSW2	Higher-order width scaling of RS	—	0
RVPOLY	Vertical poly resistance	$\Omega \text{ m}^2$	0
RWELL	Well resistance between node BI and B	$\Omega$	0
RWELLO	Well resistance between node BI and B	$\Omega$	0
SAREF	Reference distance between OD-edge and poly from one side	m	1e-06

Table 2.71: PSP103VA MOSFET Device Model Parameters

Parameter	Description	Units	Default
SBREF	Reference distance between OD-edge and poly from other side	m	1e-06
SCREF	Distance between OD-edge and well edge of a reference device	m	1e-06
STA2	Temperature dependence of A2	V	0
STA20	Temperature dependence of A2	V	0
STBET	Temperature dependence of BETN	–	1
STBETL	Length dependence of temperature dependence of BETN	–	0
STBETLW	Area dependence of temperature dependence of BETN	–	0
STBETO	Geometry independent temperature dependence of BETN	–	1
STBETW	Width dependence of temperature dependence of BETN	–	0
STBGIDL	Temperature dependence of BGIDL	V/K	0
STBGIDLD	Temperature dependence of BGIDL for drain side	V/K	0
STBGIDLDO	Temperature dependence of BGIDL for drain side	V/K	0
STBGIDL0	Temperature dependence of BGIDL	V/K	0
STCS	Temperature dependence of CS	–	0
STCS0	Temperature dependence of CS	–	0
STETA0	eta0 shift factor related to VTHO change	m	0
STFBBTBOT	Temperature scaling parameter for band-to-band tunneling of bottom component for source-bulk junction	–	-0.001
STFBBTBOTD	Temperature scaling parameter for band-to-band tunneling of bottom component for drain-bulk junction	–	-0.001
STFBBTGAT	Temperature scaling parameter for band-to-band tunneling of gate-edge component for source-bulk junction	–	-0.001
STFBBTGATD	Temperature scaling parameter for band-to-band tunneling of gate-edge component for drain-bulk junction	–	-0.001
STFBBTSTI	Temperature scaling parameter for band-to-band tunneling of STI-edge component for source-bulk junction	–	-0.001
STFBBTSTID	Temperature scaling parameter for band-to-band tunneling of STI-edge component for drain-bulk junction	–	-0.001
STIG	Temperature dependence of IGINV and IGOV	–	2

Table 2.71: PSP103VA MOSFET Device Model Parameters

Parameter	Description	Units	Default
STIGO	Temperature dependence of IGINV and IGOV	–	2
STMUE	Temperature dependence of MUE	–	0
STMUEO	Temperature dependence of MUE	–	0
STRS	Temperature dependence of RS	–	1
STRSO	Temperature dependence of RS	–	1
STTHEMU	Temperature dependence of THEMU	–	1.5
STTHEMUO	Temperature dependence of THEMU	–	1.5
STTHESAT	Temperature dependence of THESAT	–	1
STTHESATL	Length dependence of temperature dependence of THESAT	–	0
STTHESATLW	Area dependence of temperature dependence of THESAT	–	0
STTHESATO	Geometry independent temperature dependence of THESAT	–	1
STTHESATW	Width dependence of temperature dependence of THESAT	–	0
STVFB	Temperature dependence of VFB	V/K	0.0005
STVFBL	Length dependence of temperature dependence of VFB	V/K	0
STVFBLW	Area dependence of temperature dependence of VFB	V/K	0
STVFB0	Geometry-independent temperature dependence of VFB	V/K	0.0005
STVFBW	Width dependence of temperature dependence of VFB	V/K	0
STXCOR	Temperature dependence of XCOR	–	0
STXCORO	Temperature dependence of XCOR	–	0
SWDELVTAC	Flag for separate capacitance calculation; 0=off, 1=on	–	0
SWGEO	Flag for geometrical model, 0=local, 1=global, 2=binning	–	1
SWGIDL	Flag for GIDL current, 0=turn off IGIDL	–	0
SWIGATE	Flag for gate current, 0=turn off IG	–	0
SWIMPACT	Flag for impact ionization current, 0=turn off II	–	0
SWJUNASYM	Flag for asymmetric junctions; 0=symmetric, 1=asymmetric	–	0
SWJUNCAP	Flag for juncap, 0=turn off juncap	–	0
SWJUNEXP	Flag for JUNCAP-express; 0=full model, 1=express model	–	0

Table 2.71: PSP103VA MOSFET Device Model Parameters

Parameter	Description	Units	Default
SWNUD	Flag for NUD-effect; 0=off, 1=on, 2=on+CV-correction	–	0
THEMU	Mobility reduction exponent at TR	–	1.5
THEMU0	Mobility reduction exponent at TR	–	1.5
THESAT	Velocity saturation parameter at TR	$V^{-1}$	1
THESATB	Back-bias dependence of velocity saturation	$V^{-1}$	0
THESATB0	Back-bias dependence of velocity saturation	$V^{-1}$	0
THESATG	Gate-bias dependence of velocity saturation	$V^{-1}$	0
THESATG0	Gate-bias dependence of velocity saturation	$V^{-1}$	0
THESATL	Length dependence of THESAT	$V^{-1}$	0.05
THESATLEXP	Exponent for length dependence of THESAT	–	1
THESATLW	Area dependence of velocity saturation parameter	–	0
THESATO	Geometry independent velocity saturation parameter at TR	$V^{-1}$	0
THESATW	Width dependence of velocity saturation parameter	–	0
TKU0	Temperature dependence of KU0	–	0
TOX	Gate oxide thickness	m	2e-09
TOX0	Gate oxide thickness	m	2e-09
TOXOV	Overlap oxide thickness	m	2e-09
TOXOVD	Overlap oxide thickness for drain side	m	2e-09
TOXOVDO	Overlap oxide thickness for drain side	m	2e-09
TOXOVO	Overlap oxide thickness	m	2e-09
TR	nominal (reference) temperature	°C	21
TRJ	reference temperature	–	21
TYPE	Channel type parameter, +1=NMOS -1=PMOS	–	1
U0	Zero-field mobility at TR	$m^2/(Vs)$	0.05
VBIRBOT	Built-in voltage at the reference temperature of bottom component for source-bulk junction	–	1
VBIRBOTD	Built-in voltage at the reference temperature of bottom component for drain-bulk junction	–	1
VBIRGAT	Built-in voltage at the reference temperature of gate-edge component for source-bulk junction	–	1
VBIRGATD	Built-in voltage at the reference temperature of gate-edge component for drain-bulk junction	–	1
VBIRSTI	Built-in voltage at the reference temperature of STI-edge component for source-bulk junction	–	1

Table 2.71: PSP103VA MOSFET Device Model Parameters

Parameter	Description	Units	Default
VBIRSTID	Built-in voltage at the reference temperature of STI-edge component for drain-bulk junction	–	1
VBRBOT	Breakdown voltage of bottom component for source-bulk junction	–	10
VBRBOTD	Breakdown voltage of bottom component for drain-bulk junction	–	10
VBRGAT	Breakdown voltage of gate-edge component for source-bulk junction	–	10
VBRGATD	Breakdown voltage of gate-edge component for drain-bulk junction	–	10
VBRSTI	Breakdown voltage of STI-edge component for source-bulk junction	–	10
VBRSTID	Breakdown voltage of STI-edge component for drain-bulk junction	–	10
VFB	Flat band voltage at TR	V	-1
VFBL	Length dependence of flat-band voltage	V	0
VFBLW	Area dependence of flat-band voltage	V	0
VFBO	Geometry-independent flat-band voltage at TR	V	-1
VFBW	Width dependence of flat-band voltage	V	0
VJUNREF	Typical maximum source-bulk junction voltage; usually about 2*VSUP	–	2.5
VJUNREFD	Typical maximum drain-bulk junction voltage; usually about 2*VSUP	–	2.5
VNSUB	Effective doping bias-dependence parameter	V	0
VNSUB0	Effective doping bias-dependence parameter	V	0
VP	CLM logarithm dependence factor	V	0.05
VPO	CLM logarithmic dependence parameter	V	0.05
VSBNUD	Lower Vsb value for NUD-effect	V	0
VSBNUDO	Lower Vsb value for NUD-effect	V	0
WBET	Characteristic width for width scaling of BETN	m	1e-09
WEB	Coefficient for SCB	–	0
WEC	Coefficient for SCC	–	0
WKUO	Width dependence of KUO	–	0
WKVTH0	Width dependence of KVTH0	–	0
WLOD	Width parameter	m	0
WLODKUO	Width parameter for UO stress effect	–	0
WLODVTH	Width parameter for VTH-stress effect	–	0
WMAX	Dummy parameter to label binning set	m	1

Table 2.71: PSP103VA MOSFET Device Model Parameters

Parameter	Description	Units	Default
WMIN	Dummy parameter to label binning set	m	0
WOT	Effective channel width reduction per side	m	0
WSEG	Char. length of segregation of background doping NSUBO	m	1e-08
WSEGP	Char. length of segregation of pocket doping NPCK	m	1e-08
WVARL	Length dependence of WVAR	—	0
WVARO	Geom. independent difference between actual and programmed field-oxide opening	m	0
WVARW	Width dependence of WVAR	—	0
XCOR	Non-universality factor	$V^{-1}$	0
XCORL	Length dependence of non-universality parameter	—	0
XCORLW	Area dependence of non-universality parameter	—	0
XCORO	Geometry independent non-universality parameter	$V^{-1}$	0
XCORW	Width dependence of non-universality parameter	—	0
XJUNGAT	Junction depth of gate-edge component for source-bulk junction	—	1e-07
XJUNGATD	Junction depth of gate-edge component for drain-bulk junction	—	1e-07
XJUNSTI	Junction depth of STI-edge component for source-bulk junction	—	1e-07
XJUNSTID	Junction depth of STI-edge component for drain-bulk junction	—	1e-07

## Level 107 MOSFET Tables (BSIM CMG version 107.0.0)

**Xyce** includes the BSIM CMG Common Multi-gate model version 107. The code in **Xyce** was generated from the BSIM group's Verilog-A input using the default "ifdef" lines provided, and therefore supports only the subset of BSIM CMG features those defaults enable. Instance and model parameters for the BSIM CMG model are given in tables 2.72 and 2.73. Details of the model are documented in the BSIM-CMG technical report[25], available from the BSIM web site at <http://www-device.eecs.berkeley.edu/bsim/?page=BSIMCMG>.

Table 2.72: BSIM-CMG FINFET v107.0.0 Device Instance Parameters

Parameter	Description	Units	Default
ADEJ	Drain junction area (BULKMOD=1)	—	0
ADEO	Drain to substrate overlap area through oxide	—	0
ASEJ	Source junction area (BULKMOD=1)	—	0
ASEO	Source to substrate overlap area through oxide	—	0
CDSP	Constant drain-to-source fringe capacitance (All CGEOMOD)	—	0
CGDP	Constant gate-to-drain fringe capacitance (CGEOMOD=1)	—	0
CGSP	Constant gate-to-source fringe capacitance (CGEOMOD=1)	—	0
COVD	Constant g/d overlap capacitance (CGEOMOD=1)	—	0
COVS	Constant g/s overlap capacitance (CGEOMOD=1)	—	0
D	Diameter of the cylinder (GEOMOD=3)	—	4e-08
FPITCH	Fin pitch	—	8e-08
L	Designed Gate Length	—	3e-08
LRSD	Length of the source/drain	—	0
NFIN	Number of fins per finger (real number enables optimization)	—	1
NGCON	number of gate contact (1 or 2 sided)	—	1
NRD	Number of source diffusion squares	—	0
NRS	Number of source diffusion squares	—	0
PDEJ	Drain to substrate PN junction perimeter (BULKMOD=1)	—	0
PDEO	Perimeter of drain to substrate overlap region through oxide	—	0
PSEJ	Source to substrate PN junction perimeter (BULKMOD=1)	—	0
PSEO	Perimeter of source to substrate overlap region through oxide	—	0

Table 2.72: BSIM-CMG FINFET v107.0.0 Device Instance Parameters

Parameter	Description	Units	Default
TFIN	Body (Fin) thickness	–	1.5e-08

Table 2.73: BSIM-CMG FINFET v107.0.0 Device Model Parameters

Parameter	Description	Units	Default
A1	Non-saturation effect parameter for strong inversion region	–	0
A11	Temperature dependence of A1	–	0
A2	Non-saturation effect parameter for moderate inversion region	–	0
A21	Temperature dependence of A2	–	0
ADEJ	Drain junction area (BULKMOD=1)	–	0
ADEO	Drain to substrate overlap area through oxide	–	0
AEU		–	0
AGIDL	pre-exponential coeff. for GIDL in mho	–	0
AGISL	pre-exponential coeff. for GISL in mho	–	6.055e-12
AIGBACC	parameter for Igb in accumulation	–	0.0136
AIGBACC1	parameter for Igb in accumulation	–	0
AIGBINV	parameter for Igb in inversion	–	0.0111
AIGBINV1	parameter for Igb in inversion	–	0
AIGC	parameter for Igc in inversion	–	0.0136
AIGC1	parameter for Igc in inversion	–	0
AIGD	parameter for Igd in inversion	–	0
AIGD1	parameter for Igd in inversion	–	0
AIGEN	Thermal Generation Current Parameter	–	0
AIGS	parameter for Igs in inversion	–	0.0136
AIGS1	parameter for Igs in inversion	–	0
ALPHA0	first parameter of Iii	m/V	0
ALPHA01	Temperature dependence of ALPHA0, m/V/degrees	–	0
ALPHA1	L scaling parameter of Iii	V <sup>-1</sup>	0
ALPHA11	Temperature dependence ALPHA1, 1/V/degree	–	0
ALPHAII0	first parameter of Iii for IIMOD=2, m/V	–	0
ALPHAII01	Temperature dependence of ALPHAII0, m/V/degrees	–	0

Table 2.73: BSIM-CMG FINFET v107.0.0 Device Model Parameters

Parameter	Description	Units	Default
ALPHAIII1	L scaling parameter of $I_{ii}$ for IIMOD=2	$V^{-1}$	0
ALPHAIII11	Temperature dependence of ALPHAIII1, $1/V/\text{degrees}$	—	0
AMEXP		—	0
AMEXPR		—	0
APCLM		—	0
APSAT		—	0
APSATCV		—	0
APTWG		—	0
AQMTCCN	Parameter for Geometric dependence of $T_{cen}$ on $R/T_{FIN}/H_{FIN}$	—	0
ARDSW		—	0
ARDW		—	0
ARSDEND		—	0
ARSW		—	0
ASEJ	Source junction area (BULKMOD=1)	—	0
ASEO	Source to substrate overlap area through oxide	—	0
ASILIEND		—	0
ASYMMOD	Asymmetric model selector	—	0
AT		—	-0.00156
AUA		—	0
AUD		—	0
AVSAT		—	0
AVSAT1		—	0
AVSATCV		—	0
BETA0	$V_{ds}$ dependent parameter of $I_{ii}$	$V^{-1}$	0
BETAI0	$V_{ds}$ dependent parameter of $I_{ii}$	$V^{-1}$	0
BETAI1	$V_{ds}$ dependent parameter of $I_{ii}$	—	0
BETAI2	$V_{ds}$ dependent parameter of $I_{ii}$ , V	—	0.1
BEU		—	1e-07
BGOSUB	Band gap of substrate at 300.15K, eV	—	1.12
BGIDL	exponential coeff. for GIDL	V/m	0
BGISL	exponential coeff. for GISL	V/m	3e+08
BIGBACC	parameter for $I_{gb}$ in accumulation	—	0.00171
BIGBINV	parameter for $I_{gb}$ in inversion	—	0.000949

Table 2.73: BSIM-CMG FINFET v107.0.0 Device Model Parameters

Parameter	Description	Units	Default
BIGC	parameter for $I_{gc}$ in inversion	—	0.00171
BIGD	parameter for $I_{gd}$ in inversion	—	0
BIGEN	Thermal Generation Current Parameter	—	0
BIGS	parameter for $I_{gs}$ in inversion	—	0.00171
BMEXP		—	1
BMEXPR		—	0
BPCLM		—	1e-07
BPSAT		—	1
BPSATCV		—	0
BPTWG		—	1e-07
BQMTCEN	Parameter for Geometric dependence of $T_{cen}$ on $R/T_{FIN}/H_{FIN}$	—	1.2e-08
BRDSW		—	1e-07
BRDW		—	1e-07
BRSW		—	1e-07
BUA		—	1e-07
BUD		—	5e-08
BULKMOD	Bulk model	—	0
BVD	Drain diode breakdown voltage	—	0
BVS	Source diode breakdown voltage	—	10
BVSAT		—	1e-07
BVSAT1		—	0
BVSATCV		—	0
CAPMOD	Accumulation region capacitance model selector	—	0
CDSC	coupling capacitance between S/D and channel	—	0.007
CDSCD	drain-bias sensitivity of CDSC	—	0.007
CDSCDN1	NFIN dependence of CDSCD	—	0
CDSCDN2	NFIN dependence of CDSCD	—	100000
CDSCDR	Reverse-mode drain-bias sensitivity of CDSC (Experimental)	—	0
CDSCDRN1	NFIN dependence of CDSCD	—	0
CDSCDRN2	NFIN dependence of CDSCD	—	0
CDSCN1	NFIN dependence of CDSC	—	0
CDSCN2	NFIN dependence of CDSC	—	100000

Table 2.73: BSIM-CMG FINFET v107.0.0 Device Model Parameters

Parameter	Description	Units	Default
CDSP	Constant drain-to-source fringe capacitance (All CGEOMOD)	—	0
CFD	Outer Fringe Cap (drain side)	—	0
CFS	Outer Fringe Cap (source side)	—	2.5e-11
CGBL	Bias dependent component of Gate to substrate overlap cap	—	0
CGBN	Gate to substrate overlap cap per unit channel length per fin per finger	—	0
CGB0	Gate to substrate overlap cap per unit channel length per finger per NGCON	—	0
CGDL		—	0
CGD0	Non LDD region drain-gate overlap capacitance per unit channel width	—	0
CGDP	Constant gate-to-drain fringe capacitance (CGEOMOD=1)	—	0
CGEO1SW		—	0
CGEOA	Fitting parameter for CGEOMOD=2	—	1
CGEOB	Fitting parameter for CGEOMOD=2	—	0
CGEOC	Fitting parameter for CGEOMOD=2	—	0
CGEOD	Fitting parameter for CGEOMOD=2	—	0
CGEOE	Fitting parameter for CGEOMOD=2	—	1
CGEOMOD	parasitic capacitance model selector	—	0
CGIDL	parameter for body-effect of GIDL	V <sup>3</sup>	0
CGISL	parameter for body-effect of GISL	V <sup>3</sup>	0.5
CGSL		—	0
CGS0	Non LDD region source-gate overlap capacitance per unit channel width	—	0
CGSP	Constant gate-to-source fringe capacitance (CGEOMOD=1)	—	0
CHARGEWF	Average Channel Charge Weighting Factor, +1:source-side, 0:middle, -1:drain-side	—	0
CIGBACC	parameter for Igb in accumulation	—	0.075
CIGBINV	parameter for Igb in inversion	—	0.006
CIGC	parameter for Igc in inversion	—	0.075
CIGD	parameter for Igd in inversion	—	0
CIGS	parameter for Igs in inversion	—	0.075
CIT	parameter for interface trap	—	0

