GSS User's Guide Ver 0.46.01

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Contents

1	Intr	roduction	2
	1.1	The format of input card	2
	1.2		3
	1.3		3
2	Glo	bal Specification	4
	2.1	SET	4
3	Mes	sh Generation	5
	3.1	Introduction	5
	3.2	Coordinate System	5
	3.3	· · · · · · · · · · · · · · · · · · ·	6
	3.4		7
	3.5		8
	3.6		9
	3.7	REGION	_
	٠		
	3.8	SEGMENT	
	3.9	REFINE	4
4	Dop	ping Profile 1	5
5	Vol	tage and Current Source 1	6
	5.1	Introduction	6
	5.2	ISOURCE	6
	5.3	VSOURCE	

1 INTRODUCTION 2

6	Bou	indary Condition	20
	6.1	BOUNDARY and CONTACT	20
	6.2	ATTACH	23
7	Phy	rsical Model Interface	24
8	Solv	ve Specification	25
	8.1	Introduction	25
	8.2	METHOD	25
	8.3	SOLVE	28
	8.4	AC Sweep Solver	30
	8.5	EM FEM Solver	31
	8.6	IV File Format	33
9	File	· I/O	34
		Introduction	34
	9.2	IMPORT and EXPORT	34
10	Pos	t Process	35
	10.1	Plot	35
		Probe	36
11	Con	avergence Problem	37
12	Mei	mory and CPU requirement	38
A	GN	U Free Documentation License	39

1 Introduction

1.1 The format of input card

Like PISCES and MEDICI, GSS takes its command cards from a user specified disk file. The input is read by GSS's build-in command parser. Each line is recognized as a particular statement, identified by the first word (named as keyword) on the card. The remaining parts of the line are the parameters of that keyword. The statement has the format as follow:

KEYWORD [parameters]

The words on a line are separated by blanks or tabs. If more than one line of input is necessary for a particular statement, it may be continued on subsequent lines by placing a backslash sign '\' as the last non-blank character on the current line. Parameters may be one of four types: float, integer, bool or string. The float point number supports C style double precision real number. The bool value can be True, On, False and Off. String value is made up of lower line, dot, blank, number and alpha characters. The string should not begin with number and quotation marks are only needed if it contains blank. At last, the length of string is limited to 31 characters. All the parameter specification has the same format as

1 INTRODUCTION 3

```
parameter_name = [number|integer|bool|string]
```

In the card descriptions, keywords and parameters are not case sensitive. But user input strings do, because file name may be specified by the string. Comments must begin with '#' and can be either an separated line or locate at the end of current statement.

1.2 The sequence of input deck

Most of the cards GSS used are sequence insensitive. The order of occurrence of cards is significant in only two cases. The mesh generation cards must have the right order, or it can't work properly. GSS will execute the 'driven' cards sequently. So the placement order of 'driven' cards will affect simulation result.

1.3 Statement Description Format

Syntax of Parameter Lists

The following special characters are used in the formatted parameter list:

```
Angle brackets < > - parameter type

Square brackets [ ] - optional group

Vertical bar | - alternate choice

Parentheses ( ) - group hierarchy

Braces { } - group hierarchy with high level
```

Value Types

Besides some string parameters which have fixed values, most of the parameters need a user defined value. A lower case letter in angle brackets represents a value of a given type. The following types of values are represented:

```
<n> - double precision numerical value
<i> - integer value
<b> - bool value
<s> - string value
```

2 Global Specification

2.1 SET

Description

Some global definitions such as the unit scale and environment temperature must be set before the initiation of GSS's build-in data. The SET command will do the definition.

Syntax

```
set Carrier=(p|n|pn)
set Z.Width=<n>
set LatticeTemp=<n>
set DopingScale=<n>
```

parameter Carrier	type string	default pn	unit -	description The Carrier parameter specifies whether single or dual carriers will be modeled during the simulation. But at present, GSS only supports dual carriers, so the parameter value must always be "pn"
Z.Width	number	1	$\mu\mathrm{m}$	Z.Width is needed by current calculation. Because GSS is a two-dimensional simulator, the length in Z direction must be given if GSS simulates transistor with external circuit.
LatticeTemp	number	300	K	LatticeTemp defines external temperature.
DopingScale	number	1e18	${ m cm^{-3}}$	DopingScale will effect GSS's inner unit scale procedure which shows great influence to the convergence of nonlinear solver. In most case, set this value to max(Nd,Na) is a good choice. But sometimes, a smaller value may be better.

```
set Carrier = pn  # specify carrier type.
set Z.Width = 2  # device width in Z dimension. Unit:um
set LatticeTemp = 3e2  # specify initial temperature of device. Unit:K
set DopingScale = 1e16  # set carrier scale reference value
```

3 Mesh Generation

3.1 Introduction

The early version of GSS was designed as a pure solver. It uses CGNS(CFD General Notation System) as semiconductor device model file. This file format provides the ability to store grid, solution data, material information, boundary condition and connectivity in a single, well-defined and easy-to-use form. More important, CGNS has been accepted and supported by most of the commercial CFD corporations. So users have various ways to create their models. For example, models can be created by SGFramework, converted from MEDICI TIF file by TIFTOOL (shipped with GSS) or generated by ICEMCFD, which is a commercial CFD pre-processor.

Until very recently, the PISCES like model description language had been introduced to GSS. The mesh generation arithmetic works as follows. First, GSS builds the rectangle skeleton mesh by the model description statements; Then, GSS employs Triangle (developed by Jonathan Richard Shewchuk) to form the triangulate mesh and output the mesh to an initial CGNS file. At last, GSS reads the CGNS file again, computes the doping profile and finishes the remaining calculations.

Triangle uses delaunay arithmetic, which forms a high quality isotropic mesh. At the same time, MEDICI uses quadtree arithmetic to generate its mesh, which often gives a regular mesh but the mesh quality may be poor near the irregular boundary.

3.2 Coordinate System

The mesh generator uses a Cartesian coordinate system, in which the top horizontal line has the maximal y coordinate and left vertical line has the minimal x coordinate.

Note: This setting is different from PISCES and its commercial versions like MEDICI and ATLAS.

3.3 MESH

This statement indicates the beginning of the mesh generator.

Syntax

parameter	\mathbf{type}	$\mathbf{default}$	\mathbf{unit}	description
Type	string	-	-	Type indicates which mesh generator is to be
				used. But at present it is useless since GSS
				only has one mesh generator.
ModelFile	string	-	-	ModelFile gives the name of temporary
				CGNS file.
Triangle	string	pzq30AD	-	Triangle passes parameters to Triangle
				code. The detailed description of this
				string can be found at Triangle's home page
				http://www.cs.cmu.edu/ quake/triangle.html.