Table 2.73: BSIM-CMG FINFET v107.0.0 Device Model Parameters

Parameter	Description	Units	Default
CJD	Unit area drain-side junction capacitance at zero bias	—	0
CJS	Unit area source-side junction capacitance at zero bias	—	0.0005
CJSWD	Unit length drain-side sidewall junction capacitance at zero bias	—	0
CJSWGD	Unit length drain-side gate sidewall junction capacitance at zero bias	—	0
CJSWGS	Unit length source-side gate sidewall junction capacitance at zero bias	—	0
CJSWS	Unit length source-side sidewall junction capacitance at zero bias	—	5e-10
CKAPPAB		—	0.6
CKAPPAD		—	0
CKAPPAS		—	0.6
COREMOD	Surface potential algorithm	—	0
COVD	Constant g/d overlap capacitance (CGEOMOD=1)	—	0
COVS	Constant g/s overlap capacitance (CGEOMOD=1)	—	0
CRATIO		—	0.5
CSDESW	Coefficient for source/drain to substrate sidewall cap	—	0
CTHO	Thermal capacitance	—	1e-05
D	Diameter of the cylinder (GEOMOD=3)	—	4e-08
DELTAPRSD		—	0
DELTA VSAT		—	1
DELTA VSATCV		—	0
DELTA W	change of effective width due to shape of fin/cylinder	—	0
DELTA WCV	CV change of effective width due to shape of fin/cylinder	—	0
DEL VFBACC	Change in Flatband Voltage; Vfb_accumulation-Vfb_inversion	—	0
DEL VTRAND	Variability in Vth	—	0
DEVTYPE		—	1
DLBIN	Delta L for Binning	—	0
DLC	Delta L for C-V model	—	0

Table 2.73: BSIM-CMG FINFET v107.0.0 Device Model Parameters

Parameter	Description	Units	Default
DLCACC	Delta L for C-V model in accumulation region (CAPMOD=1, BULKMOD=1)	–	0
DLCIGD	Delta L for Igd model	–	0
DLCIGS	Delta L for Igs model	–	0
DROUT		–	1.06
DSUB	DIBL exponent coefficient	–	1.06
DTEMP	Variability in Device Temperature	–	0
DVTO	SCE coefficient	–	0
DVT1	SCE exponent coefficient, after binning should be in (0:inf)	–	0.6
DVT1SS	Subthreshold Swing exponent coefficient, after binning should be in (0:inf)	–	0
DVTP0	Coefficient for Drain-Induced Vth Shift (DITS)	–	0
DVTP1	DITS exponent coefficient	–	0
DVTSHIFT	Vth shift handle	–	0
EASUB	Electron affinity of substrate, eV	–	4.05
EF		–	1
EGIDL	band bending parameter for GIDL	V	0
EGISL	band bending parameter for GISL	V	0.2
EIGBINV	parameter for Igb in inversion	–	1.1
EM		–	4.1e+07
EMOBT		–	0
EOT	equivalent oxide thickness in meters	–	1e-09
EOTACC	equivalent oxide thickness for accumulation region in meters	m	0
EOTBOX	equivalent oxide thickness of the buried oxide (SOI FinFET) or STI (bulk FinFET) in meters	–	1.4e-07
EPSROX	Relative dielectric constant of the gate dielectric	–	3.9
EPSRSP	Relative dielectric constant of the spacer	–	3.9
EPSRSUB	Relative dielectric constant of the channel material	–	11.9
ESATII	Saturation channel E-Field for Iii	V/m	1e+07
ETA0	DIBL coefficient	–	0.6
ETAON1	NFIN dependence of ETA0	–	0
ETAON2	NFIN dependence of ETA0	–	100000
ETAOR	Reverse-mode DIBL coefficient (Experimental)	–	0

Table 2.73: BSIM-CMG FINFET v107.0.0 Device Model Parameters

Parameter	Description	Units	Default
ETAMOB		—	2
ETAQM	Bulk charge coefficient for Tcen	—	0.54
EU		—	2.5
FECH	End-channel factor, for different orientation/shape	—	1
FECHCV	CV end-channel factor, for different orientation/shape	—	1
FPITCH	Fin pitch	—	8e-08
GEOMOD	Geometry mode selector	—	1
GIDLMOD	GIDL/GISL current switcher	—	0
HEPI	Height of the raised source/drain on top of the fin	—	1e-08
HFIN	Fin height in meters	—	3e-08
IDSOMULT	Variability in Drain current for misc. reasons	—	1
IGBMOD	model selector for Igb	—	0
IGCMOD	model selector for Igc, Igs, and Igd	—	0
IGT	Gate Current Temperature Dependence	—	2.5
IIMOD	Impact ionization model switch	—	0
IIT	Impact Ionization Temperature Dependence, IIMOD=1	—	-0.5
IJTHDFWD	Forward drain diode breakdown limiting current	—	0
IJTHDREV	Reverse drain diode breakdown limiting current	—	0
IJTHSFWD	Forward source diode breakdown limiting current	—	0.1
IJTHSREV	Reverse source diode breakdown limiting current	—	0.1
IMIN	Parameter for Vgs Clamping for inversion region calc. in accumulation	—	1e-15
JSD	Bottom drain junction reverse saturation current density	—	0
JSS	Bottom source junction reverse saturation current density	—	0.0001
JSWD	Unit length reverse saturation current for sidewall drain junction	—	0
JSWGD	Unit length reverse saturation current for gate-edge sidewall drain junction	—	0
JSWGS	Unit length reverse saturation current for gate-edge sidewall source junction	—	0
JSWS	Unit length reverse saturation current for sidewall source junction	—	0

Table 2.73: BSIM-CMG FINFET v107.0.0 Device Model Parameters

Parameter	Description	Units	Default
JTSD	Bottom drain junction trap-assisted saturation current density	—	0
JTSS	Bottom source junction trap-assisted saturation current density	—	0
JTSSWD	Unit length trap-assisted saturation current for sidewall drain junction	—	0
JTSSWGD	Unit length trap-assisted saturation current for gate-edge sidewall drain junction	—	0
JTSSWGS	Unit length trap-assisted saturation current for gate-edge sidewall source junction	—	0
JTSSWS	Unit length trap-assisted saturation current for sidewall source junction	—	0
JTWEFF	Trap assisted tunneling current width dependence	—	0
K0	Lateral NUD voltage parameter, V	—	0
K01	Temperature dependence of lateral NUD voltage parameter, V/K	—	0
K0SI	Correction factor for strong inversion, used in Mnud, after binnig should be from (0:inf)	—	1
K0SI1	Temperature dependence of K0SI, 1/K	—	0
K1	Body effect coefficient for sub-threshold region	—	0
K11	Temperature dependence of K1	—	0
K1RSCE	K1 for reverse short channel effect calculation	—	0
K1SAT	Correction factor for K1 in saturation (high Vds)	—	0
K1SAT1	Temperature dependence of K1SAT1	—	0
K1SI	Correction factor for strong inversion, used in Mob	—	0
K1SI1	Temperature dependence of K1SI, 1/K	—	0
KSATIV		—	1
KT1	Vth Temperature Coefficient (V)	—	0
KT1L	Vth Temperature L Coefficient (m-V)	—	0
L	Designed Gate Length	—	3e-08
LA1		—	0
LA11		—	0
LA2		—	0
LA21		—	0
LAGIDL		—	0

Table 2.73: BSIM-CMG FINFET v107.0.0 Device Model Parameters

Parameter	Description	Units	Default
LAGISL		—	0
LAIGBACC		—	0
LAIGBACC1		—	0
LAIGBINV		—	0
LAIGBINV1		—	0
LAIGC		—	0
LAIGC1		—	0
LAIGD		—	0
LAIGD1		—	0
LAIGEN		—	0
LAIGS		—	0
LAIGS1		—	0
LALPHA0		—	0
LALPHA1		—	0
LALPHAIIO		—	0
LALPHAI11		—	0
LAT		—	0
LBETA0		—	0
LBETAIIO		—	0
LBETAI11		—	0
LBETAI12		—	0
LBGIDL		—	0
LBGISL		—	0
LBIGBACC		—	0
LBIGBINV		—	0
LBIGC		—	0
LBIGD		—	0
LBIGEN		—	0
LBIGS		—	0
LCDSC		—	0
LCDSCD		—	0
LCDSCDR		—	0
LCFD		—	0
LCFS		—	0

Table 2.73: BSIM-CMG FINFET v107.0.0 Device Model Parameters

Parameter	Description	Units	Default
LCGBL		—	0
LCGDL		—	0
LCGIDL		—	0
LCGISL		—	0
LCGSL		—	0
LCIGBACC		—	0
LCIGBINV		—	0
LCIGC		—	0
LCIGD		—	0
LCIGS		—	0
LCIT		—	0
LCKAPPAB		—	0
LCKAPPAD		—	0
LCKAPPAS		—	0
LCOVD		—	0
LCOVS		—	0
LDELTAVSAT		—	0
LDELTAVSATCV		—	0
LDROUT		—	0
LDSUB		—	0
LDVT0		—	0
LDVT1		—	0
LDVT1SS		—	0
LDVTB		—	0
LDVTSHIFT		—	0
LEGIDL		—	0
LEGISL		—	0
LEIGBINV		—	0
LEMOBT		—	0
LESATII		—	0
LETA0		—	0
LETAOR		—	0
LETAMOB		—	0
LEU		—	0

Table 2.73: BSIM-CMG FINFET v107.0.0 Device Model Parameters

Parameter	Description	Units	Default
LIGT		—	0
LII	Channel length dependent parameter of lii	Vm	5e-10
LIIT		—	0
LINT	Length reduction parameter (dopant diffusion effect)	—	0
LINTIGEN	Lint for Thermal Generation Current	—	0
LINTNOI		—	0
LK0		—	0
LK01		—	0
LK0SI		—	0
LK0SI1		—	0
LK1		—	0
LK11		—	0
LK1RSCE		—	0
LK1SAT		—	0
LK1SAT1		—	0
LK1SI		—	0
LK1SI1		—	0
LKSATIV		—	0
LKT1		—	0
LL	Length reduction parameter (dopant diffusion effect)	—	0
LLC	Length reduction parameter (dopant diffusion effect)	—	0
LLII		—	0
LLN	Length reduction parameter (dopant diffusion effect)	—	1
LLPEO		—	0
LLPEB		—	0
LMEXP		—	0
LMEXPR		—	0
LNBODY		—	0
LNGATE		—	0
LNIGBACC		—	0
LNIGBINV		—	0

Table 2.73: BSIM-CMG FINFET v107.0.0 Device Model Parameters

Parameter	Description	Units	Default
LNTGEN		—	0
LNTOX		—	0
LPA		—	1
LPCLM		—	0
LPCLMCV		—	0
LPCLMG		—	0
LPDIBL1		—	0
LPDIBL1R		—	0
LPDIBL2		—	0
LPEO	Equivalent length of pocket region at zero bias	—	5e-09
LPGIDL		—	0
LPGISL		—	0
LPHIBE		—	0
LPHIG		—	0
LPHIN		—	0
LPIGCD		—	0
LPOXEDGE		—	0
LPRT		—	0
LPRWGD		—	0
LPRWGS		—	0
LPSAT		—	0
LPSATCV		—	0
LPTWG		—	0
LPTWGR		—	0
LPTWGT		—	0
LPVAG		—	0
LQMFACTOR		—	0
LQMTCENCV		—	0
LQMTCENCVA		—	0
LQMTCENIV		—	0
LRDSW		—	0
LRDW		—	0
LRSD	Length of the source/drain	—	0
LRSW		—	0

Table 2.73: BSIM-CMG FINFET v107.0.0 Device Model Parameters

Parameter	Description	Units	Default
LSII0		—	0
LSII1		—	0
LSII2		—	0
LSIID		—	0
LSP		—	0
LSTTHETASAT		—	0
LTGIDL		—	0
LTII		—	0
LTSS		—	0
LU0		—	0
LUA		—	0
LUA1		—	0
LUC		—	0
LUC1		—	0
LUCS		—	0
LUCSTE		—	0
LUD		—	0
LUD1		—	0
LUP		—	0
LUTE		—	0
LUTL		—	0
LVSAT		—	0
LVSAT1		—	0
LVSAT1R		—	0
LVSATCV		—	0
LWR		—	0
LXRCRG1		—	0
LXRCRG2		—	0
MEXP		—	4
MEXPR		—	0
MJD	Drain bottom junction capacitance grading coefficient	—	0
MJD2	Drain bottom two-step second junction capacitance grading coefficient	—	0

Table 2.73: BSIM-CMG FINFET v107.0.0 Device Model Parameters

Parameter	Description	Units	Default
MJS	Source bottom junction capacitance grading coefficient	—	0.5
MJS2	Source bottom two-step second junction capacitance grading coefficient	—	0.125
MJSWD	Drain sidewall junction capacitance grading coefficient	—	0
MJSWD2	Drain sidewall two-step second junction capacitance grading coefficient	—	0
MJSWGD	Drain-side gate sidewall junction capacitance grading coefficient	—	0
MJSWGD2	Drain-side gate sidewall two-step	—	0
MJSWGS	Source-side gate sidewall junction capacitance grading coefficient	—	0
MJSWGS2	Source-side gate sidewall two-step	—	0
MJSWS	Source sidewall junction capacitance grading coefficient	—	0.33
MJSWS2	Source sidewall two-step second junction capacitance grading coefficient	—	0.083
NA1		—	0
NA11		—	0
NA2		—	0
NA21		—	0
NAGIDL		—	0
NAGISL		—	0
NAIGBACC		—	0
NAIGBACC1		—	0
NAIGBINV		—	0
NAIGBINV1		—	0
NAIGC		—	0
NAIGC1		—	0
NAIGD		—	0
NAIGD1		—	0
NAIGEN		—	0
NAIGS		—	0
NAIGS1		—	0
NALPHA0		—	0
NALPHA1		—	0

Table 2.73: BSIM-CMG FINFET v107.0.0 Device Model Parameters

Parameter	Description	Units	Default
NALPHAII0		—	0
NALPHAII1		—	0
NAT		—	0
NBETA0		—	0
NBETAII0		—	0
NBETAII1		—	0
NBETAII2		—	0
NBGIDL		—	0
NBGISL		—	0
NBIGBACC		—	0
NBIGBINV		—	0
NBIGC		—	0
NBIGD		—	0
NBIGEN		—	0
NBIGS		—	0
NBODY	channel (body) doping	—	1e+22
NBODYN1	NFIN dependence of channel (body) doping	—	0
NBODYN2	NFIN dependence of channel (body) doping	—	100000
NCOSUB	Conduction band density of states, m-3	—	2.86e+25
NCDSC		—	0
NCDSCD		—	0
NCDSCDR		—	0
NCFD		—	0
NCFS		—	0
NCGBL		—	0
NCGDL		—	0
NCGIDL		—	0
NCGISL		—	0
NCGSL		—	0
NCIGBACC		—	0
NCIGBINV		—	0
NCIGC		—	0
NCIGD		—	0
NCIGS		—	0

Table 2.73: BSIM-CMG FINFET v107.0.0 Device Model Parameters

Parameter	Description	Units	Default
NCIT		—	0
NCKAPPAB		—	0
NCKAPPAD		—	0
NCKAPPAS		—	0
NCOVD		—	0
NCOVS		—	0
NDELTAVSAT		—	0
NDELTAVSATCV		—	0
NDROUT		—	0
NDSUB		—	0
NDVTO		—	0
NDVT1		—	0
NDVT1SS		—	0
NDVTB		—	0
NDVTSHIFT		—	0
NEGIDL		—	0
NEGISL		—	0
NEIGBINV		—	0
NEMOBT		—	0
NESATII		—	0
NETAO		—	0
NETAOR		—	0
NETAMOB		—	0
NEU		—	0
NF	Number of fingers	—	1
NFIN	Number of fins per finger (real number enables optimization)	—	1
NGATE	Parameter for Poly Gate Doping, for metal gate please set NGATE = 0	—	0
NGCON	number of gate contact (1 or 2 sided)	—	1
NIOSUB	Intrinsic carrier constant at 300.15K, m-3	—	1.1e+16
NIGBACC	parameter for Igb in accumulation	—	1
NIGBINV	parameter for Igb in inversion	—	3
NIGT		—	0
NIIT		—	0

Table 2.73: BSIM-CMG FINFET v107.0.0 Device Model Parameters

Parameter	Description	Units	Default
NJD	Drain junction emission coefficient	—	0
NJS	Source junction emission coefficient	—	1
NJTS	Non-ideality factor for JTSS	—	20
NJTSD	Non-ideality factor for JTSD	—	0
NJTSSW	Non-ideality factor for JTSSWS	—	20
NJTSSWD	Non-ideality factor for JTSSWD	—	0
NJTSSWG	Non-ideality factor for JTSSWGS	—	20
NJTSSWGD	Non-ideality factor for JTSSWGD	—	0
NK0		—	0
NK01		—	0
NKOSI		—	0
NKOSI1		—	0
NK1		—	0
NK11		—	0
NK1RSCE		—	0
NK1SAT		—	0
NK1SAT1		—	0
NK1SI		—	0
NK1SI1		—	0
NKSATIV		—	0
NKT1		—	0
NLII		—	0
NLPEO		—	0
NLPEB		—	0
NMEXP		—	0
NMEXPR		—	0
NNBODY		—	0
NNGATE		—	0
NNIGBACC		—	0
NNIGBINV		—	0
NNTGEN		—	0
NNTOX		—	0
NOIA		—	6.25e+39
NOIB		—	3.125e+24

Table 2.73: BSIM-CMG FINFET v107.0.0 Device Model Parameters

Parameter	Description	Units	Default
NOIC		—	8.75e+07
NPCLM		—	0
NPCLMCV		—	0
NPCLMG		—	0
NPDI BL1		—	0
NPDI BL1R		—	0
NPDI BL2		—	0
NPGIDL		—	0
NPGISL		—	0
NPHIBE		—	0
NPHIG		—	0
NPHIN		—	0
NPIGCD		—	0
NPOXEDGE		—	0
NPRT		—	0
NPRWGD		—	0
NPRWGS		—	0
NPSAT		—	0
NPSATCV		—	0
NPTWG		—	0
NPTWGR		—	0
NPTWGT		—	0
NPVAG		—	0
NQMFACTOR		—	0
NQMTCENCV		—	0
NQMTCENCVA		—	0
NQMTCENIV		—	0
NQSMOD		—	0
NRD	Number of source diffusion squares	—	0
NRDSW		—	0
NRDW		—	0
NRS	Number of source diffusion squares	—	0
NRSW		—	0
NSD	Source/drain active doping concentration in m-3	—	2e+26

Table 2.73: BSIM-CMG FINFET v107.0.0 Device Model Parameters

Parameter	Description	Units	Default
NSDE	Source/drain active doping concentration at $L_{eff}$ edge	—	$2e+25$
NSEG	Number of segments for NQSMOD=3 (3,5 and 10 supported)	—	4
NSII0		—	0
NSII1		—	0
NSII2		—	0
NSIID		—	0
NSTHETASAT		—	0
NTGEN	Thermal Generation Current Parameter	—	1
NTGIDL		—	0
NTII		—	0
NTNOI		—	1
NTOX	Exponent for $T_{ox}$ ratio	—	1
NTSS		—	0
NU0		—	0
NUA		—	0
NUA1		—	0
NUC		—	0
NUC1		—	0
NUCS		—	0
NUCSTE		—	0
NUD		—	0
NUD1		—	0
NUP		—	0
NUTE		—	0
NUTL		—	0
NVSAT		—	0
NVSAT1		—	0
NVSAT1R		—	0
NVSATCV		—	0
NWR		—	0
NXRCRG1		—	0
NXRCRG2		—	0
PA1		—	0

Table 2.73: BSIM-CMG FINFET v107.0.0 Device Model Parameters

Parameter	Description	Units	Default
PA11		—	0
PA2		—	0
PA21		—	0
PAGIDL		—	0
PAGISL		—	0
PAIGBACC		—	0
PAIGBACC1		—	0
PAIGBINV		—	0
PAIGBINV1		—	0
PAIGC		—	0
PAIGC1		—	0
PAIGD		—	0
PAIGD1		—	0
PAIGEN		—	0
PAIGS		—	0
PAIGS1		—	0
PALPHA0		—	0
PALPHA1		—	0
PALPHAII0		—	0
PALPHAII1		—	0
PAT		—	0
PBD	Drain-side bulk junction built-in potential	—	0
PBETA0		—	0
PBETAII0		—	0
PBETAII1		—	0
PBETAII2		—	0
PBGIDL		—	0
PBGISL		—	0
PBIGBACC		—	0
PBIGBINV		—	0
PBIGC		—	0
PBIGD		—	0
PBIGEN		—	0
PBIGS		—	0