Example

MESH Type=GSS ModelFile=pn.cgns Triangle="pzA"

3.4 XMESH and YMESH

The **XMESH** and **YMESH** cards specify the location of lines of nodes in a rectangular mesh. The original mesh can be modified by following mesh cards like **ELIMINATE** and **SPREAD**.

Syntax

\mathbf{type}	default	\mathbf{unit}	description
number	-	$\mu\mathrm{m}$	The distance of the grid section in x direction.
number	-	$\mu\mathrm{m}$	The distance of the grid section in y direction.
number	-	$\mu\mathrm{m}$	The x location of the left edge of the grid
			section. synonym: X.LEFT . The value of
			X.MIN will be set to right edge of the previ-
			ous grid section automatically.
number	-	$\mu\mathrm{m}$	The x location of the right edge of the grid
			section. synonym: X.RIGHT .
number	-	$\mu\mathrm{m}$	The y location of the bottom edge of the grid
			section. synonym: Y.BOTTOM.
number	-	$\mu\mathrm{m}$	The y location of the top edge of the grid
			section. synonym: Y.TOP . The value of
			Y.MAX will be set to bottom edge of the
			previous grid section automatically.
integer	1	-	The number of grid spaces in the grid section.
number	1.0	-	The ratio between the sizes of adjacent grid
			spaces in the grid section. RATIO should
			usually lie between 0.667 and 1.5.
number	-	$\mu\mathrm{m}$	The size of the grid space at the begin edge of
			the grid section.
number	-	$\mu\mathrm{m}$	The size of the grid space at the end edge of
			the grid section.
	number number number number number number number	number - number - number - number - number - number - number 1 number 1.0	number - μ m integer 1 - number 1.0 - μ m

XMESH	X.MIN=0.0	X.MAX=0.50	N.SPACES=8
YMESH	DEPTH=0.1	N.SPACES=8	RATIO=0.8
YMESH	DEPTH=0.1	N.SPACES=20	0
YMESH	DEPTH=0.6	H1=0.005 I	H2=0.050

3.5 ELIMINATE

The **ELIMINATE** statement eliminates mesh points along planes in a rectangular grid over a specified volume. This statement is useful for eliminating nodes in regions of the device structure where the grid is more dense than necessary. Points along every second line in the chosen direction within the chosen range are removed, except the first and last line. Successive eliminations of the same range remove points along every fourth line, eighth line, and so on.

Syntax

parameter	type	default	\mathbf{unit}	description
DIRECTION	string	-	-	Specifies that horizontal or vertical lines of nodes are eliminated.
X.MIN	number	XMIN	$\mu\mathrm{m}$	The minimum x location of the rectangular volume in which nodes are eliminated. synonym: X.LEFT .
X.MAX	number	XMAX	$\mu\mathrm{m}$	The maximum x location of the rectangular volume in which nodes are eliminated. synonym: X.RIGHT .
IX.MIN	integer	0	-	The minimum x node index of the rectangular volume in which nodes are eliminated. syn-
IX.MAX	integer	IXMAX-1	-	onym: IX.LEFT . The maximum x node index of the rectangular volume in which nodes are eliminated. syn-
Y.MIN	number	YMIN	$\mu\mathrm{m}$	onym: IX.RIGHT. The minimum y location of the rectangular volume in which nodes are eliminated. syn-
Y.MAX	number	YMAX	$\mu\mathrm{m}$	onym: Y.BOTTOM . The maximum y location of the rectangular volume in which nodes are eliminated. syn-
IY.MIN	integer	0	-	onym: Y.TOP . The minimum y node index of the rectangular volume in which nodes are eliminated. synonym: IY.TOP .
IY.MAX	integer	IYMAX-1	-	The maximum y node index of the rectangular volume in which nodes are eliminated. synonym: IY.BOTTOM.

ELIMINATE	Direction=COLUMNS	Y.TOP=-1.0
ELIMINATE	Direction=ROWS	IX.MAX=8

3.6 SPREAD

The **SPREAD** statement provides a way to adjust the y position of nodes along grid lines parallel to the x-axis in a rectangular mesh to follow surface and junction contours.

Syntax

parameter	\mathbf{type}	default	\mathbf{unit}	description
LOCATION	string	-	-	Specifies which side of the grid is distorted.
WIDTH	number	0.0	$\mu\mathrm{m}$	The width of the distorted region measured from the left or right edge of the structure.
UPPER	integer	0	-	The index of the upper y-grid line of the distorted region.
LOWER	integer	0	-	The index of the lower y-grid line of the distorted region.
ENCROACH	number	1.0	-	The factor which defines the abruptness of the transition between distorted and undis- torted grid. The transition region becomes more abrupt with smaller ENCROACH fac- tors. The minimum allowed value is 0.1.
Y.LOWER	number	-	$\mu\mathrm{m}$	The vertical location in the distorted region where the line specified by LOWER is moved. The grid line specified by UPPER does not move if this parameter is specified.
THICKNESS	number	-	$\mu\mathrm{m}$	The thickness of the distorted region. Specifying THICKNESS usually causes the positions of both the UPPER and LOWER grid lines to move.
VOL.RAT	number	0.44	-	The ratio of the displacement of the lower grid line to the net change in thickness. If VOL.RAT is 0, the location of the lower grid line does not move. If VOL.RAT is 1, the upper grid line does not move.
GRADING	number	1.0	-	The vertical grid spacing ratio in the distorted region between the y-grid lines specified with UPPER and LOWER The spacing grows or shrinks by GRADING in each interval between lines. GRADING should usually lie between 0.667 and 1.5.

SPREAD	Location=Left	Width=0.625	Upper=0	Lower=2	Thickness=0.1
SPREAD	Location=Right	Width=0.625	Upper=0	Lower=2	Thickness=0.1

3.7 REGION

The **REGION** statement defines the location of materials in the mesh. Currently, GSS supports following materials: null space including Vacuum and Air; semiconductor material including Si, Ge, GaAs, $Si_{1-x}Ge_x$, $Al_xGa_{1-x}As$ and $In_xGa_{1-x}As$; insulator material including SiO_2 and electrode region including Elec, Al and PolySi.

Syntax

Shape string Specifies the shape of the region. Can be Rectangle or Ellipse. Label string Specifies the identifier of this region, limited to 12 chars. Material string Specifies the material of the region. Material strings can be Vacuum, Air, Si, Ge, GaAs, SiGe, AlGaAs, InGaAs, SiO2, Elec, Al and PolySi. X.MOLE number 0.0 - The mole fraction to use in the region for compound materials. For graded compounds, X.MOLE represents
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be Vacuum, Air, Si, Ge, GaAs, SiGe, AlGaAs, InGaAs, SiO2, Elec, Al and PolySi. X.MOLE number 0.0 - The mole fraction to use in the region for compound materials. For graded compounds, X.MOLE represents
SiO2, Elec, Al and PolySi. X.MOLE number 0.0 - The mole fraction to use in the region for compound materials. For graded compounds, X.MOLE represents
X.MOLE number 0.0 - The mole fraction to use in the region for compound materials. For graded compounds, X.MOLE represents
the initial male fraction at the left ten or front adm of
the initial mole fraction at the left, top, or front edge of
the region depending on whether X.Linear, or Y.Linear, respectively, is specified.
MOLE.SLOPE number 0.0 μm^{-1} The slope of the mole fraction for graded compounds.
If this parameter is used, the mole fraction has a value
of X.MOLE at the left, top or front edge of the region
and a value of $\mathbf{X.MOLE}$ + width * $\mathbf{X.SLOPE}$ at the
right, bottom or back edge of the region, where width
is the width or depth of the region.
MOLE.END number 0.0 - The mole fraction for graded compounds at the right,
bottom, or backedge of the region depending on whether X.Linear, or Y.Linear, respectively, is specified.
MOLE.GRAD string Y.Linear - Specifies that the mole fraction grading is in the x or y
direction.
X.MIN number XMIN μ m The minimum x location of the region. synonym:
X.LEFT.
X.MAX number XMAX μ m The maximum x location of the region. synonym:
X.RIGHT.
IX.MIN integer 0 - The minimum x node index of the region. synonym:
IX.LEFT.
IX.MAX integer IXMAX-1 - The maximum x node index of the region. synonym: IX.RIGHT.
Y.MIN number YMIN μ m The minimum y location of the region. synonym:

Y.BOTTOM.

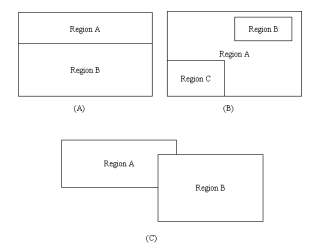
Y.MAX	number	YMAX	$\mu\mathrm{m}$	The maximum y location of the region. synonym:
				Y.TOP.
IY.MIN	integer	0	-	The minimum y node index of the region. synonym:
				IY.TOP.
IY.MAX	integer	IYMAX-1	-	The maximum y node index of the region. synonym:
				IY.BOTTOM.
CentreX	number	0.0	$\mu\mathrm{m}$	The x location of the center of ellipse.
CentreY	number	0.0	$\mu\mathrm{m}$	The y location of the center of ellipse.
MajorRadii	number	1.0	$\mu\mathrm{m}$	The length of the major radii of ellipse.
MinorRadii	number	MajorRadii	$\mu\mathrm{m}$	The length of the minor radii of ellipse.
Theta	number	0.0	degree	The angle of the first division point located on the
				boundary of ellipse region.
Division	integer	12	-	The number of points which divide the boundary of el-
				lipse into small segments.

Example

REGION	Label=Si1	Material=Si	Y.TOP= 0.000	Y.BOTTOM=-C	0.100
REGION	Label=SiGe1	Material=SiGe	Y.TOP=-0.100	Y.BOTTOM=-C).125 \
	X.MOLE=0.0	Mole.End=0.2			
REGION	Label=Hole	Material=SiO2	Shape=Ellipse	CentreX=2.0	<pre>CentreY=-0.5 \</pre>
	Division=24	MajorRadii=0.3	B MinorRadii=0.	.3	

Hint

Several regions can be defined one by one. But users should be careful that regions can't get cross each other. The situations showed by Fig1 (A) and (B) are allowed, but (C) will break the mesh generator of GSS. The ellipse region is used for photon crystal simulation. By choosing different division number, GSS can build triangle, rectangle, hexagon as well as ellipse (circle). Fig2 shows different shapes of polygons build by ellipse.





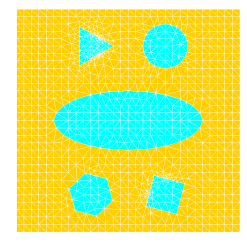


Figure 2: Define shapes of ellipse region

3.8 SEGMENT

Segment is a group of boundary edges which have the same attribute. This statement specifies the label of a special segment. User can assign the segment with a special boundary type by **BOUNDARY** statement.

Syntax

```
SEGMENT Label=<s> { Location=<s> | ( Direction=<s> X=<n> | Y=<n> ) }
        [ {X.MIN=<n> | IX.MIN=<n>} ] [ {X.MAX=<n> | IX.MAX=<n>} ]
        [ {Y.MIN=<n> | IY.MIN=<n>} ] [ {Y.MAX=<n> | IY.MAX=<n>} ]
```

parameter	\mathbf{type}	default	\mathbf{unit}	description
Label	string	-	-	Specifies the identifier of this segment, limited to 31 chars.
Location	string	-	-	Specifies which side the segment lies along. Allowed: TOP, BOTTOM, LEFT or RIGHT.
Direction	string	-	-	Specifies the dimensional orientation of the segment. Allowed: Horizontal or Vertical.
X	number	0.0	$\mu\mathrm{m}$	Specifies the X coordinate of the vertical segment.
Y	number	0.0	$\mu\mathrm{m}$	Specifies the Y coordinate of the horizontal segment.
X.MIN	number	XMIN	$\mu\mathrm{m}$	The minimum x location of the segment. synonym: X.LEFT .
X.MAX	number	XMAX	$\mu\mathrm{m}$	The maximum x location of the segment. synonym: X.RIGHT .
IX.MIN	integer	0	-	The minimum x node index of the segment. synonym: IX.LEFT .
IX.MAX	integer	IXMAX-1	-	The maximum x node index of the segment. synonym: IX.RIGHT .
Y.MIN	number	YMIN	$\mu\mathrm{m}$	The minimum y location of the segment. synonym: Y.BOTTOM .
Y.MAX	number	YMAX	$\mu\mathrm{m}$	The maximum y location of the segment. synonym: Y.TOP .
IY.MIN	integer	0	-	The minimum y node index of the segment. synonym: IY.TOP .
IY.MAX	integer	IYMAX-1	-	The maximum y node index of the segment. synonym: IY.BOTTOM .

SEGMENT	Label=Anode	Direction=Horizontal X.MIN=0.0 X.MAX=1.0 Y=0.0
SEGMENT	Label=Cathode	Direction=Horizontal X.MIN=0.0 X.MAX=3.0 Y=-3.0
SEGMENT	Label=Anode	Location=TOP X.MIN=0.0 X.MAX=1.0
SEGMENT	Label=Cathode	Location=BOTTOM

Hint

Here, I have to mention the naming principle of segments. Beside labeled segments, the interface edges between two regions will be assigned by IF_name1_to_name2 in which the name1 and name2 is the labels of the two regions by alpha order. The remain edges of a region will be assigned by name_Neumann and the name is the label of the region.