Table 2.73: BSIM-CMG FINFET v107.0.0 Device Model Parameters

Parameter	Description	Units	Default
PBS	Source-side bulk junction built-in potential	—	1
PBSWD	Built-in potential for Drain-side sidewall junction capacitance	—	0
PBSWGD	Built-in potential for Drain-side gate sidewall junction capacitance	—	0
PBSWGS	Built-in potential for Source-side gate sidewall junction capacitance	—	0
PBSWS	Built-in potential for Source-side sidewall junction capacitance	—	1
PCDSC		—	0
PCDSCD		—	0
PCDSCDR		—	0
PCFD		—	0
PCFS		—	0
PCGBL		—	0
PCGDL		—	0
PCGIDL		—	0
PCGISL		—	0
PCGSL		—	0
PCIGBACC		—	0
PCIGBINV		—	0
PCIGC		—	0
PCIGD		—	0
PCIGS		—	0
PCIT		—	0
PCKAPPAB		—	0
PCKAPPAD		—	0
PCKAPPAS		—	0
PCLM		—	0.013
PCLMCV	CLM parameter for Short Channel CV	—	0
PCLMG		—	0
PCOVD		—	0
PCOVS		—	0
PDEJ	Drain to substrate PN junction perimeter (BULKMOD=1)	—	0
PDELTAVSAT		—	0

Table 2.73: BSIM-CMG FINFET v107.0.0 Device Model Parameters

Parameter	Description	Units	Default
PDELTAVSATCV		—	0
PDEO	Perimeter of drain to substrate overlap region through oxide	—	0
PDIBL1	DIBL Output Conductance parameter - forward mode	—	1.3
PDIBL1R	DIBL Output Conductance parameter - reverse mode	—	0
PDIBL2	DIBL Output Conductance parameter	—	0.0002
PDROUT		—	0
PDSUB		—	0
PDVT0		—	0
PDVT1		—	0
PDVT1SS		—	0
PDVTB		—	0
PDVTSHIFT		—	0
PEGIDL		—	0
PEGISL		—	0
PEIGBINV		—	0
PEMOBT		—	0
PESATII		—	0
PETAO		—	0
PETAOR		—	0
PETAMOB		—	0
PEU		—	0
PGIDL	parameter for body-bias effect on GIDL	—	0
PGISL	parameter for body-bias effect on GISL	—	1
PHIBE	Body effect voltage parameter, V, after binnig should be from [0.2:1.2]	—	0.7
PHIG	Gate workfunction, eV	—	4.61
PHIGL	Length dependence of Gate workfunction, eV/m	—	0
PHIGN1	NFIN dependence of Gate workfunction	—	0
PHIGN2	NFIN dependence of Gate workfunction	—	100000
PHIN	Nonuniform vertical doping effect on surface potential, V	—	0.05
PIGCD	parameter for Igc partition	—	1
PIGT		—	0

Table 2.73: BSIM-CMG FINFET v107.0.0 Device Model Parameters

Parameter	Description	Units	Default
PIIT		—	0
PK0		—	0
PK01		—	0
PKOSI		—	0
PKOSI1		—	0
PK1		—	0
PK11		—	0
PK1RSCE		—	0
PK1SAT		—	0
PK1SAT1		—	0
PK1SI		—	0
PK1SI1		—	0
PKSATIV		—	0
PKT1		—	0
PLII		—	0
PLPE0		—	0
PLPEB		—	0
PMEXP		—	0
PMEXPR		—	0
PNBODY		—	0
PNGATE		—	0
PNIGBACC		—	0
PNIGBINV		—	0
PNTGEN		—	0
PNTOX		—	0
POXEDGE	Factor for the gate edge Tox	—	1
PPCLM		—	0
PPCLMCV		—	0
PPCLMG		—	0
PPDIBL1		—	0
PPDIBL1R		—	0
PPDIBL2		—	0
PPGIDL		—	0
PPGISL		—	0

Table 2.73: BSIM-CMG FINFET v107.0.0 Device Model Parameters

Parameter	Description	Units	Default
PPHIBE		—	0
PPHIG		—	0
PPHIN		—	0
PPIGCD		—	0
PPOXEDGE		—	0
PPRT		—	0
PPRWGD		—	0
PPRWGS		—	0
PPSAT		—	0
PPSATCV		—	0
PPTWG		—	0
PPTWGR		—	0
PPTWGT		—	0
PPVAG		—	0
PQM	Slope of normalized Tcen in inversion	—	0.66
PQMACC	Slope of normalized Tcen in accumulation	—	0.66
PQMFACTOR		—	0
PQMTCENCV		—	0
PQMTCENCVA		—	0
PQMTCENIV		—	0
PRDDR	Drain side quasi-saturation parameter	—	0
PRDSW		—	0
PRDW		—	0
PRSDEND		—	0
PRSDR	Source side quasi-saturation parameter	—	1
PRSW		—	0
PRT		—	0.001
PRWGD	Gate bias dependence of drain extension resistance	$V^{-1}$	0
PRWGS	Gate bias dependence of source extension resistance	$V^{-1}$	0
PSAT	Velocity saturation exponent, after binnig should be from [2.0:inf)	—	2
PSATCV	Velocity saturation exponent for C-V	—	0

Table 2.73: BSIM-CMG FINFET v107.0.0 Device Model Parameters

Parameter	Description	Units	Default
PSEJ	Source to substrate PN junction perimeter (BULKMOD=1)	—	0
PSEO	Perimeter of source to substrate overlap region through oxide	—	0
PSII0		—	0
PSII1		—	0
PSII2		—	0
PSIID		—	0
PSTTHETASAT		—	0
PTGIDL		—	0
PTII		—	0
PTSS		—	0
PTWG	Gmsat degradation parameter - forward mode	—	0
PTWGR	Gmsat degradation parameter - reverse mode	—	0
PTWGT		—	0.004
PU0		—	0
PUA		—	0
PUA1		—	0
PUC		—	0
PUC1		—	0
PUCS		—	0
PUCSTE		—	0
PUD		—	0
PUD1		—	0
PUP		—	0
PUTE		—	0
PUTL		—	0
PVAG		—	1
PVSAT		—	0
PVSAT1		—	0
PVSAT1R		—	0
PVSATCV		—	0
PWR		—	0
PXRCRG1		—	0
PXRCRG2		—	0

Table 2.73: BSIM-CMG FINFET v107.0.0 Device Model Parameters

Parameter	Description	Units	Default
QMO	Knee-Point for Tcen in inversion (Charge normalized to Cox)	—	0.001
QMOACC	Knee-Point for Tcen in accumulation (Charge normalized to Cox)	—	0.001
QMFACTOR	Prefactor + switch for QM Vth correction	—	0
QMTCENCV	Prefactor + switch for QM Width and Toxeff correction for CV	—	0
QMTCENCVA	Prefactor + switch for QM Width and Toxeff correction for CV (accumulation region)	—	0
QMTCENIV	Prefactor + switch for QM Width correction for IV	—	0
RDDR	Drain side drift resistance parameter - forward mode	—	0
RDDRR	Drain side drift resistance parameter - reverse mode	—	0
RDSMOD	Resistance model selector	—	0
RDSW		—	100
RDSWMIN		—	0
RDW		—	50
RDWMIN		—	0
RGATEMOD	Gate electrode resistor and ge node switcher — NOT USED IN XYCE	—	0
RGEOA	Fitting parameter for RGEOMOD=1	—	1
RGEOB	Fitting parameter for RGEOMOD=1	—	0
RGEOC	Fitting parameter for RGEOMOD=1	—	0
RGEOD	Fitting parameter for RGEOMOD=1	—	0
RGEOE	Fitting parameter for RGEOMOD=1	—	0
RGEOMOD	Bias independent parasitic resistance model selector	—	0
RGEXT	Effective gate electrode external resistance	—	0
RGFIN	Effective gate electrode per finger per fin resistance	—	0.001
RHOC		—	1e-12
RHORSD		—	1
RSDR	Source side drift resistance parameter - forward mode	—	0
RSDRR	Source side drift resistance parameter - reverse mode	—	0
RSHD	Drain-side sheet resistance	—	0

Table 2.73: BSIM-CMG FINFET v107.0.0 Device Model Parameters

Parameter	Description	Units	Default
RSHS	Source-side sheet resistance	—	0
RSW		—	50
RSWMIN		—	0
RTHO	Thermal resistance	—	0.01
SDTERM		—	0
SHMOD	Self heating and T node switcher — NOT USED IN XYCE	—	0
SIIO	Vgs dependent parameter of lii	$V^{-1}$	0.5
SI11	1st Vgs dependent parameter of lii	$V^{-1}$	0.1
SI12	2nd Vgs dependent parameter of lii	—	0
SIID	3rd Vds dependent parameter of lii	$V^{-1}$	0
SJD	Constant for drain-side two-step second junction	—	0
SJS	Constant for source-side two-step second junction	—	0
SJSWD	Constant for drain-side sidewall two-step second junction	—	0
SJSWGD	Constant for source-side gate sidewall two-step second junction	—	0
SJSWGS	Constant for source-side gate sidewall two-step second junction	—	0
SJSWS	Constant for source-side sidewall two-step second junction	—	0
TBGASUB	Bandgap Temperature Coefficient (eV / degrees)	—	0.000702
TBGBSUB	Bandgap Temperature Coefficient (degrees)	—	1108
TCJ	Temperature coefficient for CJS/CJD	—	0
TCJSW	Temperature coefficient for CJSWS/CJSWD	—	0
TCJSWG	Temperature coefficient for CJSWGS/CJSWGD	—	0
TETA0	Temperature dependence of DIBL coefficient, 1/K	—	0
TETAOR	Temperature dependence of Reverse-mode DIBL coefficient, 1/K	—	0
TFIN	Body (Fin) thickness	—	1.5e-08
TGATE	Gate height on top of the hard mask	—	3e-08
TGIDL	GIDL/GISL Temperature Dependence	—	-0.003
TI1	Impact Ionization Temperature Dependence, IIMOD=2	—	0
TMASK	Height of hard mask on top of the fin	—	3e-08

Table 2.73: BSIM-CMG FINFET v107.0.0 Device Model Parameters

Parameter	Description	Units	Default
TMEXP		–	0
TMEXPR		–	0
TNJT	Temperature coefficient for NJTS	–	0
TNJTSD	Temperature coefficient for NJTSD	–	0
TNJTSSW	Temperature coefficient for NJTSSW	–	0
TNJTSSWD	Temperature coefficient for NJTSSWD	–	0
TNJTSSWG	Temperature coefficient for NJTSSWG	–	0
TNJTSSWGD	Temperature coefficient for NJTSSWGD	–	0
TNOM	Temperature at which the model is extracted (degrees)	–	27
TOXG	oxide thickness for gate current model in meters, Introduced in BSIM-CMG106.1.0	m	0
TOXP	physical oxide thickness in meters	–	1.2e-09
TOXREF	Target tox value [m]	–	1.2e-09
TPB	Temperature coefficient for PBS/PBD	–	0
TPBSW	Temperature coefficient for PBSWS/PBSWD	–	0
TPBSWG	Temperature coefficient for PBSWGS/PBSWGD	–	0
TRDDR		–	0
TRSDR		–	0
TSILI	Thickness of the silicide on top of the raised source/drain	–	1e-08
TSS	SSwing Temperature Coefficient (/ degrees)	–	0
U0		–	0.03
UOMULT	Variability in carrier mobility	–	1
UON1	NFIN dependence of U0	–	0
UON2	NFIN dependence of U0	–	100000
UA		–	0.3
UA1		–	0.001032
UC	Body effect for mobility degradation parameter - BULKMOD=1	–	0
UC1		–	5.6e-11
UCS		–	1
UCSTE		–	-0.004775
UD		–	0
UD1		–	0

Table 2.73: BSIM-CMG FINFET v107.0.0 Device Model Parameters

Parameter	Description	Units	Default
UP		–	0
UTE		–	0
UTL		–	-0.0015
VSAT		–	85000
VSAT1	Velocity Saturation parameter for I <sub>on</sub> degradation - forward mode	–	0
VSAT1N1	NFIN dependence of VSAT1	–	0
VSAT1N2	NFIN dependence of VSAT1	–	0
VSAT1R	Velocity Saturation parameter for I <sub>on</sub> degradation - reverse mode	–	0
VSAT1RN1	NFIN dependence of VSAT1R	–	0
VSAT1RN2	NFIN dependence of VSAT1R	–	0
VSATCV	Velocity Saturation parameter for CV	–	0
VSATN1	NFIN dependence of VSAT	–	0
VSATN2	NFIN dependence of VSAT	–	100000
VTSD	Bottom drain junction trap-assisted current voltage dependent parameter	–	0
VTSS	Bottom source junction trap-assisted current voltage dependent parameter	–	10
VTSSWD	Unit length trap-assisted current voltage dependent parameter for sidewall drain junction	–	0
VTSSWGD	Unit length trap-assisted current voltage dependent parameter for gate-edge sidewall drain junction	–	0
VTSSWGS	Unit length trap-assisted current voltage dependent parameter for gate-edge sidewall source junction	–	10
VTSSWS	Unit length trap-assisted current voltage dependent parameter for sidewall source junction	–	10
WR		–	1
WTH0	Width dependence coefficient for R <sub>th</sub> and C <sub>th</sub>	–	0
XJBVD	Fitting parameter for drain diode breakdown current	–	0
XJBVS	Fitting parameter for source diode breakdown current	–	1
XL	L offset for channel length due to mask/etch effect	–	0
XRCRG1		–	12

Table 2.73: BSIM-CMG FINFET v107.0.0 Device Model Parameters

Parameter	Description	Units	Default
XRCRG2		–	1
XTID	Drain junction current temperature exponent	–	0
XTIS	Source junction current temperature exponent	–	3
XTSD	Power dependence of JTSD on temperature	–	0
XTSS	Power dependence of JTSS on temperature	–	0.02
XTSSWD	Power dependence of JTSSWD on temperature	–	0
XTSSWGD	Power dependence of JTSSWGD on temperature	–	0
XTSSWGS	Power dependence of JTSSWGS on temperature	–	0.02
XTSSWS	Power dependence of JTSSWS on temperature	–	0.02

## Level 301 MOSFET Tables (EKV version 3.0.1)

**Xyce** includes the EKV MOSFET model, version 3.0.1 [14][26][27], the EKV3 model. Full documentation for the EKV3 model is available on the **Xyce** internal web site; the documentation for the EKV3 model available there may be freely redistributed. Instance and model parameters for the EKV model are given in tables 2.74 and 2.75.

The EKV3 model is developed by the EKV Team of the Electronics Laboratory-TUC (Technical University of Crete). It is included in **Xyce** under license from Technical University of Crete. The official web site of the EKV model is <http://ekv.epfl.ch/>.

**Due to licensing restrictions, the EKV3 mosfet is not available in open-source versions of Xyce. The license for EKV3 authorizes Sandia National Laboratories only to distribute binary versions of code with EKV3 included.**

Table 2.74: EKV3 MOSFET Device Instance Parameters

Parameter	Description	Units	Default
AD	DRAIN'S AREA	—	0
AS	SOURCE'S AREA	—	0
L	GATE'S LENGTH	—	1e-05
M	NUMBER OF DEVICES IN PARALLEL	—	1
NF	NUMBER OF FINGERS	—	1
PD	DRAIN'S PERIMETER	—	0
PS	SOURCE'S PERIMETER	—	0
SA	STI PARAMETER; DISTANCE FROM STI	—	0
SB	STI PARAMETER; DISTANCE FROM STI	—	0
SD	STI PARAMETER; DISTANCE BETWEEN GATES	—	0
W	GATE'S WIDTH	—	1e-05

Table 2.75: EKV3 MOSFET Device Model Parameters

Parameter	Description	Units	Default
ACLM		—	0.83
AF		—	1
AGAM		—	0
AGAMMA	MATCHING PARAMETER FOR BODY FACTOR (GAMMA)	—	0
AGIDL		—	0
AKP	MATCHING PARAMETER FOR MOBILITY (KP)	—	0
AQMA		—	0.5
AQMI		—	0.4

Table 2.75: EKV3 MOSFET Device Model Parameters

Parameter	Description	Units	Default
AVT		—	0
AVTO	MATCHING PARAMETER FOR THRESHOLD VOLTAGE (VTO)	—	0
BEX		—	-1.5
BGIDL		—	2.3e+09
BVD		—	10
BVS		—	10
CGB0		—	0
CGD0		—	0
CGIDL		—	0.5
CGS0		—	0
CJD		—	0
CJF		—	0
CJS		—	0
CJSWD		—	0
CJSWGD		—	0
CJSWGS		—	0
CJSWS		—	0
COX		—	0.012
DDITS		—	0.3
DELTA		—	2
DFR		—	0.001
DGAMMAEDGE		—	0
DL		—	-1e-08
DLC		—	0
DPHIEDGE		—	0
DW		—	-1e-08
DWC		—	0
E0		—	1e+10
E1		—	3.1e+08
EB		—	2.9e+10
EF		—	2
EGIDL		—	0.8
ETA		—	0.5
ETAD		—	1

Table 2.75: EKV3 MOSFET Device Model Parameters

Parameter	Description	Units	Default
ETAQM		—	0.75
FLR		—	0
FPROUT		—	1e+06
GAMMA		—	0.3
GAMMAG		—	4.1
GAMMAGOV		—	10
GAMMAOV		—	1.6
GC		—	1
GMIN		—	0
HDIF		—	0
IBA		—	0
IBB		—	3e+08
IBBT		—	0.0008
IBN		—	1
INFO_LEVEL		—	0
JSD		—	0
JSS		—	0
JSSWD		—	0
JSSWGD		—	0
JSSWGS		—	0
JSSWS		—	0
JTSD		—	0
JTSS		—	0
JTSSWD		—	0
JTSSWGD		—	0
JTSSWGS		—	0
JTSSWS		—	0
KA		—	0
KB		—	0
KETAD		—	0
KF		—	0
KG		—	0
KGAMMA		—	0
KGFN		—	0
KJF		—	0

Table 2.75: EKV3 MOSFET Device Model Parameters

Parameter	Description	Units	Default
KKP		—	0
KP		—	0.0005
KRGL1		—	0
KUCRIT		—	0
KVTO		—	0
LA		—	1
LAMBDA		—	0.5
LB		—	1
LDIF		—	0
LDPHIEDGE		—	0
LDW		—	0
LETA		—	0.5
LETA0		—	0
LETA2		—	0
LGAM		—	1
LKKP		—	0
LKVTO		—	0
LL		—	0
LLN		—	1
LLODKKP		—	1
LLODKVTO		—	1
LNWR		—	0
LODKETAD		—	1
LODKGAMMA		—	1
LOV		—	2e-08
LOVIG		—	2e-08
LQWR		—	0
LR		—	5e-08
LVT		—	1
LWR		—	0
MJD		—	0.9
MJS		—	0.9
MJSWD		—	0.7
MJSWGD		—	0.7
MJSWGS		—	0.7

Table 2.75: EKV3 MOSFET Device Model Parameters

Parameter	Description	Units	Default
MJSWS		—	0.7
NO		—	1
NCS		—	1
NFVTA		—	0
NFVTB		—	10000
NJD		—	1
NJS		—	1
NJTSD		—	1
NJTSS		—	1
NJTSSWD		—	1
NJTSSWGD		—	1
NJTSSWGS		—	1
NJTSSWS		—	1
NLR		—	0.01
NQS_NOI		—	1
NWR		—	0.005
PBD		—	0.8
PBS		—	0.8
PBSWD		—	0.6
PBSWGD		—	0.6
PBSWGS		—	0.6
PBSWS		—	0.6
PDITS		—	0
PDITSD		—	1
PDITSL		—	0
PHIF	FERMI BULK POTENTIAL	—	0.45
PKKP		—	0
PKVTO		—	0
QLR		—	0.0005
QOFF		—	0
QWR		—	0.0003
RBN		—	0
RBWSH		—	0.003
RD		—	0
RDBN		—	0

Table 2.75: EKV3 MOSFET Device Model Parameters

Parameter	Description	Units	Default
RDBWSH		—	0.001
RDSBSH		—	1000
RDX		—	-1
RGSH		—	3
RINGTYPE		—	1
RLX	EXTERNAL SERIES RESISTANCE	—	-1
RS		—	0
RSBN		—	0
RSBWSH		—	0.001
RSH		—	0
RSX		—	-1
SAREF		—	0
SBREF		—	0
SCALE		—	1
SIGMAD		—	1
SIGN	SIGN = 1 FOR NMOS; SIGN = -1 FOR PMOS	—	1
TCJ		—	0
TCJSW		—	0
TCJSWG		—	0
TCV		—	0.0006
TCVL		—	0
TCVW		—	0
TCVWL		—	0
TEOEX		—	0.5
TE1EX		—	0.5
TETA		—	-0.0009
TG	TYPE OF GATE: -1 ENHANCEMENT TYPE; 1 DEPLETION TYPE	—	-1
TH_NOI		—	0
THC		—	0
TKKP		—	0
TLAMBDA		—	0
TNJTSD		—	0
TNJTSS		—	0
TNJTSSWD		—	0

Table 2.75: EKV3 MOSFET Device Model Parameters

Parameter	Description	Units	Default
TNJTSSWGD		—	0
TNJTSSWGS		—	0
TNJTSSWS		—	0
TNOM		—	27
TPB		—	0
TPBSW		—	0
TPBSWG		—	0
TR		—	0
TR2		—	0
UCEX		—	1.5
UCRIT		—	5e+06
VBI		—	0
VFBOV		—	0
VFR		—	0
VOV		—	1
VTO	THRESHOLD VOLTAGE	—	0.3
VTSD		—	0
VTSS		—	0
VTSSWD		—	0
VTSSWGD		—	0
VTSSWGS		—	0
VTSSWS		—	0
WDL		—	0
WDPHIEDGE		—	0
WEO		—	0
WE1		—	0
WEDGE		—	0
WETA		—	0.2
WETAD		—	0
WGAM		—	1
WKKP		—	0
WKP1		—	1e-06
WKP2		—	0
WKP3		—	1
WKVT0		—	0

Table 2.75: EKV3 MOSFET Device Model Parameters

Parameter	Description	Units	Default
WLAMBDA		—	0
WLDGAMMAEDGE		—	0
WLDPHIEDGE		—	0
WLOD		—	0
WLODKKP		—	1
WLODKVTO		—	1
WLR		—	0
WNLR		—	0
WQLR		—	0
WR		—	9e-08
WRLX		—	0
WUCEX		—	0
WUCRIT		—	0
WVT		—	1
XB		—	3.1
XJ		—	2e-08
XJBVD		—	0
XJBVS		—	0
XL		—	0
XTID		—	3
XTIS		—	3
XTSD		—	0
XTSS		—	0
XTSSWD		—	0
XTSSWGD		—	0
XTSSWGS		—	0
XTSSWS		—	0
XW		—	0
ZC		—	1e-06

## 2.3.18 Lossy Transmission Line (LTRA)

Symbol



**Instance Form**    0<name> <A port (+) node> <A port (-) node>  
+ <B port (+) node> <B port (-) node> [model name]

**Model Form**    .MODEL <model name> LTRA R=<value> L=<value> C=<value>  
+ G=<value> LEN=<value> [model parameters]

**Examples**    0line1 inp inn outp outn cable1  
0line2 inp inn outp outn cable1

**Comments**    The lossy transmission line, or LTRA, device is a two port (A and B), bi-directional device. The (+) and (-) nodes define the polarity of a positive voltage at a port. R, L, C, and G are the resistance, inductance, capacitance, and conductance of the transmission line per unit length, respectively. LEN is the total length of the transmission line. Supported configurations for the LTRA are RLC, RC, LC (lossless) and RG.

### Device Parameters

Table 2.76: Lossy Transmission Line Device Instance Parameters

Parameter	Description	Units	Default
I1	Initial current at end 1	A	0
I2	Initial current at end 2	A	0
V1	Initial voltage at end 1	V	0
V2	Initial voltage at end 2	V	0

### Model Parameters

Table 2.77: Lossy Transmission Line Device Model Parameters

Parameter	Description	Units	Default
ABS	Abs. rate of change of deriv. for bkpt	—	1
C	Capacitance per unit length	F/m	0
COMPACTABS	special abstol for straight line checking	—	1e-12

Table 2.77: Lossy Transmission Line Device Model Parameters

Parameter	Description	Units	Default
COMPACTREL	special reltol for straight line checking	—	0.001
COMPLEXSTEPCONTROL	do complex time step control using local truncation error estimation	logical (T/F)	false
G	Conductance per unit length	$\Omega^{-1} \text{m}^{-1}$	0
L	Inductance per unit length	$\text{Hm}^{-1}$	0
LEN	length of line	m	0
LININTERP	use linear interpolation	logical (T/F)	false
MIXEDINTERP	use linear interpolation if quadratic results look unacceptable	logical (T/F)	false
NOSTEPLIMIT	don't limit timestep size based on the time constant of the line	logical (T/F)	false
QUADINTERP	use quadratic interpolation	logical (T/F)	true
R	Resistance per unit length	$\Omega/\text{m}$	0
REL	Rel. rate of change of deriv. for bkpt	—	1
STEPLIMIT	limit timestep size based on the time constant of the line	logical (T/F)	true
TRUNCDONTCUT	don't limit timestep to keep impulse response calculation errors low	logical (T/F)	false
TRUNCNR	use N-R iterations for step calculation in LTRATrunc	logical (T/F)	false

By default time step limiting is on in the LTRA. This means that simulation step sizes will be reduced if required by the LTRA to preserve accuracy. This can be disabled by setting NOSTEPLIMIT=1 and TRUNCDONTCUT=1 on the .MODEL line.

The option most worth experimenting with for increasing the speed of simulation is REL. The default value of 1 is usually safe from the point of view of accuracy but occasionally increases computation time. A value greater than 2 eliminates all breakpoints and may be worth trying depending on the nature of the rest of the circuit, keeping in mind that it might not be safe from the viewpoint of accuracy. Breakpoints may be entirely eliminated if the circuit does not exhibit any sharp discontinuities. Values between 0 and 1 are usually not required but may be used for setting many breakpoints.

COMPACTREL and COMPACTABS are tolerances that control when the device should attempt to compact past history. This can significantly speed up the simulation, and reduce memory usage, but can negatively impact accuracy and in some cases may cause problems with the nonlinear solver. In general this capability should be used with linear type signals, such as square-wave-like voltages. In order to activate this capability the general device option TRYTOCOMPACT=1 must be set, if

it is not no history compaction will be performed and the COMPACT options will be ignored.

Example:

```
.OPTIONS DEVICE TRYTOCOMPACT=1
```

## References

See references [28] and [29] for more information about the model.

## 2.3.19 Voltage- or Current-controlled Switch

---

<b>Instance Form</b>	<pre>S&lt;name&gt; &lt;(+) switch node&gt; &lt;(-) switch node&gt; + &lt;(+) control node&gt; &lt;(-) control node&gt; + &lt;model name&gt; [ON] [OFF]  W&lt;name&gt; &lt;(+) switch node&gt; &lt;(-) switch node&gt; + &lt;control node voltage source&gt; + &lt;model name&gt; [ON] [OFF]</pre>
----------------------	---

---

<b>Model Form</b>	<pre>.MODEL &lt;model name&gt; VSWITCH [model parameters] .MODEL &lt;model name&gt; ISWITCH [model parameters]</pre>
-------------------	--

---

<b>Examples</b>	<pre>S1 21 23 12 10 SMOD1 SSET 15 10 1 13 SRELAY W1 1 2 VCLOCK SWITCHMOD1 W2 3 0 VRAMP SM1 ON</pre>
-----------------	---

---

<b>Comments</b>	<p>The voltage- or current-controlled switch is a particular type of controlled resistor. This model is designed to help reduce numerical issues. See Special considerations below.</p>
-----------------	---

The resistance between the <(+) switch node> and the <(-) switch node> is dependent on either the voltage between the <(+) control node> and the <(-) control node> or the current through the control node voltage source. The resistance changes in a continuous manner between the RON and ROFF model parameters.

No resistance is inserted between the control nodes. It is up to the user to make sure that these nodes are not floating.

Even though evaluating the switch model is computationally inexpensive, for transient analysis **Xyce** steps through the transition section using small time-steps in order to calculate the waveform accurately. Thus, a circuit with many switch transitions can result in lengthy run times.

The ON and OFF parameters are used to specify the initial state of the switch at the first step of the operating point calculation; this does not force the switch to be in that state, it only gives the operating point solver an initial state to work with. If it is known that the switch should be in a particular state in the operating point it could help convergence to specify one of these keywords.

## Model Parameters

Table 2.78: Controlled Switch Device Model Parameters

Parameter	Description	Units	Default
IOFF	Off current	A	0
ION	On current	A	0.001
OFF	Off control value	–	0
ON	On control value	–	1
ROFF	Off resistance	$\Omega$	1e+06
RON	On resistance	$\Omega$	1
VOFF	Off voltage	V	0
VON	On voltage	V	1

## Special Considerations

- Due to numerical limitations, **Xyce** can only manage a dynamic range of approximately 12 decades. Thus, it is recommended the user limit the ratio **ROFF/RON** to less than  $10^{12}$ .
- Furthermore, it is a good idea to limit the narrowness of the transition region. This is because in the transition region, the switch has gain and the narrower the region, the higher the gain and the more potential for numerical problems. The smallest value allowed for  $\|\mathbf{VON} - \mathbf{VOFF}\|$  or  $\|\mathbf{ION} - \mathbf{IOFF}\|$  is  $1 \times 10^{-12}$ .

## Controlled switch equations

The equations in this section use the following variables:

$R_s$	=	switch resistance	
$V_c$	=	voltage across control nodes	
$I_c$	=	current through control node voltage source	
$L_m$	=	log-mean of resistor values	$= \ln(\sqrt{\mathbf{RON} \cdot \mathbf{ROFF}})$
$L_r$	=	log - ratio of resistor values	$= \ln(\mathbf{RON}/\mathbf{ROFF})$
$V_d$	=	difference of control voltages	$= \mathbf{VON} - \mathbf{VOFF}$
$I_d$	=	difference of control currents	$= \mathbf{ION} - \mathbf{IOFF}$

### Switch Resistance

To compute the switch resistance, **Xyce** first calculates the “switch state”  $S$  as  $S = (V_c - \mathbf{VOFF})/V_d$  or  $S = (I_c - \mathbf{IOFF})/I_d$ . The switch resistance is then:

$$R_s = \begin{cases} \mathbf{RON}, & S \geq 1.0 \\ \mathbf{ROFF}, & S \leq 0.0 \\ \exp(L_m + 0.75L_r(2S - 1) - 0.25L_r(2S - 1)^3), & 0 < S < 1 \end{cases}$$

## 2.3.20 Generic Switch

---

<b>Instance Form</b>	SW<name> <(+) switch node> <(-) switch node> <model name> [ON] [OFF] <control = expression >
----------------------	---

---

<b>Model Form</b>	.MODEL <model name> VSWITCH [model parameters] .MODEL <model name> ISWITCH [model parameters] .MODEL <model name> SWITCH [model parameters]
-------------------	---

---

<b>Examples</b>	SW 1 2 SWI OFF CONTROL={I(VMON)} SW 1 2 SWV OFF CONTROL={V(3)-V(4)} SW 1 2 SW OFF CONTROL={if(time>0.001,1,0)}
-----------------	--

---

<b>Comments</b>	The generic switch is similar to the voltage- or current-controlled switch except that the control variable is anything that can be written as an expression. The examples show how a voltage- or current-controlled switch can be implemented with the generic switch. Also shown is a relay that turns on when a certain time is reached. Model parameters are given in Table 2.78.
-----------------	---

## 2.3.21 Lossless (Ideal) Transmission Line

Symbol



**Instance Form**    T<name> <A port (+) node> <A port (-) node>  
                         + <B port (+) node> <B port (-) node>  
                         + Z0=<value> [TD=<value>] [F=<value> [NL=<value>]]

**Examples**            Tline inp inn outp outn Z0=50 TD=1us  
                         Tline2 inp inn outp outn Z0=50 F=1meg NL=1.0

**Comments**            The lossless transmission line device is a two port (A and B), bi-directional delay line. The (+) and (-) nodes define the polarity of a positive voltage at a port.

                             Z0 is the characteristic impedance. The transmission line's length is specified by either TD (a delay in seconds) or by F and NL (a frequency and relative wavelength at F). NL defaults to 0.25 (F is the quarter-wave frequency). If F is given, the time delay is computed as  $\frac{NL}{F}$ . While both TD and F are optional, at least one of them must be given.

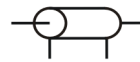
### Instance Parameters

Table 2.79: Ideal Transmission Line Device Instance Parameters

Parameter	Description	Units	Default
F	Frequency	Hz	0
NL	Length in wavelengths	—	0
TD	Time delay	s	0
Z0	Characteristic Impedance	$\Omega$	0
Z0	Characteristic Impedance	$\Omega$	0

## 2.3.22 Lumped Transmission Line

Symbol



**Instance Form**     `ytransline <name> <Input port> <Output port> testLine`  
                         `+ len=<value> lumps=<value>`

**Model Form**        `.model testLine transline r=<value> l=<value>`  
                         `+ c=<value> [model parameters]`

**Examples**           `ytransline line1 inn out testLine len=12.0 lumps=1440`

**Comments**         The lumped transmission line, device is a two port bi-directional device. The specification is patterned, loosely, from the netlist specification for the LTRA device.

R, L, and C are the resistance, inductance, and capacitance of the transmission line per unit length, respectively. LEN is the total length of the transmission line, and LUMPS is the number of lumped elements used to discretize the line. Supported configurations for this device are RLC and LC.

Unlike the LTRA device, which is based on an analytic solution, this device is based on assembling chains of linear R,L and C devices to approximate the solution to the Telegraph equations. It is the functional equivalent of building a transmission line in the netlist using subcircuits of linear elements. The advantage of using this approach is that it automates the mechanics of this process, and thus is less prone to error. It can be used with all analysis types, including harmonic balance (HB).

The model is based on the assumption that the segments of the line are evenly spaced. The number of segments is specified by the parameter LUMPS and the larger this number, the more accurate the calculation.

### Device Parameters

Table 2.80: Lumped Transmission Line Device Instance Parameters

Parameter	Description	Units	Default
LEN	length of line	m	0
LUMPS		—	1

### Model Parameters

Table 2.81: Lumped Transmission Line Device Model Parameters

Parameter	Description	Units	Default
C	Capacitance per unit length	F/m	0
L	Inductance per unit length	Hm <sup>-1</sup>	0
R	Resistance per unit length	Ω/m	0

## 2.3.23 Behavioral Digital Devices

---

**Instance Form**     U<name> <type>(<num inputs>) [digital power node]  
                      + [digital ground node] <input node>\* <output node>\*  
                      + <model name> [device parameters]

---

**Model Form**        .MODEL <model name> DIG [model parameters]

---

**Examples**           UMYAND AND(2) DPWR DGND in1 in2 out DMOD IC=TRUE  
                      UTHEINV INV DPWR DGND in out DMOD  
                      .model DMOD DIG (  
                      + CLO=1e-12 CHI=1e-12  
                      + SORLO=5 SORHI=5 SOTSW=5e-9  
                      + SOVLO=-1 SOVHI=1.8  
                      + S1RLO=200 S1RHI=5 S1TSW=5e-9  
                      + S1VLO=1 S1VHI=3  
                      + RLOAD=1000  
                      + CLOAD=1e-12  
                      + DELAY=20ns )

---

### Parameters and Options

type

Type of digital device. Supported devices are: INV, BUF, AND, NAND, OR, NOR, XOR, NXOR, DFF, DLTCH and ADD. (Note: NOT is an allowed synonym for INV, but will be deprecated in future **Xyce** releases.)

The following gates have a fixed number of inputs. INV and BUF have only one input and one output node. XOR and NXOR have two inputs and one output. ADD has three inputs (in1, in2, carryIn) and two outputs (sumOut and carryOut). DFF has four inputs (PREB, CLRB, Clock and Data) and two outputs ( $Q$  and  $\bar{Q}$ ). DLTCH has four inputs (PREB, CLRB, Enable and Data) and two outputs ( $Q$  and  $\bar{Q}$ ).

The AND, NAND, OR and NOR gates have one output but a variable number of inputs. There is no limit on the number of inputs for AND, NAND, OR and NOR gates, but there must be at least two inputs.

num inputs

For AND, NAND, OR and NOR gates, with N inputs, the syntax is (N), as shown for the MYAND example given above, where AND(2) is specified. The inclusion of (N) is mandatory for gates with a variable number of inputs, and both the left and right parentheses must be used to enclose N.

This parameter is optional, and typically omitted, for gates with a fixed number of inputs, such as INV, BUF, XOR, NXOR, DFF, DLTCH and

ADD. This is illustrated by the THEINV example given above, where the device type is INV rather than INV(1).

**digital power node**

Dominant node to be connected to the output node(s) to establish high output state. This node is connected to the output by a resistor and capacitor in parallel, whose values are set by the model. This node must be specified on the instance line.

**digital ground node**

This node serves two purposes, and must be specified on the instance line. It is the dominant node to be connected to the output node(s) to establish low output state. This node is connected to the output by a resistor and capacitor in parallel, whose values are set by the model. This node is also connected to the input node by a resistor and capacitor in parallel, whose values are set by the model. Determination of the input state is based on the voltage drop between the input node and this node.

**input nodes, output nodes**

Input and output nodes that connect to the circuit.

**model name**

Name of the model defined in a .MODEL line.

**device parameters**

Parameter listed in Table 2.82 may be provided as <parameter>=<value> specifications as needed. For devices with more than one output, multiple output initial states may be provided as Boolean values in either a comma separated list (e.g. IC=TRUE,FALSE for a device with two outputs) or individually (e.g. IC1=TRUE IC2=FALSE or IC2=FALSE). Finally, the IC specification must use TRUE and FALSE rather than T and F.

## Device Parameters

Table 2.82: Behavioral Digital Device Instance Parameters

Parameter	Description	Units	Default
IC1	Vector of initial values for output(s)	logical (T/F)	false
IC2		—	false

## Model Parameters

Table 2.83: Behavioral Digital Device Model Parameters

Parameter	Description	Units	Default
CHI	Capacitance between output node and high reference	F	1e-06
CLO	Capacitance between output node and low reference	F	1e-06
CLOAD	Capacitance between input node and input reference	F	1e-06
DELAY	Delay time of device	s	1e-08
RLOAD	Resistance between input node and input reference	$\Omega$	1000
SORHI	Low state resistance between output node and high reference	$\Omega$	100
SORLO	Low state resistance between output node and low reference	$\Omega$	100
SOTSW	Switching time transition to low state	s	1e-08
S0VHI	Maximum voltage to switch to low state	V	1.7
S0VLO	Minimum voltage to switch to low state	V	-1.5
S1RHI	High state resistance between output node and high reference	$\Omega$	100
S1RLO	High state resistance between output node and low reference	$\Omega$	100
S1TSW	Switching time transition to high state	s	1e-08
S1VHI	Maximum voltage to switch to high state	V	7
S1VLO	Minimum voltage to switch to high state	V	0.9

## Model Description

The input interface model consists of the input node connected with a resistor and capacitor in parallel to the digital ground node. The values of these are: **RLOAD** and **CLOAD**.

The logical state of any input node is determined by comparing the voltage relative to the reference to the range for the low and high state. The range for the low state is **S0VLO** to **S0VHI**. Similarly, the range for the high state is **S1VLO** to **S1VHI**. The state of an input node will remain fixed as long as its voltage stays within the range for its current state. That input node will transition to the other state only when its state goes outside the voltage range of its current state.

The output interface model is more complex than the input model, but shares the same basic configuration of a resistor and capacitor in parallel to simulate loading. For the output case, there are such parallel RC connections to two nodes, the digital ground node and the digital power node. Both of these nodes must be specified on the instance line.

The capacitance to the high node is specified by **CHI**, and the capacitance to the low node is **CLO**. The resistors in parallel with these capacitors are variable, and have values that depend

on the state. In the low state (S0), the resistance values are: **S0RLO** and **S0RHI**. In the high state (S1), the resistance values are: **S1RLO** and **S1RHI**. Transition to the high state occurs exponentially over a time of **S1TSW**, and to the low state **S0TSW**.