One can define a segment for probing data. Please refer to **PROBE** statement. This kind of segment should be placed inside a region. Equally, NO intersection to any other segment.

3.9 REFINE

The **REFINE** statement allows refinement of a coarse mesh.

Syntax

parameter	\mathbf{type}	default	\mathbf{unit}	description
Variable	string	-	-	Specifies that the grid refinement is based on
				the potential or doping quantity.
Measure	string	Linear	-	Specifies that refinement is based on the orig-
				inal value or logarithm of the specified quan-
				tity.
Dispersion	number	3.0	-	The numerical criterion for refining a triangle.
				If the specified quantity differs by more than
				this parameter at the nodes of a triangle, the
				triangle is divided.
DivisionRatio	number	0.25	-	The area of divided triangle over area of orig-
				inal triangle. The default value suggests Tri-
				angle code divide one triangle into 4 small tri-
				angles. It is a suggestion value, Triangle code
				will adjust it for mesh quality reason.
Triangle	string	praq30Dz	-	Passes parameters to Triangle code.

Example

REFINE Variable=Doping Measure=SignedLog Dispersion=1
REFINE Variable=Potential Measure=Linear Dispersion=0.1

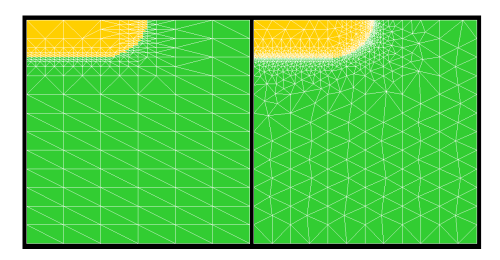


Figure 3: Mesh refinement for a PN diode.

4 DOPING PROFILE 15

4 Doping Profile

The **PROFILE** statement defines profiles for impurities to be used in the device structure. At present, GSS supports analytic profiles such as uniform, gauss distribution in both x-y directions and error function distribution in x direction while gauss distribution in y direction.

Syntax

```
PROFILE { Type=Uniform |
Type=Gauss [YCHAR=<n> | Y.Junction=<n>] [XCHAR=<n>] |
Type=ErrorFunc [YCHAR=<n>] [XCHAR=<n>] }
Ion=(Donor|Acceptor) { N.Peak=<n> | Dose=<n> }
[ X.MIN=<n> ] [ X.MAX=<n> ] [ Y.MIN=<n> ] [ Y.MAX=<n> ]
```

parameter	\mathbf{type}	default	\mathbf{unit}	description
Type	string	-	-	Specifies that the profile has a uniform, gauss
				or error function distribution.
Ion	string	-	-	Specifies the impurity ionization.
N.Peak	number	0.0	cm^{-3}	The peak impurity concentration for an impurity profile.
Dose	number	0.0	cm^{-2}	The dose of the impurity profile assuming a
				full Gaussian distribution.
X.MIN	number	0.0	$\mu\mathrm{m}$	The minimum x location of the doping profile.
				synonym: X.LEFT .
X.MAX	number	XMIN	$\mu\mathrm{m}$	The maximum x location of the doping profile.
				synonym: X.RIGHT.
Y.MIN	number	YMAX	$\mu\mathrm{m}$	The minimum y location of the doping profile.
				synonym: Y.BOTTOM .
Y.MAX	number	0.0	$\mu\mathrm{m}$	The maximum y location of the doping profile.
				synonym: Y.TOP.
YCHAR	number	0.25	$\mu\mathrm{m}$	The y characteristic length of the profile out-
TT 077 1 D				side the range of $\mathbf{Y.MIN} < \mathbf{y} < \mathbf{Y.MAX}$.
XCHAR	number	0.25	$\mu\mathrm{m}$	The x characteristic length of the profile out-
77.7	,	0.0		side the range of $\mathbf{X}.\mathbf{MIN} < \mathbf{x} < \mathbf{X}.\mathbf{MAX}$.
Y.Junction	number	0.0	$\mu\mathrm{m}$	The y location under the center of the pro-
				file where the magnitude of the profile being
				added equals the magnitude of the background
				profile.

```
PROFILE Type=Uniform Ion=Donor N.PEAK=1E15 \
X.MIN=0.0 X.MAX=3.0 Y.TOP=0.0 Y.BOTTOM=-3.0

PROFILE Type=Gauss Ion=Acceptor N.PEAK=1E18 X.CHAR=0.2 Y.JUNCTION=-0.5 \
X.MIN=0.0 X.MAX=0.7 Y.TOP=0.0 Y.BOTTOM=0.0

PROFILE Type=ErrorFunc Ion=Acceptor N.PEAK=2E17 X.CHAR=0.25 Y.CHAR=0.25 \
X.MIN=0.5 X.MAX=1.0 Y.TOP=0.0 Y.BOTTOM=0.0
```

5 Voltage and Current Source

5.1 Introduction

For simulation the transient response of device, GSS supports several types of voltage and current source. The original models of these sources come from SPICE, a famous circuit simulation program. Several sources may be defined in one disk file. And the placement of these definitions are not critical. The sources can be assigned to electrode by **ATTACH** statement when needed.

5.2 ISOURCE

Syntax

parameter Type	type string	default -	unit -	description This parameter declares which type of current
V 1				source is defined here. Only four types of current source listed as previous are supported at present.
ID	string	-	-	A unique string which identifies the current
				source.
Tdelay	number	0	\mathbf{s}	A proper delay time before the activation of
				this current source.
Iconst	number	0	mA	The current of the IDC.
Iamp	number	0	mA	The amplitude current of the ISIN.
Freq	number	0	$_{\mathrm{Hz}}$	The frequency of the ISIN.
TRC	number	0	\mathbf{S}	The rise time constant of the IEXP.
TFD	number	0	\mathbf{S}	The fall delay time of the IEXP.
TFC	number	0	\mathbf{S}	The fall time constant of the IEXP.
Tr	number	0	\mathbf{S}	The raise edge of the IPULSE.
Tf	number	0	\mathbf{S}	The fall edge of the IPULSE.
Pw	number	0	\mathbf{S}	The pulse with of the IPULSE.
\Pr	number	0	\mathbf{S}	The period of the IPULSE.
Ilo	number	0	mA	The low current for both IEXP and IPULSE.
Ihi	number	0	mA	The high current for both IEXP and IPULSE.
DLL	string	-	-	The name of dynamic library file.
Func	string	-	-	The name of the function loaded from dynamic library file.