The device's delay is given by the model parameter **DELAY**. Any input changes that affect the device's outputs are propagated after this delay.

As a note, the model parameters **VREF**, **VLO** and **VHI** are used by the now deprecated Y-type digital device, but are ignored by the U device. A warning message is emitted if any of these three parameters are used in the model card for a U device.

The DCOP Calculations for flip-flops and latches may differ between **Xyce** and PSpice. As an example, the **Xyce** DLTCH model enforces  $Q$  and  $\bar{Q}$  being different at DCOP. This behavior differs from PSpice in some cases (such as when Enable is low at DCOP). In that case, PSpice would have  $Q$  and  $\bar{Q}$  in an indeterminate state that is about halfway between **V\_L0** and **V\_HI**.

Another caveat is that closely spaced input transitions to the **Xyce** digital behavioral models may not be accurately reflected in the output states. In particular, input-state changes spaced by more than **DELAY** seconds have independent effects on the output states. However, two input-state changes (S1 and S2) that occur within **DELAY** seconds (e.g., at time= $t_1$  and time= $t_1+0.5*\text{DELAY}$ ) have the effect of masking the effects of S1 on the device's output states, and only the effects of S2 are propagated to the device's output states.

## 2.3.24 Y-Type Behavioral Digital Devices (Deprecated)

---

**Instance Form**    Y<type> <name> [low output node] [high output node]  
+ [input reference node] <input node>\* <output node>\*  
+ <model name> [device parameters]

---

**Model Form**    .MODEL <model name> DIG [model parameters]

---

**Examples**

```
YAND MYAND in1 in2 out DMOD IC=TRUE
YNOT THENOT in out DMOD
YNOR ANOR2 vlo vhi vref in1 in2 out DDEF
.model DMOD DIG (
+ CLO=1e-12 CHI=1e-12
+ SORLO=5 SORHI=5 SOTSW=5e-9
+ SOVLO=-1 SOVHI=1.8
+ S1RLO=200 S1RHI=5 S1TSW=5e-9
+ S1VLO=1 S1VHI=3
+ RLOAD=1000
+ CLOAD=1e-12
+ VREF=0 VLO=0 VHI=3
+ DELAY=20ns )
.MODEL DDEF DIG
```

---

### Parameters and Options

type

Type of digital device. Supported devices are: NOT, BUF, AND, NAND, OR, NOR, XOR, NXOR, DFF, DLTCH and ADD. (Note: INV is now the preferred synonym for NOT. The NOT device type will be deprecated in future **Xyce** releases.) For Y-type digital devices, all devices have two input nodes and one output node, except for NOT, DFF and ADD. NOT has one input and one output. ADD has three inputs (in1, in2, carryIn) and two outputs (sumOut and carryOut). DFF has four inputs (PREB, CLRB, Enable and Data) and two outputs ( $Q$  and  $\bar{Q}$ ). DLTCH has four inputs (PREB, CLRB, Enable and Data) and two outputs ( $Q$  and  $\bar{Q}$ ).

name

Name of the device instance. This must be present, and when combined with the Y<type>, must be unique in the netlist. In the examples, MYAND, THENOT and ANOR2 have been used as names for the three devices.

low output node

Dominant node to be connected to the output node(s) to establish low output state. This node is connected to the output by a resistor and

capacitor in parallel, whose values are set by the model. If specified by the model, this node must be omitted from the instance line and a fixed voltage **VLO** is used instead.

#### high output node

Dominant node to be connected to the output node(s) to establish high output state. This node is connected to the output by a resistor and capacitor in parallel, whose values are set by the model. If specified by the model, this node must be omitted from the instance line and a fixed voltage **VHI** is used instead.

#### input reference node

This node is connected to the input node by a resistor and capacitor in parallel, whose values are set by the model. Determination if the input state is based on the voltage drop between the input node and this node. If specified by the model, this node must be omitted from the instance line and a fixed voltage **VREF** is used instead.

#### input nodes, output nodes

Nodes that connect to the circuit.

#### model name

Name of the model defined in a .MODEL line.

#### device parameters

Parameter listed in Table 2.84 may be provided as <parameter>=<value> specifications as needed. For devices with more than one output, multiple output initial states may be provided as Boolean values in either a comma separated list (e.g. IC=TRUE,FALSE for a device with two outputs) or individually (e.g. IC1=TRUE IC2=FALSE or IC2=FALSE). Finally, the IC specification must use TRUE and FALSE rather than T and F.

## Device Parameters

Table 2.84: Behavioral Digital Device Instance Parameters

Parameter	Description	Units	Default
IC1	Vector of initial values for output(s)	logical (T/F)	false
IC2		–	false

## Model Parameters

Table 2.85: Behavioral Digital Device Model Parameters

Parameter	Description	Units	Default
CHI	Capacitance between output node and high reference	F	1e-06

Table 2.85: Behavioral Digital Device Model Parameters

Parameter	Description	Units	Default
CLO	Capacitance between output node and low reference	F	1e-06
CLOAD	Capacitance between input node and input reference	F	1e-06
DELAY	Delay time of device	s	1e-08
RLOAD	Resistance between input node and input reference	$\Omega$	1000
SORHI	Low state resistance between output node and high reference	$\Omega$	100
SORLO	Low state resistance between output node and low reference	$\Omega$	100
SOTSW	Switching time transition to low state	s	1e-08
SOVHI	Maximum voltage to switch to low state	V	1.7
SOVLO	Minimum voltage to switch to low state	V	-1.5
S1RHI	High state resistance between output node and high reference	$\Omega$	100
S1RLO	High state resistance between output node and low reference	$\Omega$	100
S1TSW	Switching time transition to high state	s	1e-08
S1VHI	Maximum voltage to switch to high state	V	7
S1VLO	Minimum voltage to switch to high state	V	0.9
VHI	Internal high state supply voltage	V	0
VLO	Internal low state supply voltage	V	0
VREF	Internal reference voltage for inputs	V	0

## Model Description

The input interface model consists of the input node connected with a resistor and capacitor in parallel to the digital ground node. The values of these are: **RLOAD** and **CLOAD**.

The logical state of any input node is determined by comparing the voltage relative to the reference to the range for the low and high state. The range for the low state is **S0VLO** to **S0VHI**. Similarly, the range for the high state is **S1VLO** to **S1VHI**. The state of an input node will remain fixed as long as its voltage stays within the voltage range for its current state. That input node will transition to the other state only when its state goes outside the range of its current state.

The output interface model is more complex than the input model, but shares the same basic configuration of a resistor and capacitor in parallel to simulate loading. For the output case, there are such connections to two nodes, the digital ground node and the digital power node. Both of these nodes must be specified on the instance line.

The capacitance to the high node is specified by **CHI**, and the capacitance to the low node is **CLO**. The resistors in parallel with these capacitors are variable, and have values that depend on the state. In the low state (S0), the resistance values are: **S0RLO** and **S0RHI**. In the high state (S1), the resistance values are: **S1RLO** and **S1RHI**. Transition to the high state occurs exponentially over a time of **S1TSW**, and to the low state **S0TSW**.

The device's delay is given by the model parameter **DELAY**. Any input changes that affect the device's outputs are propagated after this delay.

The DCOP Calculations for flip-flops and latches may differ between **Xyce** and PSpice. As an example, the **Xyce** DLTCH model enforces  $Q$  and  $\bar{Q}$  being different at DCOP. This behavior differs from PSpice in some cases (such as when Enable is low at DCOP). In that case, PSpice would have  $Q$  and  $\bar{Q}$  in an indeterminate state that is about halfway between **V<sub>LO</sub>** and **V<sub>HI</sub>**.

Another caveat is that closely spaced input transitions to the **Xyce** digital behavioral models may not be accurately reflected in the output states. In particular, input-state changes spaced by more than **DELAY** seconds have independent effects on the output states. However, two input-state changes (S1 and S2) that occur within **DELAY** seconds (e.g., at time=t1 and time=t1+0.5\***DELAY**) have the effect of masking the effects of S1 on the device's output states, and only the effects of S2 are propagated to the device's output states.

## Converting Y-Type Digital Devices to U-Type Digital Devices

**Xyce** is migrating the digital behavioral devices to U devices. The goal is increased compatibility with PSpice netlists. This subsection gives four examples of how to convert an existing **Xyce** netlist using Y-type digital devices to the corresponding U device syntaxes. The conversion process depends on whether the device has a fixed number of inputs or a variable number of inputs. In all cases, the the model parameters **VREF**, **VLO** and **VHI** should be omitted from the U device model card. For U devices, the nodes **vlo** and **vhi** are always specified on the instance line.

Example 1: Fixed number of inputs, Y-device model card contains **VREF**, **VLO** and **VHI**. Assume **VREF=VLO**.

```
YNOT THENOT in out DMOD
.model DMOD DIG (
+ CLO=1e-12 CHI=1e-12
+ SORLO=5 SORHI=5 S0TSW=5e-9
+ SOVLO=-1 SOVHI=1.8
+ S1RLO=200 S1RHI=5 S1TSW=5e-9
+ S1VLO=1 S1VHI=3
+ RLOAD=1000
+ CLOAD=1e-12
+ VREF=0 VLO=0 VHI=3
+ DELAY=20ns )

* Digital power node. Assume digital ground node = GND
V1 DPWR 0 3V
UTHENOT INV DPWR 0 in out DMOD1
```

```
.model DMOD1 DIG (
+ CLO=1e-12 CHI=1e-12
+ SORLO=5 SORHI=5 SOTSW=5e-9
+ SOVLO=-1 SOVHI=1.8
+ S1RLO=200 S1RHI=5 S1TSW=5e-9
+ S1VLO=1 S1VHI=3
+ RLOAD=1000
+ CLOAD=1e-12
+ DELAY=20ns )
```

Example 2: Fixed number of inputs, Y-device instance line contains **vlo**, **vhi** and **vref**. Assume **vref=vlo**.

```
YNOT THENOT vlo vhi vref in out DMOD1
UTHENOT INV vhi vlo in out DMOD1
```

Example 3: Variable number of inputs, Y-device model card contains **VREF**, **VLO** and **VHI**. Assume **VREF=VLO**.

```
YAND MYAND in1 in2 out DMOD
UMYAND AND(2) DPWR 0 in1 in2 out DMOD1
```

Example 4: Variable number of inputs, Y-device instance line contains **vlo**, **vhi** and **vref**. Assume **vref=vlo**.

```
YAND MYAND vlo vhi vref in1 in2 out DMOD1
UMYAND AND(2) vhi vlo in1 in2 out DMOD1
```

## 2.3.25 Accelerated mass

Simulation of electromechanical devices or magnetically driven machines may require that **Xyce** simulate the movement of an accelerated mass, that is, to solve the second order initial value problem

$$\begin{aligned}\frac{d^2x}{dt^2} &= a(t) \\ x(0) &= x_0 \\ \dot{x}_0 &= v_0\end{aligned}$$

where  $x$  is the position of the object,  $\dot{x}$  its velocity, and  $a(t)$  the acceleration. In **Xyce**, this simulation capability is provided by the accelerated mass device.

---

<b>Instance Form</b>	YACC <name> <acceleration node> <velocity node> <position node> + [v0=<initial velocity>] [x0=<initial position>]
----------------------	--

---

<b>Examples</b>	<pre>* Simulate a projectile thrown upward against gravity V1 acc 0 -9.8 R1 acc 0 1 YACC acc1 acc vel pos v0=10 x0=0 .print tran v(pos) .tran 1u 10s .end  * Simulate a damped, forced harmonic oscillator * assuming K, c, mass, amplitude and frequency * are defined in .param statements B1 acc 0 V={(-K * v(pos) - c*v(vel))/mass +          + amplitude*sin(frequency*TIME)} R1 acc 0 1 YACC acc2 acc vel pos v0=0 x0=0.4 .print tran v(pos) .tran 1u 10s .end</pre>
-----------------	--

---

<b>Comments</b>	When used as in the examples, <b>Xyce</b> will emit warning messages about the <code>pos</code> and <code>vel</code> nodes not having a DC path to ground. This is normal and should be ignored. The position and velocity nodes should not be connected to any real circuit elements. Their values may, however, be used in behavioral sources; this is done in the second example.
-----------------	--

## 2.3.26 Subcircuit

A subcircuit can be introduced into the circuit netlist using the specified nodes to substitute for the argument nodes in the definition. It provides a building block of circuitry to be defined a single time and subsequently used multiple times in the overall circuit netlists. See Section 2.1.25 for more information about subcircuits.

---

**Instance Form**    X<name> [node]\* <subcircuit name> [PARAMS: [<name> = <value>]\*]

---

**Examples**

```
X12 100 101 200 201 DIFFAMP
XBUFF 13 15 UNITAMP
XFOLLOW IN OUT VCC VEE OUT OPAMP
XFELT 1 2 FILTER PARAMS: CENTER=200kHz
XNANDI 25 28 7 MYPWR MYGND PARAMS: IO_LEVEL=2
```

---

### Parameters and Options

subcircuit name

The name of the subcircuit's definition.

PARAMS:

Passed into subcircuits as arguments and into expressions inside the subcircuit.

---

### Comments

There must be an equal number of nodes in the subcircuit call and in its definition.

Subcircuit references may be nested to any level. However, the nesting cannot be circular. For example, if subcircuit A's definition includes a call to subcircuit B, then subcircuit B's definition cannot include a call to subcircuit A.

## 2.4 TCAD Devices

Semiconductor device simulation, which is based on a coupled set of partial differential equations (PDE's) is supported in **Xyce**. Such devices can be invoked from the circuit netlist, in a similar manner to traditional SPICE-style analog devices. One dimensional and two dimensional devices are supported, with the dimensionality determined by the device model level.

**1D Device Form**    YPDE <name> <node> [node] [model name]  
                      + [na=<value>] [nd=<value>]  
                      + [nx=<value>] [area=<value>]  
                      + [graded=<value>]  
                      + [wj=<value>] [l=<value>] [w=<value>]  
                      + [tecplotlevel=<value>]  
                      + [gnuplotlevel=<value>]  
                      + [node=<tabular data>]  
                      + [region=<tabular data>]  
                      + [bulkmaterial=<string>]  
                      + [temp=<value>]

---

**2D Device Form**    YPDE <name> <node> <node> [node] [node] [model name] |  
                      + [na=<value>] [nd=<value>]  
                      + [meshfile=<filename.msh>]  
                      + [nx=<value>] [ny=<value>]  
                      + [l=<value>] [w=<value>]  
                      + [node=<tabular data>]  
                      + [region=<tabular data>]  
                      + [x0=<value>] [cyl=<value>]  
                      + [tecplotlevel=<value>]  
                      + [gnuplotlevel=<value>] [txtdatalevel=<value>]  
                      + [bulkmaterial=<string>]  
                      + [temp=<value>]

---

**Model Form**        .MODEL <model name> ZOD [model parameters]

---

**Comments**        Most of the PDE parameters are specified on the instance level. At this point the model statement is only used for specifying if the device is 1D or 2D, via the level parameter. Both the 1D and the 2D devices can construct evenly spaced meshes, internally. The 2D device also has the option of reading in an unstructured mesh from an external mesh file, but this is currently an alpha-level capability.

The electrode tabular data specification is explained in detail in table 2.91 Similarly, the doping region tabular data specification is explained in detail in table 2.89.

## TCAD Device Parameters

Most TCAD device parameters are specified on the instance level. There is only one TCAD device model parameter, the level.

Table 2.86: 1D PDE (level 1) Device Instance Parameters

Parameter	Description	Units	Default
AUGER	Flag to turn on/off Auger recombination	logical (T/F)	true
BULKMATERIAL	Bulk semiconductor material	–	'SI'
FIELDDEP	If true, use field dependent mobility.	logical (T/F)	false
MOBMODEL	Mobility model.	–	'ARORA'
NX	Number of mesh points	–	11
SRH	Flag to turn on/off Shockley-Read-Hall recombination.	logical (T/F)	true
<b><i>Doping Parameters</i></b>			
DOPING_FILE	File containing doping profile.	–	'NOFILE'
GRADED	Flag for graded junction vs. abrupt junction. (1/true=graded, 0/false=abrupt)	logical (T/F)	false
NA	Acceptor doping level	cm <sup>-3</sup>	1e+15
ND	Donor doping level	cm <sup>-3</sup>	1e+15
WJ	Junction width, if graded junction enabled.	cm	0.0001
<b><i>Geometry Parameters</i></b>			
ANODE.AREA	Anode area (used for two-terminal devices)	cm <sup>-2</sup>	0
AREA	Cross sectional area of the device.	cm <sup>-2</sup>	1
BASE.AREA	Base area (used for three-terminal (BJT) devices)	cm <sup>-2</sup>	0
BASE.LOC	Location of base contact (necessary if running with three terminals).	cm	0.0005
CATHODE.AREA	Cathode area (used for two-terminal devices)	cm <sup>-2</sup>	0
COLLECTOR.AREA	Collector area (used for three-terminal (BJT) devices)	cm <sup>-2</sup>	0
EMITTER.AREA	Emitter area (used for three-terminal (BJT) devices)	cm <sup>-2</sup>	0
L	Device width.	cm	0.001
W	Device width.	cm	0.001
<b><i>Temperature Parameters</i></b>			
TEMP	Device temperature	°C	27
<b><i>Model Output Parameters</i></b>			

Table 2.86: 1D PDE (level 1) Device Instance Parameters

Parameter	Description	Units	Default
FIRSTELECTRODEOFFSET	This is an output parameter. It is only used if OFFSETOUTPUTVOLTAGE=true. (see description of that parameter)	logical (T/F)	false
GNUPLOTLEVEL	Flag for gnuplot output. 0 - no gnuplot files. 1 - gnuplot files. gnuplot is an open source plotting program that is usually installed on Linux systems. gnuplot files will have the *Gnu.dat suffix, and the prefix will be the name of the device instance.	–	1
OFFSETOUTPUTVOLTAGE	This is an output parameter that determines the “zero” of the potential at output. If OFFSETOUTPUTVOLTAGE=true (default) it will adjust the voltages at output so that the minimum voltage is zero. If true and also FIRSTELECTRODEOFFSET=true, then the voltage of the first electrode is the zero point. If OFFSETOUTPUTVOLTAGE=false, the output voltage sets the intrinsic Fermi level to zero. Depending on circumstances each of these may be more or less convenient for plotting.	logical (T/F)	true
OUTPUTINTERVAL	Time interval for tecplot output (if tecplot is enabled).	s	0
OUTPUTNLPOISSON	Flag to determine if the results of the nonlinear Poisson calculation is included in the output files. Normally, this calculation is used to initialize a drift-diffusion calculation and isn't of interest.	logical (T/F)	false
TECLOTLEVEL	Setting for Tecplot output: 0 - no Tecplot files 1 - Tecplot files, each output in a separate file. 2 - Tecplot file, each output appended to a single file. Tecplot files will have the .dat suffix, and the prefix will be the name of the device instance	–	1
<b>Scaling Parameters</b>			
C0	Density scalar; adjust to mitigate convergence problems. The model will do all of its scaling automatically, so it is generally not necessary to specify it manually.	cm <sup>-3</sup>	1e+15
DENSITYSCALARFRACTION	Fraction of the maximum doping by which density will be scaled. The model will do all of its scaling automatically, so it is generally not necessary to specify it manually.	logical (T/F)	0.1
SCALEDENSITYTOMAXDOPING	If set the density will be scaled by a fraction of the maximum doping. The model will do all of its scaling automatically, so it is generally not necessary to specify it manually.	logical (T/F)	true

Table 2.86: 1D PDE (level 1) Device Instance Parameters

Parameter	Description	Units	Default
t0	Time scalar; adjust to mitigate convergence problems. The model will do all of its scaling automatically, so it is generally not necessary to specify it manually.	s	1e-06
X0	Length scalar; adjust to mitigate convergence problems. The model will do all of its scaling automatically, so it is generally not necessary to specify it manually.	cm	1e-07

Table 2.87: 2D PDE (level 2) Device Instance Parameters

Parameter	Description	Units	Default
BULKMATERIAL	Material of bulk material.	–	'SI'
MESHFILE	This is a required field for a 2D simulation. If the user specifies meshfile=internal.mesh, the model will create a Cartesian mesh using the parameters L,W,NX and NY. If the user specifies anything else (for example meshfile=diode.msh), the model will attempt to read in a mesh file of that name. The format is assumed to be that of the SG Framework.	–	'internal.msh'
MOBMODEL	Mobility model.	–	'ARORA'
NX	Number of mesh points, x-direction.	–	11
NY	Number of mesh points, y-direction.	–	11
<b><i>Doping Parameters</i></b>			
GRADED	Flag for graded junction vs. abrupt junction. (1/true=graded, 0/false=abrupt)	logical (T/F)	false
NA	Acceptor doping level	cm <sup>-3</sup>	1e+15
ND	Donor doping level	cm <sup>-3</sup>	1e+15
WJ	Junction width, if graded junction enabled.	cm	0.0001
<b><i>Geometry Parameters</i></b>			
AREA	Cross sectional area of the device.	cm <sup>-2</sup>	1
CYL	Flag to enable cylindrical geometry	logical (T/F)	false
L	Device length	cm	0.001
W	Device width	cm	0.001
<b><i>Temperature Parameters</i></b>			
TEMP	Device temperature	°C	27
<b><i>Model Output Parameters</i></b>			

Table 2.87: 2D PDE (level 2) Device Instance Parameters

Parameter	Description	Units	Default
GNUPLOTLEVEL	Flag for gnuplot output. 0 - no gnuplot files. 1 - gnuplot files. gnuplot is an open source plotting program that is usually installed on Linux systems. gnuplot files will have the *Gnu.dat suffix, and the prefix will be the name of the device instance.	—	0
OUTPUTINTERVAL	Time interval for tecplot output (if tecplot is enabled).	s	0
OUTPUTNLPOISSON	Flag to determine if the results of the nonlinear Poisson calculation is included in the output files. Normally, this calculation is used to initialize a drift-diffusion calculation and isn't of interest.	logical (T/F)	false
TECLOTLEVEL	Setting for Tecplot output: 0 - no Tecplot files 1 - Tecplot files, each output in a separate file. 2 - Tecplot file, each output appended to a single file. Tecplot files will have the .dat suffix, and the prefix will be the name of the device instance	—	1
TXTDATALEVEL	Flag for volume-averaged text output. 0 - no text files. 1 - text files. txtdataplot files will have the *.txt suffix, and the prefix will be the name of the device instance.	—	1
<b><i>Scaling Parameters</i></b>			
X0	Length scalar; adjust to mitigate convergence problems. The model will do all of its scaling automatically, so it is generally not necessary to specify it manually.	cm	0.0001

Table 2.88: PDE Device Model Parameters

Parameter	Description	Units	Default
LEVEL	The level determines if this is a 1D or a 2D device. 1=1D, 2=2D	—	1

## Doping Parameters

Table 2.89: PDE Device Doping Region Parameters. These correspond to the region instance parameter.




Parameter	Description	Units	Default	Device Type
<b><i>All Levels</i></b>				
function	functional form of doping region. Options are uniform, gaussian, and step.		uniform	1D,2D

Table 2.89: PDE Device Doping Region Parameters. These correspond to the region instance parameter.

Parameter	Description	Units	Default	Device Type
type	Ntype or Ptype		ntype	1D,2D
nmax	Maximum value of impurity concentration.	$\text{cm}^{-3}$	1.0e15	1D,2D
nmin	Minimum value of impurity concentration.	$\text{cm}^{-3}$	1.0e15	1D,2D
xloc	Peak location	cm	0.0	1D,2D
xwidth	Distance from nmax to nmin, if applicable. This is only applicable for the function=gaussian case.		1.0e-3	1D,2D
flatx	This parameter determines if we're doing a half gaussian or a full gaussian. See table 2.90	-	0	1D,2D
<b>Level 2 (2D) only</b>				
yloc	Same as xloc, but for the y-direction.	cm	0.0	2D
ywidth	Same as xwidth, but for the y-direction.	cm	1.0e-3	2D
flaty	Same as flatx, but for the y-direction.	-	0	2D

## Flat Parameters

Table 2.90: Description of the flatx, flaty doping parameters

Flatx or Flaty view	Description	1D Cross Section
0	Gaussian on both sides of the peak (xloc) location.	
+1	Gaussian if $x > x_{loc}$ , flat (constant at the peak value) if $x < x_{loc}$ .	
-1	Gaussian if $x < x_{loc}$ , flat (constant at the peak value) if $x > x_{loc}$ .	

## Exectrode Parameters

Table 2.91: PDE Device Electrode Parameters.

Parameter	Description	Units	Default
<b>Level 2 (2D) only</b>			
name	Electrode name	-	anode
bc	Carrier Density Boundary condition type (dirichlet or neumann)	-	dirichlet

Table 2.91: PDE Device Electrode Parameters.

Parameter	Description	Units	Default
start	Starting location	cm	0.0
end	Ending location	cm	0.0
side	Side specification (top, bottom, left or right)	-	top
material	Contact material		neutral
oxidebndryflag	Oxide layer boolean	-	false (0)
oxthick	Oxide thickness	cm	0.0
oxcharge	Oxide charge	C	0.0

## 2.4.1 Physical Models

This section contains information about physical models used in **Xyce** for TCAD devices. This includes various mobility models, expressions for calculating the effective mass for electrons and holes, an expression for intrinsic carrier concentration as a function of temperature, expressions which describe contacts to metal as well as contacts to metal-oxide-semiconductor devices.

### Material Models and Parameters

This section describes some of the basic material properties that are available in **Xyce**. Described here are the models for effective mass, intrinsic carrier concentration, and the bandgap. This information is needed for the more complex models described in the mobility section (section 2.4.2) and the boundary condition section (section 2.4.2).

#### Effective Mass

**Xyce** includes functions which return the effective mass of electrons and holes for a number of semiconductor materials.

#### Electron Effective Mass

The electron effective mass is calculated as

$$m_{de} = (m_l^* m_t^{*2})^{1/3} \quad (2.21)$$

where  $m_l$  and  $m_t$  are the effective masses along the longitudinal and transverse directions of the ellipsoidal energy surface.

#### Hole Effective Mass

The hole effective mass is calculated as

$$m_{dh} = (m_{lh}^{*3/2} + m_{hh}^{*3/2})^{2/3} \quad (2.22)$$

where  $m_{lh}$  and  $m_{hh}$  are the "light" and "heavy" hole masses, respectively.

#### Intrinsic Carrier Concentration

The intrinsic carrier concentration in a semiconductor is obtained from the "np" product

$$np = n_i^2 = N_C N_V \exp(-E_g/kT) \quad (2.23)$$

or

$$n_i = \sqrt{N_C N_V} e^{-E_g/2kT} \quad (2.24)$$

The expression used in **Xyce** to calculate the intrinsic carrier concentration comes from this and is given by

$$n_i = 4.9 \times 10^{15} \left( \frac{m_{de} m_{dh}}{m_0^2} \right)^{3/4} M_c^{1/2} T^{3/2} e^{-E_g/2kT} \quad (2.25)$$

where  $M_c$  is the number of equivalent minima in the conduction band for the semiconductor,  $m_{de}$  is the density-of-state effective mass for electrons,  $m_{dh}$  is the density-of-state effective mass for holes, and  $m_0$  is the free-electron mass.

Table 2.92: Intrinsic Carrier Concentration Parameters

Semiconductor	Symbol	$M_c^{1/2}$	$n_i$ at room temperature
Silicon	si	$\sqrt{6.00}$	$1.25 \times 10^{10}$
Germanium	ge	2.00	$2.5 \times 10^{13}$
Galium Arsenide	gaas	1.00	$2.0 \times 10^6$

## Bandgap

The bandgap is a material and temperature-dependent quantity. The bandgap model for semiconductor materials, is based on Thurmond [30]. This model is given by:

$$E_g = E_{g0} - A * \left( \frac{T^{2.0}}{T + T_{off}} \right) \quad (2.26)$$

where  $E_g$  is the bandgap (eV) and  $T$  is the temperature (K).  $A$ ,  $E_{g0}$ , and  $T_{off}$  are all material-dependent constants. Insulating materials, such as silicon dioxide, are assumed to have constant bandgaps, so their bandgaps are given by:

$$E_g = E_{g0} \quad (2.27)$$

where  $E_{g0}$  is a material-dependent constant. The values for the material-dependent constants used by equations 2.26 and 2.27 are given in Table 2.93.

Table 2.93: Bandgap constants

Material	Symbol	$E_{g0}$ (eV)	$A$	$T_{off}$ (K)
----------	--------	---------------	-----	---------------

Table 2.93: Bandgap constants

Material	Symbol	$E_{g0}$ (eV)	$A$	$T_{off}$ (K)
Silicon	si	1.17	4.73e-4	636.0
Germanium	ge	0.7437	4.774e-4	235.0
Galium Arsenide	gaas	1.519	5.405e-4	204.0
Silicon Dioxide	sio2	9.00	NA	NA
Silicon Nitride	wdi	4.7	NA	NA
Sapphire	cu	4.7	NA	NA

## 2.4.2 Mobility Models

A number of mobility models are included in **Xyce**. The analytic, arora, and carrier-carrier scattering models are considered to be low-field mobility models. The Lombardi surface mobility model is a transverse-field dependent model which also incorporates the mobility of the bulk silicon.

### Analytic Mobility

This is a concentration- and temperature-dependent empirical mobility model, based on the work of Caughey and Thomas [31], which combines the effects of lattice scattering and ionized impurity scattering. The equation for the mobility of electrons is:

$$\mu_{0n} = \mu_{nmin} + \frac{\mu_{nmax}(\frac{T}{T_{ref}})^{nun} - \mu_{nmin}}{1 + (\frac{T}{T_{ref}})^{xin}(N_{total}/N_n^{ref})^{\alpha_n}} \quad (2.28)$$

and the equation for the mobility of holes is:

$$\mu_{0p} = \mu_{pmin} + \frac{\mu_{pmax}(\frac{T}{T_{ref}})^{nup} - \mu_{pmin}}{1 + (\frac{T}{T_{ref}})^{xip}(N_{total}/N_p^{ref})^{\alpha_p}} \quad (2.29)$$

where  $N_{total}$  is the local total impurity concentration (in  $\#/cm^3$ ),  $T_{ref}$  is a reference temperature (300.15K), and T is the temperature (in degrees K). The parameters  $N_n^{ref}$  and  $N_p^{ref}$  are reference values for the doping concentration. The analytic mobility model can be selected by using the statement "mobmodel=analytic" in the netlist.

The parameters for the analytic mobility model are given in Table 2.94.

Table 2.94: Analytic Mobility Parameters

Parameter	Silicon	GaAs
$\mu_{nmin}$	55.24	0.0
$\mu_{nmax}$	1429.23	8500.0
$N_n^{ref}$	1.072e17	1.69e17
nun	-2.3	-1.0
xin	-3.8	0.0
$\alpha_n$	0.73	0.436
$\mu_{pmin}$	49.70	0.0
$\mu_{pmax}$	479.37	400.0
$N_p^{ref}$	1.606e17	2.75e17
nup	-2.2	-2.1
xip	-3.7	0.0
$\alpha_p$	0.70	0.395

### Arora Mobility

This mobility model is also an analytic model which depends on impurity concentration and temperature. It comes from the work of Arora, *et al.* [32] and is based on both experimental data and the modified Brooks-Herring theory of mobility. The equation for the mobility of electrons is:

$$\mu_{0n} = \mu_{n1} \left( \frac{T}{T_{ref}} \right)^{exn1} + \frac{\mu_{n2} \left( \frac{T}{T_{ref}} \right)^{exn2}}{1 + \left( \frac{N_{total}}{C_n \left( \frac{T}{T_{ref}} \right)^{exn3}} \right)^{\alpha_n}} \quad (2.30)$$

and the equation for the mobility of holes is:

$$\mu_{0p} = \mu_{p1} \left( \frac{T}{T_{ref}} \right)^{exp1} + \frac{\mu_{p2} \left( \frac{T}{T_{ref}} \right)^{exp2}}{1 + \left( \frac{N_{total}}{C_p \left( \frac{T}{T_{ref}} \right)^{exp3}} \right)^{\alpha_p}} \quad (2.31)$$

where

$$\alpha_n = A_n \left( \frac{T}{T_{ref}} \right)^{exn4} \quad (2.32)$$

and

$$\alpha_p = A_p \left( \frac{T}{T_{ref}} \right)^{exp4} \quad (2.33)$$

The Arora mobility model can be selected by including the statement "mobmodel=arora" in the netlist. The parameters for the arora mobility model are given in Table 2.95.

Table 2.95: Arora Mobility Parameters

Parameter	Silicon	GaAs
$\mu_{n1}$	88.0	8.5e3
$\mu_{n2}$	1252.0	0.0
Cn	1.26e17	1.26e17
An	0.88	0.0
exn1	-0.57	-0.57
exn2	-2.33	0.0
exn3	2.4	0.0
exn4	-0.146	0.0
$\mu_{p1}$	54.3	4e2
$\mu_{p2}$	407.0	0.0
Cp	2.35e17	2.35e17
Ap	0.88	0.0
exp1	-0.57	0.0
exp2	-2.23	0.0
exp3	2.4	0.0
exp4	-0.146	0.0

## Carrier-Carrier Scattering Mobility

This mobility model is based on the work of Dorkel and Leturq [33]. It incorporates carrier-carrier scattering effects, which are important when high concentrations of electrons and holes are present in the device. This model also takes lattice scattering and ionized impurity scattering into account. One important difference between the carrier-carrier scattering mobility model and the two previous mobility models (analytic and arora models) is that the carrier-carrier scattering mobility model depends upon the actual carrier concentrations in the device. This model is important for modeling breakdown as well as various radiation effects, which often result in very high carrier densities.

The expressions for the carrier-carrier model are as follows:

$$\mu_L = \mu_{L0} \left( \frac{T}{T_{ref}} \right)^{-\alpha} \quad (2.34)$$

where  $\mu_L$  is the lattice mobility, which has to do with scattering due to acoustic phonons.

$$\mu_I = \frac{AT^{3/2}}{N} \left[ \ln\left(1 + \frac{BT^2}{N}\right) - \frac{BT^2}{N + BT^2} \right]^{-1} \quad (2.35)$$

where  $\mu_I$  is the impurity mobility which is related to the interactions between the carriers and the ionized impurities.

$$\mu_{ccs} = \frac{2 \times 10^{17} T^{3/2}}{\sqrt{pn}} \left[ \ln\left(1 + 8.28 \times 10^8 T^2 (pn)^{-1/3}\right) \right]^{-1} \quad (2.36)$$

where  $\mu_{ccs}$  is the carrier-carrier scattering mobility, which is very important when both types of carriers are at high concentration.

$$X = \sqrt{\frac{6\mu_L(\mu_I + \mu_{ccs})}{\mu_I\mu_{ccs}}} \quad (2.37)$$

is an intermediate term and

$$\mu = \mu_L \left[ \frac{1.025}{1 + (X/1.68)^{1.43}} - 0.025 \right] \quad (2.38)$$

is the carrier mobility. The carrier-carrier scattering mobility can be selected by including the statement "mobmodel=carr" in the netlist. The parameters for the carrier-carrier mobility model are given in Table 2.96.

Table 2.96: Carrier-Carrier Mobility Parameters

Parameter	Carrier	Silicon	GaAs
Al	$e^-$	1430.0	8.50e3
Bl	$e^-$	-2.2	0.0
Ai	$e^-$	4.61e17	4.61e17
Bi	$e^-$	1.52e15	1.52e15
Al	$h^+$	495.0	4.0e2
Bl	$h^+$	-2.2	0.0
Ai	$h^+$	1.00e17	1.00e17
Bi	$h^+$	6.25e14	6.25e14

### Lombardi Surface Mobility Model

This mobility model combines expressions for mobility at the semiconductor-oxide interface and in bulk silicon. It is based on the work of Lombardi *et al.* [34]. The overall mobility is found using Mathiessen's rule:

$$\frac{1}{\mu} = \frac{1}{\mu_{ac}} + \frac{1}{\mu_b} + \frac{1}{\mu_{sr}} \quad (2.39)$$

where  $\mu_{ac}$  is the carrier mobility due to scattering with surface acoustic phonons,  $\mu_b$  is the carrier mobility in bulk silicon, and  $\mu_{sr}$  is the carrier mobility limited by surface roughness scattering.

The Lombardi model is a more physics-based surface mobility model. It is a semi-empirical model for carrier mobility, and the expressions for the individual scattering mechanisms were extracted from experimental data taken in appropriate experimental conditions.

The expressions used in this model are given below:

$$\mu_{ac,n} = \frac{bn}{E_{\perp}} + \frac{cnN^{exn4}}{T(E_{\perp})^{1/3}} \quad (2.40)$$

is the expression for electron mobility for acoustic phonon scattering,

$$\mu_{ac,p} = \frac{bp}{E_{\perp}} + \frac{cpN^{exp4}}{T(E_{\perp})^{1/3}} \quad (2.41)$$

is the expression for hole mobility for acoustic phonon scattering,

$$\mu_{b,n} = \mu_{n0} + \frac{\mu_{max,n} - \mu_{n0}}{1 + (N/crn)^{exn1}} - \frac{\mu_{n1}}{1 + (csn/N)^{exn2}} \quad (2.42)$$

is the expression for bulk mobility for electrons, where

$$\mu_{max,n} = \mu_{n2} \left( \frac{T}{T_{ref}} \right)^{-exn3} \quad (2.43)$$

and

$$\mu_{b,p} = \mu_{p0} \exp(-pc/N) + \frac{\mu_{max,p}}{1 + (N/crp)^{exp1}} - \frac{\mu_{p1}}{1 + (csp/N)^{exp2}} \quad (2.44)$$

is the expression for bulk mobility for holes, where

$$\mu_{max,p} = \mu_{p2} \left( \frac{T}{T_{ref}} \right)^{-exp3} \quad (2.45)$$

The expression for electrons for surface roughness scattering is

$$\mu_{sr,n} = \left( \frac{dn}{E_{\perp}^{exn8}} \right) \quad (2.46)$$

and the expression for holes for surface roughness scattering is

$$\mu_{sr,p} = \left( \frac{dp}{E_{\perp}^{exp8}} \right) \quad (2.47)$$

The parameters for the lombardi surface mobility model are given in Table2.97.

Table 2.97: Lombardi Surface Mobility Parameters

Parameter	Silicon	GaAs
$\mu_{n0}$	52.2	0.0
$\mu_{n1}$	43.4	0.0
$\mu_{n2}$	1417.0	1e6
crn	9.68e16	9.68e16
csn	3.43e20	0.0
bn	4.75e7	1e10
cn	1.74e5	0.0
dn	5.82e14	1e6
exn1	0.680	0.0
exn2	2.0	0.0
exn3	2.5	0.0
exn4	0.125	0.0
exn8	2.0	0.0
$\mu_{p0}$	44.9	0.0
$\mu_{p1}$	29.0	0.0
$\mu_{p2}$	470.5	1.0
crp	2.23e17	2.23e17
csp	6.1e20	0.0
bp	9.93e6	1e10
cp	8.84e5	0.0
dp	2.05e14	1e6
exp1	0.719	0.0
exp2	2.0	0.0
exp3	2.2	0.0
exp4	0.0317	0.0
exp8	2.0	0.0
pc	9.23e16	0.0

## Edge Mobilities

Mobility values are calculated along the edge connecting two nodes. In the case of the analytic, arora, and surface mobility models, the edge mobilities are calculated by taking the average of the mobilities at the two nodes. Then, the mobility along the edge connecting nodes 1 and 2 is:

$$\mu_{edge} = (\mu[1] + \mu[2])/2.0 \quad (2.48)$$

In the case of the carrier-carrier scattering mobility, the edge mobilities were calculated differently. The electron and hole concentrations were first calculated at the midpoint of the edge using a "product" average and then these values of "n" and "p" were used in the function to calculate the mobility at the midpoint of the edge. For example, if  $n[1]$  and  $n[2]$  are the electron concentrations at nodes 1 and 2, the electron concentration along the edge is given by:

$$n_{edge} = \sqrt{n[1] * n[2]} \quad (2.49)$$

Subsequently, the mobility at the midpoint of an edge is found by using the values of electron and hole concentration at the midpoint of the edge when calling the function which returns the mobility, `calcMob()`.

$$\mu_{n,edge}^{carrier} = f(n_{edge}) \quad (2.50)$$

This method makes more sense, especially when the electron and hole concentrations vary by several orders of magnitude. Then it approximates taking the average of the logarithms.