17

```
isource Type=IDC ID=I1 Tdelay=0 Iconst=5
isource Type=ISIN ID=I2 Tdelay=0 Iamp=0.1 Freq=1e6
isource Type=IEXP ID=I3 Tdelay=0 TRC=1E-6 TFD=3E-6 TFC=1E-6 Ilo=0 Ihi=1
isource Type=IPULSE ID=I4 Tdelay=0 Tr=1E-9 Tf=1E-9 Pw=5E-6 Pr=1E-5 Ilo=0 Ihi=1
```

5.3 VSOURCE

Syntax

parameter	\mathbf{type}	$\mathbf{default}$	\mathbf{unit}	description
Type	string	-	-	This parameter declares which type of voltage
				source is defined here. Only four types of volt-
				age source listed as previous are supported at
				present.
ID	string	-	-	A unique string which identifies the voltage
				source.
Tdelay	number	0	\mathbf{S}	A proper delay time before the activation of
				this voltage source.
Vconst	number	0	V	The voltage of the VDC.
Vamp	number	0	V	The amplitude voltage of the VSIN.
Freq	number	0	$_{\mathrm{Hz}}$	The frequency of the VSIN.
Alpha	number	0	-	The exponential attenuation parameter of the
				VSIN.
TRC	number	0	S	The rise time constant of the VEXP.
TFD	number	0	\mathbf{s}	The fall delay time of the VEXP.
TFC	number	0	\mathbf{s}	The fall time constant of the VEXP.
Tr	number	0	\mathbf{S}	The raise edge of the VPULSE.
Tf	number	0	\mathbf{s}	The fall edge of the VPULSE.
Pw	number	0	\mathbf{s}	The pulse with of the VPULSE.
\Pr	number	0	\mathbf{S}	The period of the VPULSE.
Vlo	number	0	V	The low voltage for both VEXP and VPULSE.
Vhi	number	0	V	The high voltage for both VEXP and
				VPULSE.
DLL	string	-	-	The name of dynamic library file.
Func	string	-	-	The name of the function loaded from dy-
				namic library file.

```
vsource Type=VDC    ID=GND    Tdelay=0 Vconst=0
vsource Type=VDC    ID=VCC    Tdelay=0 Vconst=5
vsource Type=VSIN    ID=Vs         Tdelay=1e-6 Vamp=0.1 Freq=1e6
vsource Type=VEXP    ID=V1         Tdelay=0 TRC=1e-6 TFD=1e-6 TFC=1e-6 Vlo=0 Vhi=1
vsource Type=VPULSE ID=V2         Tdelay=0 Tr=1e-9 Tf=1e-9 Pw=5e-6 Pr=1e-5 Vlo=0 Vhi=1
vsource Type=VSHELL ID=VGauss    DLL=foo.so Func=vsrc_gauss
```

Hint

GSS supports user defined voltage and current source by loading shared object (.so) file. The file which contains a user defined voltage source should have the function as follow. GSS will pass the argument time in the unit of second to the function $vsrc_name$ and get voltage value in the unit of volt. The current source function is almost the same except the unit of current is mA.

```
double vsrc_name(double time)
{
    /* calculate the voltage amplitude */
    return vsrc_amplitude;
}
double isrc_name(double time)
{
    /* calculate the current amplitude */
    return isrc_amplitude;
}
The c code should be linked with -shared and -fPIC option as:
    gcc -shared -fPIC -o foo.so foo.c -lm
The foo.so file should be put in the same directory as input file.
```

6 Boundary Condition

6.1 BOUNDARY and CONTACT

The **BOUNDARY** statement sets boundary information to representing segments which defined by mesh generator or read from CGNS file.

GSS now fully support electrode region (the material of this region may be metal or poly-Si). One should use **CONTACT** statement to specify the electrode type of this region(s).

Syntax

```
BOUNDARY Type=OhmicContact
                                   ID=<s>
                                           [ Res=<n> ] [ Cap=<n> ] [ Ind=<n> ]
         [ Heat.Transfer=<n> ] [EXT.Temp=<n> ] [ConnectTo=<s>]
BOUNDARY Type=SchottkyContact
                                  ID=<s> [ Res=<n> ] [ Cap=<n> ] [ Ind=<n> ]
         WorkFunction=<n> [ Heat.Transfer=<n> ] [EXT.Temp=<n> ]
BOUNDARY Type=GateContact
                                  ID=<s> WorkFunction=<n>
         [ Res=<n> ] [ Cap=<n> ] [ Ind=<n> ]
         [ Heat.Transfer=<n> ] [EXT.Temp=<n> ]
                                  ID=<s> WorkFunction=<n> [ QF=<n> ]
BOUNDARY Type=InsulatorContact
         [ Res=<n> ] [ Cap=<n> ] [ Ind=<n> ]
         Thickness=<n> Eps=<n> [ Heat.Transfer=<n> ] [EXT.Temp=<n> ]
BOUNDARY Type=InsulatorInterface ID=<s> [ QF=<n> ]
BOUNDARY Type=Heterojunction
                                  ID=<s> [QF=<n>]
BOUNDARY Type=NeumannBoundary
                                  ID=<s> [ Heat.Transfer=<n> ] [EXT.Temp=<n> ]
CONTACT Type=OhmicContact
                                  ID=<s>
         [ Res=\langle n \rangle ] [ Cap=\langle n \rangle ] [ Ind=\langle n \rangle ] [ConnectTo=\langle s \rangle]
         [ Heat.Transfer=<n> ] [EXT.Temp=<n> ]
CONTACT Type=SchottkyContact
         [ Res=<n> ] [ Cap=<n> ] [ Ind=<n> ]
         WorkFunction=<n> [ Heat.Transfer=<n> ] [EXT.Temp=<n> ]
        Type=GateContact
CONTACT
                                  ID=<s> WorkFunction=<n>
         [ Res=<n> ] [ Cap=<n> ] [ Ind=<n> ]
         [ Heat.Transfer=<n> ] [EXT.Temp=<n> ]
                                  ID=<s> [QF=<n>]
CONTACT
        Type=FloatMetal
```

parameter	\mathbf{type}	$\operatorname{default}$	${f unit}$	description
Type	string	-	-	This parameter declares which type of bound-
				ary condition is defined here.
ID	string	-	-	A unique string which identifies the corre-
				sponding segment.
Res	number	0	Ω	The lumped resistance for the electrode.
Cap	number	0	\mathbf{F}	The lumped capacitance for the electrode.
Ind	number	0	Н	The lumped inductance for the electrode.
ConnectTo	string	-	-	Specifies the ID of an ohmic electrode which
				connect to this ohmic electrode. Useful for
				CMOS structure.