## Boundary Conditions for Electrode Contacts

This section describes various boundary conditions that need to be applied to the semiconductor boundary. **Xyce** is predominantly an analog circuit simulator, and the TCAD (PDE-based) device modeling that has been implemented in **Xyce** takes external circuit information as input. This input consists of voltages and currents which are applied as boundary conditions to the semiconductor domain.

The physical connection from the circuit to the device generally includes a variety of materials, including metals and oxides. Electrical differences between the semiconductor and the contact material can result in a potential barrier that must be included in the imposed voltage boundary condition.

There are three general types of contacts between the circuit and the TCAD device that are handled by **Xyce**. The first is the "neutral" contact, in which it is simply assumed that the electrode material does not impose any additional potential barrier to that of the Fermi level differences in the semiconductor. The second is the Schottky contact, in which the electrode is a specified metal, and a potential barrier is imposed to account for the workfunction difference between the metal and the semiconductor. The last type of contact is the metal-oxide-semiconductor contact, in which the workfunction difference, and the voltage drop across the oxide must be accounted for.

## Neutral Contacts

A neutral contact refers to the case in which the contact is made to the semiconductor itself, and barrier heights due to material differences are not considered. This is the simplest type of contact in **Xyce**, and problems which use this type of contact are generally easier to solve, compared with other types of contacts. In this case, the boundary is given by

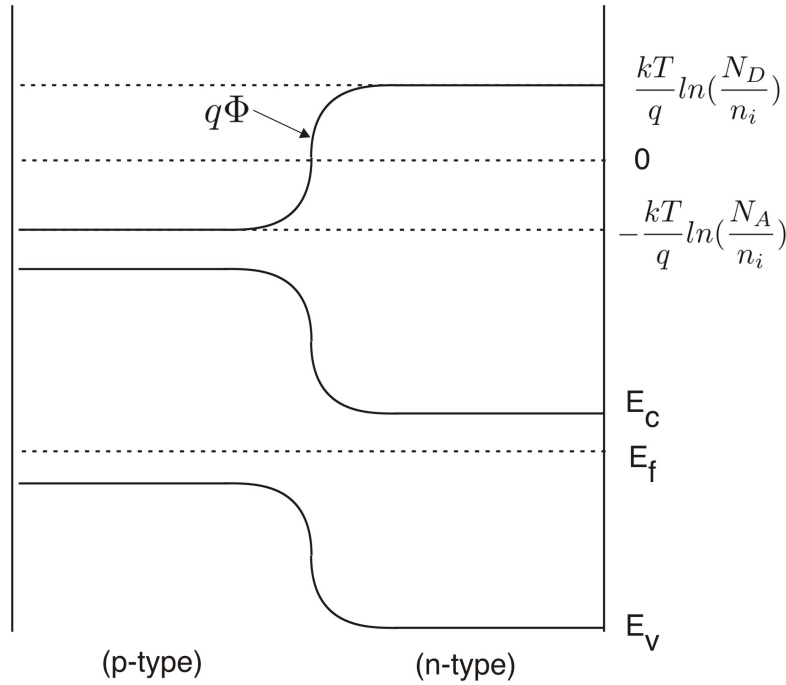
$$V_{bc} = V_{ckt} + V_{bi} \quad (2.51)$$

where  $V_{ckt}$  is the potential applied by the circuit and  $V_{bi}$  is the "built-in" potential of the semiconductor. For a p-type substrate, the built-in potential is given by

$$V_{bi} = -\frac{kT}{q} \ln\left(\frac{N_A}{n_i}\right) \quad (2.52)$$

and for an n-type substrate, the built-in potential is given by

$$V_{bi} = \frac{kT}{q} \ln\left(\frac{N_D}{n_i}\right) \quad (2.53)$$



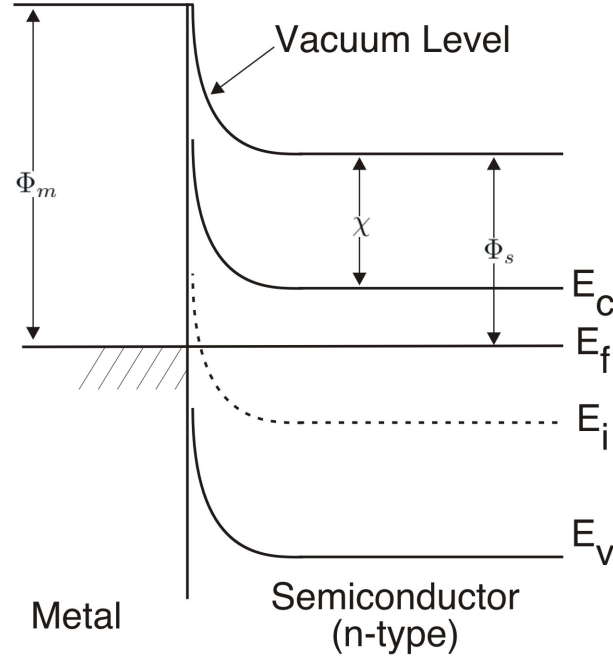
**Figure 2.2.** Neutral Contacts.

$V_{bi}$  represents the extent of the energy band bending due to the doping of a device. While most of the dramatic changes will happen away from the contact, near junctions, it is still incorporated into the voltage boundary condition to maintain a flat potential near the contacts. Figure 2.2 shows the energy band variation across a PN junction, and the corresponding electrostatic potential. This variation is due to the internal physics of the device, and needs to be there even in the event of zero applied voltage. This is partially enforced by the solution to Poisson's equation, and also by the application of equation 2.51.

## Schottky Contacts

In the case of a metal-semiconductor contact, it is necessary to add the workfunction difference,  $\Phi_{ms}$ , to the potential in the semiconductor [35].  $\Phi_m$  is a constant for a given metal, and  $\Phi_s$  is a

function of the doping in the semiconductor. The workfunction potential,  $\Phi$ , when multiplied by  $q$ , is the difference between the Fermi level and vacuum in the material. In essence, the workfunction difference represents the distance between the Fermi level in the metal and the Fermi level in the semiconductor when considering the individual band structures.



**Figure 2.3.** Schottky Contact, N-type.

In the case of an n-type semiconductor, the semiconductor workfunction can be represented as

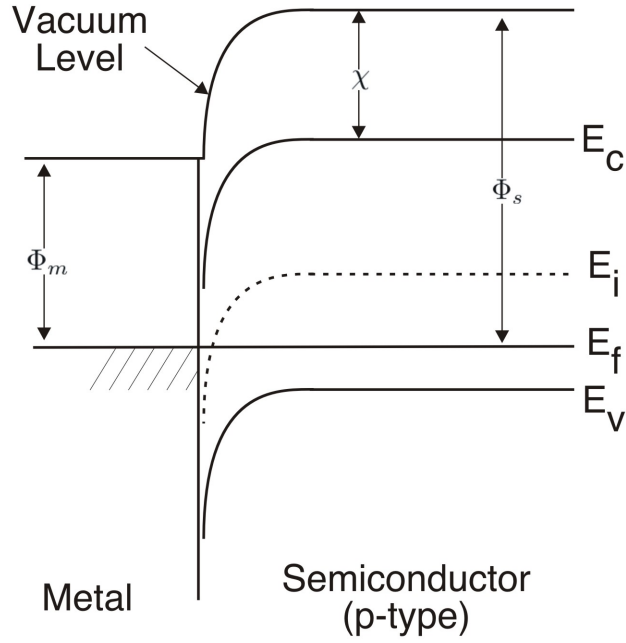
$$\Phi_s = \chi + (E_C - E_{FS})/q \quad (2.54)$$

where  $\chi$  is the electron affinity in the semiconductor and  $q\chi$  is the distance between the conduction band and vacuum in the semiconductor.  $E_C$  is the conduction band energy and  $E_{FS}$  is the Fermi level of the semiconductor. Rewriting this expression in terms of the doping concentration, it becomes

$$\Phi_s = \chi + E_g/2 - V_t \ln\left(\frac{N_d}{n_i}\right) \quad (2.55)$$

In the case of a p-type semiconductor, the semiconductor workfunction can be represented as

$$\Phi_s = \chi + E_g/2 + (E_i - E_{FS})/q \quad (2.56)$$



**Figure 2.4.** Schottky Contact, P-type.

where  $E_i$  is the intrinsic value of the Fermi level, and can be approximated as the halfway point between the conduction band ( $E_C$ ) and the valence band ( $E_V$ ). Rewriting this expression in terms of the doping concentration

$$\Phi_s = \chi + E_g/2 + V_t \ln\left(\frac{N_a}{n_i}\right) \quad (2.57)$$

For the TCAD devices in **Xyce**, for a node at a metal-semiconductor contact, the quantity  $\Phi_m - \Phi_s$  is added to the potential at the node to account for the metal-semiconductor barrier. The current values of metal workfunctions used in **Xyce** are given in Table 2.98. The values for electron affinity are given in Table 2.99. The boundary condition for a metal electrode in **Xyce** is given by

$$V_{bc} = V_{ckt} + V_{bi} + \Phi_{ms} \quad (2.58)$$

where  $V_{ckt}$  is the potential applied by the circuit to the electrode and  $V_{bi}$  is the "built-in" potential of the semiconductor, a function of the semiconductor doping.

Table 2.98: Material workfunction values

Metal	Symbol	Workfunction, $\Phi_m$ (Volts)
aluminum	al	4.10
p+-polysilicon	ppoly	5.25
n+-polysilicon	npoly	4.17
molybdenum	mo	4.53
tungsten	w	4.63
molybdenum disilicide	modi	4.80
tungsten disilicide	wdi	4.80
copper	cu	4.25
platinum	pt	5.30
gold	au	4.80

Table 2.99: Electron affinities

Semiconductor	Symbol	Electron Affinity, $\chi$ (Volts)
Silicon	si	4.17
Germanium	ge	4.00
Galium Arsenide	gaas	4.07
Silicon Dioxide	sio2	0.97
Nitride	nitride	0.97
Sapphire	sapphire	0.97

## Metal-Oxide-Semiconductor Contacts

To date in **Xyce**, only semiconductor material is included in the PDE solution domain. Metals and oxide materials are only included through boundary conditions. This is an adequate approach for a lot of problems. For some problems (such as modeling of low-dose radiation effects) modeling the oxide in more detail, as a PDE, will become necessary. However, since oxides are usually very thin, compared with the semiconductor domain, meshing both materials as part of the same simulation is difficult. Therefore, incorporating the effects of a gate oxide as part of the gate boundary condition is a reasonable approach.

In the case of a contact to a metal-oxide-semiconductor structure, the separation of the Fermi energies in the metal and the semiconductor at equilibrium is due to two effects: the workfunction difference between the metal and the semiconductor, and the effective interface charge. These two effects cause the bands to bend at the surface in equilibrium. The flatband voltage is the sum of these two terms [35]:

$$V_{FB} = \Phi_{ms} - \frac{Q_i}{C_i} \quad (2.59)$$

where  $\Phi_{ms}$  is the metal-semiconductor workfunction difference,  $Q_i$  is the value of interface charge (in  $C/cm^2$ ), and  $C_i$  is the oxide capacitance per unit area, which is given by

$$C_i = \frac{\epsilon_{ox}\epsilon_0}{x_o} \quad (2.60)$$

The voltage  $V_{FB}$  is the amount of bias which, when applied to the gate, causes the electron energy bands to be flat. This is the potential that is added to a boundary node in **Xyce** to account for a metal-oxide-semiconductor barrier. The overall boundary condition for a contact to a metal-oxide-semiconductor structure is given by

$$V_{bc} = V_{ckt} + V_{bi} + \Phi_{ms} - Q_i/C_i \quad (2.61)$$

where  $V_{ckt}$  is the potential applied by the circuit and  $V_{bi}$  is the "built-in" potential of the semiconductor.

## NMOS Device

The default NMOS device used in **Xyce** has a substrate doping concentration of  $1.0 \times 10^{16}/cm^3$  and an oxide thickness of  $1.0 \times 10^{-6}cm$ . Since the ideal threshold voltage  $V_T$  is given by

$$V_T = 2\phi_F + \frac{\epsilon_s}{\epsilon_{ox}}x_o\sqrt{\frac{2qN_A\phi_F}{\epsilon_s\epsilon_0}} \quad (2.62)$$

$V_T$  is equal to 0.892 V. for this device. Note that

$$\phi_F = \frac{1}{q}[E_i(bulk) - E_F] = \frac{kT}{q} \ln\left(\frac{N_A}{n_i}\right) \quad (2.63)$$

for a p-type semiconductor substrate and

$$\phi_F = -\frac{kT}{q} \ln\left(\frac{N_D}{n_i}\right) \quad (2.64)$$

for an n-type substrate.

# 3. Command Line Arguments

**Xyce** supports a handful of command line arguments which must be given *before* the netlist filename. While most of these are intended for general use, others simply give access to new features that, while supported, are not enabled by default. The general usage is as follows:

```
runxyce [arguments] <netlist filename>
```

Table 3.1 gives a list of supported command line options.<sup>1</sup>

Table 3.1: List of **Xyce** command line arguments.

Argument	Description	Usage	Default
-h	Help option. Prints usage and exits.	-h	-
-v	Prints the version banner and exits.	-v	-
-license	Prints the license text and exits.	-license	-
-capabilities	Prints a list of compiled-in options and exits.	-capabilities	-
-delim	Set the output file field delimiter.	-delim <TAB COMMA string>	-
-o	Place the results into specified file.	-o <file>	Results output file name based on netlist file name.
-l	Place the log output into specified file.	-l <file>	Log output sent to standard out.
-r	Output a binary rawfile.	-r <file>	No rawfile written.
-a	Use with -r to output a readable (ascii) rawfile.	-r <file> -a	Default rawfile is binary.
-nox	Use the NOX nonlinear solver.	-nox <ON OFF>	on
-linsolv	Set the linear solver.	-linsolv <KLU AZTEC00>	KLU(serial) and AztecOO(parallel)
-param	Print a terse summary of model and/or device parameters.	-param	-

<sup>1</sup>Note that the “-h” option might list command line options not present in this table. These extra options are generally deprecated and should not be used. Only the options listed in the table are considered supported features.

Table 3.1: List of **Xyce** command line arguments.

Argument	Description	Usage	Default
-syntax	Check netlist syntax then exit.	-syntax	-
-norun	Check netlist syntax and topology, then exit.	-norun	-
-maxord	Maximum time integration order.	-maxord <1..5>	-
-jacobian_test	Jacobian matrix diagnostic.	-jacobian_test	-

# 4. Runtime Environment

## 4.1 Running **Xyce** in Serial

If **Xyce** was installed from one of Sandia's binary installers, use the `runxyce` script to run serial versions of **Xyce**. This script sets the environment variables, `LD_LIBRARY_PATH` or `DYLD_LIBRARY_PATH`, and starts **Xyce**. No additional runtime configuration is necessary, as these binary installers are shipped with the shared libraries they require..

If **Xyce** was built from source and is being run on the machine where it was compiled, then generally no `LD_LIBRARY_PATH` or `DYLD_LIBRARY_PATH` settings are required, and so **Xyce** is run directly without a wrapper script. After ensuring that the directory into which **Xyce** was installed is in your `PATH` variable, one merely executes the code by running the command, `Xyce`.

## 4.2 Running **Xyce** in Parallel

Open MPI must be installed on the host machine. It may be download from

<http://www.open-mpi.org/>. Consult the documentation for help with installation.

Use the `xmpirun` to run parallel versions of **Xyce**. This script sets the environment variables, `LD_LIBRARY_PATH` or `DYLD_LIBRARY_PATH`, and starts `Xyce` using the `mpiexec` wrappers.

If **Xyce** was built from source and is being run on the machine where it was compiled, then generally no `LD_LIBRARY_PATH` or `DYLD_LIBRARY_PATH` settings are required, and so **Xyce** is run directly without a wrapper script. After ensuring that the directory into which **Xyce** was installed is in your `PATH` variable, one merely executes the code by running the command, `mpirun [mpirun options] Xyce [xyce options]`.

## 4.3 Running **Xyce** on Sandia HPC and CEE Platforms

This version of **Xyce** has been installed centrally on Sandia HPC and CEE platforms, and requires metagroup access. Contact the **Xyce** team for details on how to obtain this access.

Once you have registered for metagroup membership, the central installs of **Xyce** may be accessed by a module load.

`module load Xyce` adds all required modules and sets all required environment variables to access the normal version of **Xyce**. `module load XyceRad` does the same thing for the version **Xyce** containing Sandia proprietary models.

`module help Xyce` provides some additional information about what the module does.

Consult the system documentation for help with submitting jobs on these platforms.

<https://computing.sandia.gov/>

# 5. Setting Convergence Parameters for **Xyce**

Because the solution algorithms and methods within **Xyce** are different than those used by other circuit simulation tools (e.g., SPICE), the overall convergence behavior is sometimes different, as are the parameters which control this behavior.

## 5.1 Adjusting Transient Analysis Error Tolerances

**Xyce** uses a variable order trapezoid integration as its default scheme, and this method may also be requested explicitly with the **TIMEINT** option **METHOD=trap** or **METHOD=7**. Trapezoid time-stepping is second order accurate and does not have any numerical dissipation in its local truncation error. Variable order trapezoid integration dynamically uses Backward euler (BE) and trapezoid rule. When **ERROPTION=1** is set with **METHOD=7**, trapezoid rule is used almost exclusively (BE only used at breakpoints). See table 2.2 for details.

**Xyce** also supports a variable order Backward Differentiation Formula (BDF 1-5) time integration method (also known as a 1-5 step Gear method) for performing transient analysis [36]. The BDF integrator is selected by using the **TIMEINT** option **METHOD=BDF** or **METHOD=6**. This method starts out with Backward Euler on the first few steps and then ramps up to as high an order as will maintain stability, and which takes the largest time steps. The maximum order it can attain is five, and this can be reduced with the **MAXORD** option. It is also possible to set a minimum order that the integrator should maintain with the option **MINORD**. When **MINORD** is set, the integrator will move upward in order from Backward Euler as quickly as possible to achieve **MINORD**, and then it will adjust the order between **MINORD** and **MAXORD** to maintain stability and take large steps. See table 2.2 for details.

A third time integration option is the second-order Gear method. It offers some improvements over the BDF implementation, and may be selected with the **TIMEINT** option **METHOD=gear** or **METHOD=8**. See table 2.2 for details.

### 5.1.1 Setting **RELTOL** and **ABSTOL**

In **Xyce**, both the time integration package and the nonlinear solver package have **RELTOL** and **ABSTOL** settings. Some general guidelines for settings parameters are [36]:

- Use the *same* **RELTOL** and **ABSTOL** values for both the **TIMEINT** and the **NONLIN-TRAN .OPTIONS** statements.
- For a conservative approach (i.e., safe), set  $\text{RELTOL} = 1.0\text{E}-(m+1)$  where  $m$  is the desired number of significant digits of accuracy.
- Set **ABSTOL** to the smallest value at which the solution components (either voltage or current) are essentially insignificant.
- Note that the above suggests that  $\text{ABSTOL} < \text{RELTOL}$ .

The current defaults for these parameters are  $\text{ABSTOL} = 1.0\text{E}-6$  and  $\text{RELTOL} = 1.0\text{E}-3$ . For a complete list of the time integration parameters, see chapter 2.1.

## 5.2 Adjusting Nonlinear Solver Parameters (in transient mode)

In **Xyce**, the nonlinear solver options for transient analysis are set using the **.OPTIONS NONLIN-TRAN** line in a netlist. This subsection gives some guidelines for setting these parameters.