WorkFunction	number	4.7	V	The workfunction of the Schottky contact or
				gate material.
$_{ m QF}$	number	0	$\mathrm{C}\cdot\mathrm{cm}^{-2}$	For InsulatorContact and InsulatorInter-
				face bc: The surface charge density of
				semiconductor-insulator interface.
$_{ m QF}$	number	0	$\mathrm{C}\cdot\mathrm{cm}^{-2}$	For Heterojunction bc: The surface charge
				density of heterojunction.
$_{ m QF}$	number	0	$\mathrm{C}\cdot\mu\mathrm{m}^{-1}$	For FloatMetal bc: The free charge per micron
				in Z dimension.
Thickness	number	2e-7	$^{ m cm}$	The thickness of SiO_2 layer.
Eps	number	3.9	-	The relative permittivity of SiO_2 layer.
Heat.Transfer	number	1e3	$W/(cm \cdot K)$	The heat transfer rate of boundary.
EXT.Temp	$_{ m number}$	LatticeTemp	K	The external temperature.

Example

BOUNDARY	Type=InsulatorContract	ID=SiSiO2	Res=0	Cap=0	Ind=0	\
	Thickness=1e-6 Eps=3.9 W	VorkFunction=4	1.7 QF=	=0		
BOUNDARY	Type=InsulatorInterface	ID=IFACE	QF=0			
BOUNDARY	Type=GateContract	ID=GATE	Res=0	Cap=0	Ind=0	WorkFunction=4.7
BOUNDARY	Type=NeumannBoundary	ID=WALL	Heat. T	Transfe	er=O EX	T.Temp=300
BOUNDARY	Type=SchottkyContract	ID=sgate	Res=0	Cap=0	Ind=0	VBarrier=0.8
BOUNDARY	Type=OhmicContract	ID=OMANODE	Res=0	Cap=0	Ind=0	
BOUNDARY	Type=OhmicContract	ID=OMCATHODE	Res=0	Cap=0	Ind=0	

Hint

Four "electrode" boundary conditions are supported by GSS. The names are ended with "Contact". The OhmicContact and SchottkyContact electrodes have current flow in both steady state and transient situations. While GateContact and InsulatorContact(a simplified MOSFET Gate boundary condition) only have displacement current in transient situation.

GSS supports five interfaces which can be set automatically: semiconductor-insulator interface(InsulatorInterface), semiconductor-electrode interface(set to OhmicContract as default), interface between different semiconductor material(Heterojunction) and interface between same semiconductor material(Homojunction). These boundaries can be set automatically by GSS if user didn't set them explicitly. However, the electrode-insulator interface, may have several situations: Gate to Oxide interface, FloatMetal to Oxide interface or Source/Drain electrode to Oxide interface. As a result, this interface can only be set correctly when electrode type is known. Please refer to the following CONTACT statement.

GSS can build region with metal or poly-Si material to form an electrode. Which means, i.e. for OhmicContact bc, one can simply specify a segment as Ohmic bc or build an electrode region as Ohmic electrode. Since Version 0.45.03, GSS considers electrode region, semiconductor region and insulator region during calculation. As a result, GSS added **CONTACT** statement for fast boundaries specification of electrode region. At present, GSS support electrode with the type of Ohmic, Schottky, Gate and FloatMetal. All the electrode should be specified explicitly and GSS will set corresponding boundaries automatically.

22

The "ID" parameter of ${\bf BOUNDARY}$ statement is limited to segment label. And The "ID" parameter of ${\bf CONTACT}$ statement is limited to region name.

The NeumannBoundary, which is the default boundary type for all the non-interface segments, can also be set automatically.

6.2 ATTACH

This statement is used to add voltage or current sources to the electrode boundary. The statement first clears all the sources connected to the specified electrode and then adds source(s) defined by VApp or IApp parameter. If two or more sources are attached to the same electrode, the total effect is the summation of all sources. However, the sources attached to one electrode must have the same type.

Syntax

```
ATTACH Electrode=<s> Type=Voltage VApp=<s> [VApp=<s> ...]
ATTACH Electrode=<s> Type=Current IApp=<s> [IApp=<s> ...]
```

parameter	$_{ m type}$	default	\mathbf{unit}	description
Electrode	string	-	-	Specifies which electrode boundary is to be at-
				tached with one or more sources.
Type	string	Voltage	-	The sources are voltage or current type.
VApp	string	-	-	Specifies the ID of voltage source which is to
				be attached to this electrode.
IApp	string	-	-	Specifies the ID of current source which is to
				be attached to this electrode.

Example

A'I"I'ACH	Electrode=Collector	VApp=VCC		
ATTACH	Electrode=Emitter	VApp=GND		
ATTACH	Electrode=Base	VApp=Vb VApp	=Vs	
ATTACH	Electrode=Base	Type=Current	IApp=Ib	IApp=Is

Hint

If electrode is attached with voltage source(s), the R, C and L defined by **BOUNDARY** statement will affect later simulation. But solver will ignore those lumped elements with the electrode which stimulated by current source(s). Please refer to Fig 4.

The positive direction of current is flow into the electrode.

Only Ohmic and Schottky electrodes can be attached by current source(s).

If no source attached explicitly, the electrode is set to be attached to ground.

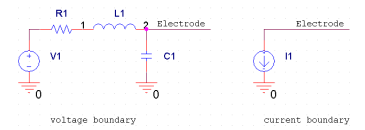


Figure 4: Voltage and current boundary.

7 Physical Model Interface

GSS use a dynamic mechanician to support various materials and physical models. Each material has a dynamic load library (.so) which contains its physical parameters. User can modify the parameters which can be found at \$(GSS_DIR)/src/material and recompile it. Experts can even offer their own physical model files.

At present, GSS has a **PMIS** statement for choosing different mobility models and impact ionization models.

Syntax

```
PMIS Region=<s> Mobility=<s> II.Model=<s>
```

parameter	\mathbf{type}	$\mathbf{default}$	\mathbf{unit}	description
Region	string		-	Specifies the semiconductor region which use
				the following physical model.
Mobility	string	Analytic	-	The mobility model name.
II.Model	string	Default	-	The impact ionization model name.

Example

```
PMIS Region=Si Mobility=Philips
PMIS Region=Si Mobility=Lucent II.Model=Valdinoci
```

Hint

One can set different physical models to individual region.

GSS has implemented Analytic, Philips and Lucent mobility model for all the supported material. The Analytic and Philips mobility model only takes parallel field effect and they can be used within all the four solvers. The author suggest to use these models for bipolar device simulations. The Lucent mobility model, which considers parallel and transverse electrical field, is an accurate model for MOS structure. But it should work with DDML1E/DDML2E solvers in which transverse electrical field is calculated. The Lombardi and HP (Hewlett-Packard) mobility model only validate for Silicon. These two mobility models include parallel and transverse electrical field corrections and can be used for MOSFET simulation. The Hypertang mobility model only validate for GaAs. It is reported that this model can avoid unrealistic drain current oscillation when applied to the simulation of GaAs MESFET.

The impact ionization model is still very limited in GSS. Only Valdinoci model for silicon is valid at present.

8 Solve Specification

8.1 Introduction

These statements instruct GSS core to perform user specified solution(s).

8.2 METHOD

The **METHOD** statement sets the solver and the parameters of the solver. At present, GSS 0.4x has basic DDM solver(DDML1E), lattice temperature corrected DDM solver(DDML2E) and EBML3E solver which base on energy balance model.

Syntax

```
METHOD Type=(DDML1E|DDMLE2|EBML3E|QDDML1E) Scheme=Newton
       HighFieldMobility=(On|Off)
                                    EJModel=(On|Off)
       ImpactIonization=(On|Off)
                                    II.Type=(EdotJ|EVector|ESide|GradQf)
       BandBandTunneling=(On|Off)
       Fermi=(On|Off)
       NS=(Basic|LineSearch|TrustRegion)
       LS=(SuperLU|LU|CGS|BICG|BCGS|GMRES|TFQMR)
       Damping=(BankRose|Potential|No)
       MaxIteration=<i> relative.tol=<n>
       possion.tol=<n> elec.continuty.tol=<n> hole.continuty.tol=<n>
       elec.energy.tol=<n> hole.energy.tol=<n>
                                                latt.temp.tol=<n>
       electrode.tol=<n> toler.relax=<n>
       QNFactor=<n> QPFactor=<n>
```

parameter	\mathbf{type}	$\operatorname{default}$	${f unit}$	description
Type	string	DDML1	-	Specifies the solver.
Scheme	string	Newton	-	At present, GSS only supports Newton's full
HighFieldMobility	bool	On	-	iterative scheme. Specifies if high field mobility should be used.
				GSS set this flag to OFF for equilibrium state.
EJModel	bool	Off	-	Specifies if EdotJ and EcrossJ should be use
				to calculate high field mobility. GSS will use
				a simpler model when this flag is set to OFF.
ImpactIonization	bool	Off	-	Specifies if impact ionization should be con-
				sidered.
II.Type	string	GradQf	-	Specifies the implement model of impact ion-
				ization.
BandBandTunneling	string	Off	-	Specifies if band to band tunneling should be
				considered.
Fermi	bool	Off	-	Specifies if Fermi-Dirac statistics should be
				considered.
NS	string	LineSearch	-	Specifies the nonlinear solver.
LS	string	GMRES	-	Specifies the linear solver.
Damping	string	No	-	Load a Newton damping method for Line-
				Search or Basic Newton nonlinear solver.

MaxIteration	integer	30	-	The max number of iteration nonlinear solver will try. But for equilibrium state calculation, the max allowed iteration number is 10 times
1 4: 4 1	1	1 F		more than this value.
relative.tol	number	1e-5	-	When relative error of solution variable less than this value, solution is considered con- verged.
possion.tol	number	1e-26	$\mathrm{C}\cdot\mu\mathrm{m}^{-1}$	The absolute converged criteria for the Poisson equation.
elec.continuty.tol	number	5e-18	$A \cdot \mu m^{-1}$	The absolute converged criteria for the electron continuity equation.
hole.continuty.tol	number	5e-18	$A \cdot \mu m^{-1}$	The absolute converged criteria for the hole continuity equation.
elec.energy.tol	number	1e-18	$\mathbf{W} \cdot \mu \mathbf{m}^{-1}$	The absolute converged criteria for the elec-
				tron energy balance equation.
hole.energy.tol	number	1e-18	$W \cdot \mu m^{-1}$	The absolute converged criteria for the hole energy balance equation.
latt.temp.tol	number	1e-11	$W \cdot \mu m^{-1}$	The absolute converged criteria for the lattice heat equation equation.
electrode.tol	number	1e-9	V	The absolute converged criteria for the electrode bias equation.
toler.relax	number	1e4	-	When relative error is used as converged criteria, the equation norm should satisfy the absolute converged criteria with a relaxation of
				this value.
ONE	1	1.0		
QNFactor	number	1.0	-	The damping quantity of electron quantum potential.
QPFactor	number	1.0	-	The damping quantity of hole quantum potential.

Example

METHOD	Type=DDML1E	Scheme=Newton	NS=LineSearch LS=GMRES
METHOD	Type=DDML1E	Scheme=Newton	NS=TrustRegion LS=LU
METHOD	Type=DDML2E	Scheme=Newton	NS=Basic LS=TFGMR Damping=Potential

Hint

All the DDML1E/DDML2E/EBML3E/QDDML1E solvers support parallel and transverse electrical field dependent mobility.

Lattice temperature equation is considered by DDML2E solver. The EBML3E solver is based on advanced energy balance method. The QDDML1E is a density-gradient solver which consists of quantum correction to classical model.

The carrier generation by impact ionization and band tunneling is really difficult for calculation. However, DDML1E/DDML2E solvers are carefully designed for impact ionization and band band tunneling calculation, i.e. diode reverse breakdown simulation. Usually, the temperature can't keep unchanged if carrier generation takes place. As a result, DDML2E solver is highly recommend for these types of situations. At present, EBML3E and QDDML1E solver don't support impact ionization.

Fermi statistics is only supported by DDML1E and DDML2E solvers.

LineSearch and TrustRegion accelerating methods work well when initial value a bit far from real solution, e.g. first time computing. Basic Newton method should only be used when initial value is near the true solution, e.g. dc sweep and transient calculation.

Each nonlinear solver should have a inner linear solver. To choose a suitable linear solver may help the convergence. The performance of LineSearch and Basic Newton methods is good when Krylov subspace linear solvers (CGS, BICG, BCGS, GMRES and TFQMR) are employed. However, the TrustRegion method prefers LU factorization linear solver to Krylov subspace linear solvers.

Newton Damping is a useful tool for helping convergence, especially for the Basic Newton method.

QNFactor and QPFactor is used to enforce the convergence property of QDDML1E solver. Since quantum solution differs much from classical solution near Si/SiO2 interface, setting these two factors with small value i.e. 1e-4 and varying it gradually to 1.0, with each step the solution can get convergence. At last, the value of QXFactor of 1.0 means that the quantum model is fully turned on and applied.

The parameters of **METHOD** statement will not be affected by previous **METHOD** statement.

The convergence is considered to be achieved when either the X norm or the function residual norm falls below certain tolerance. When every function's residual norm falls small than certain tolerance, the absolute convergence is achieved. For X norm criteria, it should fall below **relative.tol** and every function residual norm should fit the relaxed (with the relaxation value of **toler.relax**) absolute converged criteria.