- For guidelines on setting **RELTOL** and **ABSTOL**, see above.
- **RHSTOL** – This is the maximum residual error for each nonlinear solution. **Xyce** uses this as a “safety” check on nonlinear convergence. Typically,  $1.0\text{E}-2$  (the default) works well.
- **DELTAXTOL** – This is the weighted update norm tolerance and is the primary check for nonlinear convergence. Since it is weighted (i.e., normalized using **RELTOL** and **ABSTOL**), a value of 1.0 would give it the same accuracy as the time integrator. For robustness, the default is 0.33 but sometimes a value of 0.1 may help prevent “time-step too small” errors. A value of 0.01 is considered quite small.
- **MAXSTEP** – This is the maximum number of Newton (nonlinear) steps for each nonlinear solve. In transient analysis, the default is 20 but can be increased to help prevent “time-step too small” errors. This is roughly equivalent to **ITL4** in SPICE.

# 6. Quick Reference for Users of Other SPICE Circuit Simulators

This chapter describes many of the differences between **Xyce** and other SPICE-like circuit simulators. The primary focus is on the difference between Orcad PSpice and **Xyce**, with an eye towards providing the ability for those familiar with using PSpice to begin using **Xyce** quickly.

This chapter is likely not complete, and **Xyce** users might also consult specific sections of this Reference Guide about particular **Xyce** commands. Those sections may have additional information on Xyce's incompatibilities with other circuit simulators, and how to work around them.

## 6.1 Differences Between **Xyce** and PSpice

This section is focused on the differences between **Xyce** and PSpice. However, some of this discussion also applies to other SPICE-like circuit simulators.

### 6.1.1 Command Line Options

Command line arguments are supported in **Xyce** but they are different than those of PSpice. For a complete reference, see Chapter 3.

### 6.1.2 Device Support

Most, but not all, devices commonly found in other circuit simulation tools are supported. **Xyce** also contains enhanced versions of many semiconductor devices that simulate various environmental effects. For the complete list, please see the Analog Device Summary in Table 2.20.

### 6.1.3 Netlist Support

For the specific devices or models that are supported in **Xyce**, most of the standard netlist inputs are the same as those in standard SPICE. However, the **.OPTIONS** command has several additional features used to expose capabilities specific to **Xyce**. In particular, **Xyce** does not support the standard PSpice format **.OPTIONS** line in netlists. Instead, options for each supported package are called according to the following format.

**General Form**     **.OPTIONS** <pkg> [<tag>=<value>]\*

---

## Arguments and Options

DEVICE	Device Model
TIMEINT	Time Integration
NONLIN	Nonlinear Solver
NONLIN-TRAN	Transient Nonlinear Solver
NONLIN-HB	HB Nonlinear Solver
LOCA	Continuation/Bifurcation Tracking
LINSOL	Linear Solver
LINSOL-HB	HB Linear Solver
OUTPUT	Output
RESTART	Restart
HBINT	Harmonic Balance (HB)
SENSITIVITY	Direct and Adjoint sensitivity analysis

For a complete description of the supported options in **Xyce**, see section 2.1.16.

**Xyce** does not support the “**.PROBE**” statement. Output of Probe-format files, that are readable by PSpice is done using the **.PRINT** netlist statement. See section 2.1.21 for the syntax.

**Xyce** does not support PSpice style abbreviations in the **.PRINT** statement. For example, to print out the value of the voltage at node A in a transient simulation you must request **.PRINT TRAN V(A)**, not **.PRINT TRAN A**. **Xyce** also does not support **N()** as a synonym for **V()** on **.PRINT** lines.

## 6.1.4 Converting PSpice ABM Models for Use in Xyce

**Xyce** is almost fully compatible with PSpice with respect to analog behavioral models. This includes the E, F, G, and H device types. A notable exception to this compatibility is in the use of lead and device currents in expressions. These are limited to expressions in the **.PRINT** statement.

## 6.1.5 Usage of .STEP Analysis

The implementation of **.STEP** in **Xyce** is not the same as that of PSpice. See section 2.1.24 for the syntax and function of the **.STEP** function in **Xyce**.

### Global .PARAM Sweeps

PSpice also supports sweeps over variables specified in **.PARAM** lines. This is not supported in **Xyce**. For example, this block of text will not work in **Xyce**:

```
VAB 2 0 5
VAC 1 0 {variable}
.param variable=0
.step param variable 0 5 1
.dc VAB 4 5 1
```

An equivalent block of code that will work in **Xyce** replaces the **.param** with a **.global\_param**, and removes the **param** keyword from the **.step** line:

```
VAB 2 0 5
VAC 1 0 {variable}
.global_param variable=0
.step variable 0 5 1
.dc VAB 4 5 1
```

### Model Parameter Sweeps

PSpice requires extra keywords to apply a **.STEP** statement to a model parameter. **Xyce** handles model parameters differently, and is actually somewhat more flexible than PSpice. Unfortunately, this means that the two specifications are not compatible.

A model parameter in PSpice would be handled like this:

```
R1 1 2 RMOD 1
.model RMOD RES(R=30)
.step RES RMOD(R) 30 50 5
```

The equivalent way to specify this in **Xyce** would be:

```
R1 1 2 RMOD 1
.model RMOD RES(R=30)
.step RMOD:R 30 50 5
```

Note that **Xyce** does not require the RES keyword on the .STEP line. In PSpice, this keyword is needed to specify what type of model is being used. **Xyce** actually has more flexibility than PSpice in this regard—any model or instance variable can be set on the .STEP line using the same syntax.

**Example:**            .step D101:IS 1.0e-3 5.0e-3 1.0e-3

In this example, D101 is the name of a model or instance, and IS is the name of the parameter within that model or instance.

## 6.1.6 Behavioral Digital Devices

There are at least four significant differences. First, the instance line syntax for the **Xyce** digital behavioral devices differs from PSpice. Second, **Xyce** uses one model card for the timing and Input/Output (I/O) characteristics, while PSpice uses separate model cards for timing and I/O characteristics. The model cards also have different parameters. Third, the DCOP calculations for flip-flops and latches may be different between **Xyce** and PSpice. Finally, closely spaced input transitions to a gate (e.g., ones spaced by less than the DELAY parameter of the **Xyce** model) may produce different behaviors in **Xyce** and PSpice. Please consult Section 2.3.23 for more details.

## 6.1.7 Power Dissipation

PSpice supports printing the power dissipation of a device via syntax like W(<name>). This feature is not directly supported by **Xyce** on its .PRINT lines. Example work-arounds are as follows, using either the node voltage at Node 2 or the lead current through Resistor 2:

```
.DC V1 0 5 1
.param R2VAL=10
V1 1 0 5V
R1 1 2 10
R2 2 0 {R2VAL}
.PRINT DC V(2) {V(2)*V(2)/R2VAL} {I(R2)*I(R2)*R2VAL}
```

## 6.1.8 Dependent Sources with TABLE Syntax

PSpice and **Xyce** have two differences in their syntax for dependent sources that are specified with a TABLE syntax. PSpice might specify:

```
TABLE {EXPR} ((x1,y1) (x2,y2) ... (xn, yn))
```

PSpice tables might also be missing the commas between the value-pairs. For example, this is legal PSpice syntax:

```
TABLE {EXPR} ((x1 y1) (x2 y2) ... (xn yn))
```

The corresponding **Xyce** syntax is as follows, where an extra = and extra commas are needed and the outer parentheses are omitted:

TABLE {EXPR}= (x1,y1) (x2,y2) ... (xn, yn)

## 6.1.9 MODEL STATEMENTS

In PSpice, some .MODEL statements may have commas separating the list of parameters, which causes problems in **Xyce**. A simple workaround is to replace those commas with spaces in the corresponding **Xyce** .MODEL statements.

In PSpice, some .MODEL statements may not have parentheses surrounding the list of parameters. While Xyce also does not require parentheses in model cards, parentheses are accepted. The only **Xyce** requirement is that if they are used then they must be paired with a left parenthesis before all of the parameters and a right parentheses after all of the parameters. Unmatched parentheses are an error in **Xyce**.

## 6.1.10 Additional differences

Some other differences between **Xyce** and PSpice are described in Table 6.1. Users should also consult Table 6.2, since that table lists more general incompatibilities that span multiple circuit simulators.

Table 6.1: Incompatibilities with PSpice.

Issue	Comment
.VECTOR, .WATCH, and .PLOT output control analysis are not supported.	<b>Xyce</b> does not support these commands.
.NOISE and .SENS and .TF analysis types are not supported.	<b>Xyce</b> fully supports .DC, .TRAN, .AC and .OP analysis. .SENS is partially supported.
.MC and .WCASE statistical analyses are not supported.	<b>Xyce</b> does not support these commands.
.DISTRIBUTION, which defines a user distribution for tolerances, is not supported.	<b>Xyce</b> does not support this command. This command goes along with .MC and .WCASE statistical analyses, which are also not supported.
.LOADBIAS and .SAVEBIAS initial condition commands are not supported.	<b>Xyce</b> does not support these commands.
.ALIASES, .ENDALIASES, are not supported.	<b>Xyce</b> does not support these commands.
.STIMULUS is not supported.	<b>Xyce</b> does not support this command.
.TEXT is not supported.	<b>Xyce</b> does not support this command.
.PROBE does not work	<b>Xyce</b> does not support this. Use the FORMAT=PROBE option of .PRINT instead. See section 2.1.21 for syntax.

.OP only produces output in serial	.OP is supported in <b>Xyce</b> , but will not produce the extra output normally associated with the .OP statement, if running a parallel build.
Pulsed source rise time of zero	A requested pulsed source rise/fall time of zero really is zero in <b>Xyce</b> . In other simulators, requesting a zero rise/fall time causes them to use the printing interval found on the tran line.
Mutual Inductor Model	Not the same as PSpice. This is a Sandia developed model.
.PRINT line shorthand	Output variables have to be specified as a V(node) or I(source). Listing the node alone will not work.
BSIM3 level	In <b>Xyce</b> the BSIM3 level=9. In PSpice the BSIM3 is level=8.
Interactive mode	<b>Xyce</b> does not have an interactive mode.
Time integrator default tolerances	<b>Xyce</b> has much tighter default solver tolerances than some other simulators (e.g., PSpice), and thus often takes smaller time steps. As a result, it will often take a greater number of total time steps for a given time interval. To have <b>Xyce</b> take time steps comparable to those of PSpice, set the RELTOL and ABSTOL time integrator options to larger values (e.g., RELTOL=1.0E-2, ABSTOL=1.0E-6).
.OPTIONS statements	<b>Xyce</b> does not support PSpice style .OPTION statements. In <b>Xyce</b> , the various packages all (potentially) have their own separate .OPTIONS line in the netlist. For a complete description, see section 2.1.16.
DTMAX	<b>Xyce</b> does support a maximum time step-size control on the .tran line, but we discourage its use. The time integration algorithms within <b>Xyce</b> use adaptive time-stepping methods that adjust the time-step size according to the activity in the analysis. If the simulator is not providing enough accuracy, the RELTOL and ABSTOL parameters should be decreased for both the time integration package (.OPTIONS TIMEINT) and the transient nonlinear solver package (.OPTIONS NONLIN-TRAN). We have found that in most cases specifying the same maximum timestep that PSpice requires for convergence actually slows <b>Xyce</b> down by preventing it from taking larger timesteps when the behavior warrants.
.TRAN "UIC" keyword	PSpice requires the use of a keyword UIC on the .TRAN line in order to use initial conditions via IC keywords on instance lines. Doing so also tells PSpice not to perform an operating point calculation. In <b>Xyce</b> , UIC is ignored and produces a warning message. <b>Xyce</b> always uses initial conditions specified with IC keywords, and the case of inductors and capacitors automatically inserts a fictitious voltage source around the device that guarantees the correct potential drop across the device during the operating point. If the user desires that <b>Xyce</b> not perform an operating point calculation, but rather use an initial condition for a transient run of all zero voltages, then the user should specify NOOP instead.
Temperature specification	Device temperatures in <b>Xyce</b> are specified through the .OPTIONS DEVICE line. PSpice allows a .TEMP line that is not recognized (and is ignored) by <b>Xyce</b> .

## 6.1.11 Translating Between PSpice and **Xyce** Netlists

Some internal Sandia users have found the following checklist to be helpful in getting their PSpice netlists to run in **Xyce**. Additional changes may be needed in some cases.

For the .cir file:

- Change .LIB references to point to the modified libraries generated for use with **Xyce**.
- Change PROBE64 statements to PRINT <Sim Type>
- Find cases where the PSpice netlist used N() rather than V().
- .PARAM statements need to be replaced with .GLOBAL\_PARAM statements in **Xyce**.
- .DC has the keyword PARAM in PSpice. If it exists then remove it in the **Xyce** netlist.
- .OPTIONS TNOM=X is changed to .OPTIONS DEVICE TNOM=X in the **Xyce** netlist.
- .TEMP args does not exist in **Xyce**. The equivalent **Xyce** statement is .STEP TEMP LIST args
- The default time integrator tolerances can make **Xyce** take smaller timesteps on some circuits, and therefore have slower simulation times. The **Xyce** timesteps can be increased at the expense of time integration accuracy by loosening the integrator tolerances. Some users find that .OPTIONS TIMEINT RELTOL=1e-2 ABSTOL=1e-4 leads to time steps more like PSpice's.

For the .lib file:

- Add LEVEL=2 parameter to diode models.
- Fix the parentheses and comma differences between PSpice and **Xyce**.MODEL statements per Section 6.1.9.
- Find and modify any nested expression statements. This may entail replacing "{" with "(" in the expression in the **Xyce** netlist.
- Fix the table syntax for dependent sources, as discussed in Section 6.1.8.

## 6.2 Differences Between **Xyce** and Other SPICE Simulators

This section covers some known differences between **Xyce** and other SPICE-like circuit simulators, besides PSpice, as listed in Table 6.2. However, users of those other simulators (e.g., SPICE3F5, HSPICE, ngspice, ...) should also check the previous subsection on PSpice, since some of that discussion also applies here.

Table 6.2: Incompatibilities with Other Circuit Simulators.

Issue	Comment
.DC sweep output.	The .DC sweep calculation does not automatically output the sweep variable. Only variables explicitly listed on the .PRINT line are output.
MOSFET levels.	In <b>Xyce</b> the MOSFET levels are not the same. In <b>Xyce</b> , a BSIM3 is MOSEFET level 9. Other simulators have different levels for the BSIM3.
BSIM SOI v3.2 level.	In <b>Xyce</b> the BSIM SOI (v3.2) is MOSFET level 10. Other simulators have different levels for the BSIM SOI.
BSIM4 level.	In <b>Xyce</b> the BSIM4 is MOSFET levels 14 and 54. Other simulators have different levels for the BSIM4.
Syntax for .STEP is different.	The manner of specifying a model parameter to be swept is slightly different than in some other simulators. See the <b>Xyce</b> Users' and Reference Guides for details.
Switch is not the same as SPICE3F5.	The <b>Xyce</b> switches are not compatible with the simple switch implementation in SPICE3F5. The switch in <b>Xyce</b> smoothly transitions between the ON and OFF resistances over a small range between the ON and OFF values of the control signal (voltage, current, or control expression). See the <b>Xyce</b> Reference Guide for the precise equations that are used to compute the switch resistance from the control signal values. The SPICE3F5 switch has a single switching threshold voltage or current, and RON is used above threshold while ROFF is used below threshold. <b>Xyce's</b> switch is considerably less likely to cause transient simulation failures. Results similar to SPICE3F5 can be obtained by setting VON and VOFF to the same threshold value, but this is not a recommended practice.
Piecewise Linear (PWL) source not fully compatible with either HSpice or PSpice.	See Section 2.3.7 of the <b>Xyce</b> Reference Guide for more details.
Acceptable prefixes in the metric system.	The "atto" prefix, which is designated by "a", is acceptable in HSpice, but is not accepted in <b>Xyce</b> . The use of the "atto" prefix in <b>Xyce</b> must be replaced with "E-18".
Hierarchical parameters.	In <b>Xyce</b> hierarchical parameters, M (multiply) and S (scale), are not commonly supported. The M parameter is only supported by the MOSFET models.

## 7. Quick Reference for Microsoft Windows Users

**Xyce** is supported on Microsoft Windows. However, the primary targets for **Xyce** are high-performance supercomputers and workstations, which are almost always running a variant of Unix. All of **Xyce** development is done on Unix platforms. Bearing this in mind, there are occasionally issues with using a Unix application on a Windows platform. Some of these issues are described in the table below.

Table 7.1: Issues for Microsoft Windows.

Issue	Comment
File names are case-sensitive	<b>Xyce</b> will expect library files, which are referenced in the netlist, to have exactly the same case as the actual filename. If not, <b>Xyce</b> will be unable to find the library file.
<b>Xyce</b> is unable to read proprietary file formats.	Programs such as Microsoft Word by default use file formats that <b>Xyce</b> cannot recognize. It is best not to use such programs to create netlists, unless netlists are saved as *.txt files. If you must use a Microsoft editor, it is better to use Microsoft Notepad. In general, the best solution is to use a Unix-style editor, such as Vi, Gvim, or Emacs.

# 8. Rawfile Format

The rawfile format produced by **Xyce** closely follows SPICE3 conventions. Differences are noted in section 8.3. Details on the both the ASCII and binary formats are provided here for reference.

## 8.1 ASCII Format

The ASCII file format can be created using the `-a` flag on the command line. See Chapter 3 for more information.

The ASCII format standard dictates that the file consist of lines or sets of lines introduced by a keyword. The `Title` and `Date` lines should be the first in the file, and should occur only once. They are followed by the `Plotname`, `Flags`, `No. Variables`, and `No. Points` lines for each plot.

Listed next are sets of `Variables`, and `Values` lines. Let *numvars* be the number of variables (as specified in the `No. Variables` line), and *numpts* be the number of points (as shown on the `No. Points` line). After the `Variables` keyword there must be *numvars* declarations of outputs, and after the `Values` keyword, there must be *numpts* lines, each consisting of *numvars* values.

Finally, **Xyce** also allows for a `Version` line to be placed after the `No. Points` line for compatability with various software programs.

See Table 8.1 for a summary of the above.

Table 8.1: **Xyce** ASCII rawfile format.

Issue	Comment
Title:	An arbitrary string describing the circuit.
Date:	A free-format date string.
Plotname:	A string describing the analysis type.
Flags:	A string describing the data type ( <i>real</i> or <i>complex</i> ).
No. Variables:	The number of variables.
No. Points:	The number of points.
Version: (optional)	The version of <b>Xyce</b> used to generate this output. By default the version is not output in the header. It can be output with the <code>.options output outputversioninrawfile=true</code> option.
Variables:	A newline followed by multiple lines, one for each variable, of the form [tab] <index> [tab] <name> [tab] <type>.

Values:	A newline followed by multiple lines, for each point and variable, of the form [tab] <value> with an integer index preceeding each set of points. Complex values are output as [tab] <real component>, <imaginary component> .
---------	--

## 8.2 Binary Format

The binary format is similar to the ASCII format, except that strings are null terminated rather than newline terminated. In addition, all the `values` lines are stored in a binary format. The binary storage of real values as double precision floats is architecture specific.

See Table 8.2 for a summary of the binary table format.

Table 8.2: **Xyce** binary rawfile format.

Issue	Comment
Title:	An arbitrary string describing the circuit.
Date:	A free-format date string.
Plotname:	A string describing the analysis type.
Flags:	A string describing the data type ( <i>real</i> or <i>complex</i> ).
No. Variables:	The number of variables.
No. Points:	The number of points.
Version: (optional)	The version of <b>Xyce</b> used to generate this output. By default the version is not output in the header. It can be output with the <code>.options outputoutputversioninrawfile=true</code> option.
Variables:	A newline followed by multiple lines, one for each variable, of the form [tab] <index> [tab] <name> [tab] <type>.
Binary:	Each real data point is stored contiguously in <code>sizeof(double)</code> byte blocks. Complex values are output as real and imaginary components in a block of size <code>2*sizeof(double)</code> byte blocks.

## 8.3 Special Notes

- Complex data points are only output under AC analysis.
- Commands and Options lines are not used.
- Binary header is formatted ASCII.
- **Xyce** can output an optional `Version` line in the header.

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