8.3 SOLVE

The **SOLVE** statement instructs GSS to perform a solution for one or more specified bias points.

Syntax

```
SOLVE Type=EQUILIBRIUM

SOLVE Type=STEADYSTATE

SOLVE Type=DCSWEEP VScan=<s> [VScan=<s> ...] [IVRecord=<s> ...]

[IVFile=<s>] VStart=<s> VStep=<s> VStop=<n>

SOLVE Type=DCSWEEP IScan=<s> [IVRecord=<s> ...]

[IVFile=<s>] IStart=<s> IStep=<s> IStop=<n>

SOLVE Type=TRANSIENT ODE.Formula=(BDF1|BDF2) [IVRecord=<s> ...]

[IVFile=<s>] TStart=<n> TStep=<n> TStop=<n>
AutoStep=<b> Predict=<b>
```

parameter	\mathbf{type}	$\mathbf{default}$	\mathbf{unit}	description	
Type	string	-	-	Specifies the Solve condition.	
VScan	string	-	-	Specifies the voltage variational electrode	
				boundary for DCSWEEP.	
VStart	number	-	V	The initial voltage for DC sweep.	
VStep	number	-	V	The voltage step size of DC sweep.	
VStop	number	-	V	The finish voltage for DC sweep.	
IScan	string	-	-	Specifies the current variational electrode	
				boundary for DCSWEEP.	
IStart	number	-	mA	The initial current for DC sweep.	
IStep	number	-	mA	The current step size of DC sweep.	
IStop	number	-	mA	The finish current for DC sweep.	
IV.Record	string	-	-	Specifies which electrode's IV data should be	
				recorded. User can define serval electrodes	
				here.	
IV.File	string	-	-	Specifies the file which contains the IV data.	
ODE.Formula	string	BDF2	-	Specifies the time march scheme for solving	
				the time-domain ordinary differential equa-	
				tion.	
TStart	number	-	\mathbf{s}	The initial time for transient calculation.	
TStep	number	-	\mathbf{s}	The time step size of transient calculation.	
TStop	number	-	\mathbf{s}	The finish time for transient calculation.	
AutoStep	bool	-	True	Use automatically time step control based on	
				LTE.	
Predict	bool	-	True	Predict initial value for next time step.	

SOLVE	Type=EQUILIBRIUM	[
SOLVE	Type=DCSWEEP	VScan=Anode	IVRecord=Anode	IVRecord=Cathode \
	IVFile=ivfp.txt	VStart=0 VSter	=1e-2 VStop=0.6	

SOLVE	Type=DCSWEEP	IScan=Anode	IVRecord=Anode	IVRecord=Cathode	\
	IVFile=ivfp2.txt	IStart=0.02 IS	tep=1e-2 IStop=	1	
SOLVE	Type=TRANSIENT	IVRecord=Anode	IVFile=iv.txt `	\	
	TStart=0 TSten=1	=10 TSton=3e-	8		

Hint

For equilibrium state calculation, all the electrodes are set to ground.

You can't do a DC sweep with current scan to GateContact and InsulatorContact.

When STEADYSTATE or DCSWEEP solve is performed, transient 0 value of the voltage(current) source will be used as the bias of each electrode.

One can do voltage DCSWEEP with multi-electrode by specifying two or more VScan parameter. The voltage will be assigned to each electrode during the simulation. This function is useful for Double Gate MOS simulation.

The step size for DCSWEEP calculation will automatically reduce to half size if last step diverged. Then it will be multiplied by 1.1 on each step until it reaches original step size.

TRANSIENT simulation now use automatically time step control based on LTE (local truncation error).

8.4 AC Sweep Solver

In addition to DC steady state and transient analysis, GSS now allows AC small-signal analysis as a post-processing step after a DC solution.

Syntax

\mathbf{type}	$\mathbf{default}$	\mathbf{unit}	description
string	-	-	Specifies the electrode for ACSWEEP.
number	1e6	$_{\mathrm{Hz}}$	The initial frequency for AC sweep.
number	1.1	-	The multiplicative factor for incrementing fre-
number number	1e9 0.0026	Hz V	quency. The finish frequency for AC sweep. The magnitude of the applied small-signal bias.
	string number number	string - number 1e6 number 1.1 number 1e9	string number 1e6 Hz number 1.1 - number 1e9 Hz

Hint

This solver shared Jacobian Matrix with DDML1E solver. Which means one should call it directly after DDML1E, keeping all the parameters unchanged for **METHOD** statement. If a previous computed result is imported, call DDML1E to do a steady-state calculation again and run DDML1AC later.

The convergence may be difficult if frequency is very high, i.e. nearly cut off frequency, because of the poor condition number of Jacobian matrix.

8.5 EM FEM Solver

GSS has a electromagnetic solver based on finite element method. This solver calculates the distribution of electromagnetic field radiated by monochrome (light) wave. The photon generated carrier density in semiconductor region can be got at the same time.

Syntax

PHOTOGEN	WAVELEN= <n> IN</n>	TENSITY= <n> [ANGLE=<n>] WTM=<n> WTE=<n></n></n></n></n>
	[phase.diff= <n< td=""><td>>] [quan.eff=<n>]</n></td></n<>	>] [quan.eff= <n>]</n>
METHOD	Type=EMFEM	[LS=LU]
SOLVE		
LSOURCE	Type=UNIFORM	Tdelay= <n> Power=<n></n></n>
LSOURCE	Type=PULSE	Tdelay= <n> Tr=<n> Tf=<n> Pw=<n> Pr=<n></n></n></n></n></n>
	Powerhi= <n></n>	Powerlo= <n></n>
LSOURCE	Type=LSHELL	DLL= <s> Func=<s></s></s>

Syntax for PHOTOGEN

parameter	\mathbf{type}	default	${f unit}$	description
WAVELEN	number	0.532	$\mu\mathrm{m}$	The wavelength of incident monochrome wave.
INTENSITY	number	1.0	${ m W\cdot cm^{-2}}$	The power density of incident wave.
ANGLE	number	90	$_{ m degree}$	The clockwise angle of the ray direction relative to
				the horizontal axis.
WTM	number	1.0	-	The percentage of intensity of TM model.
WTE	number	0.0	-	The percentage of intensity of TE model.
phase.diff	number	0.0	$_{ m degree}$	The differentiation of phase angle between TE model
				and TM model. $\Delta \Phi = \Phi_{TM} - \Phi_{TE}$
quan.eff	number	1.0	-	The quantum efficiency (which means electron-hole
				pares generated by one photon) of photon generation.

Syntax for LSOURCE

parameter	$_{ m type}$	default	\mathbf{unit}	description
Type	string	-	-	The type of light source.
Tdelay	number	0.0	S	The delay time before the activation of the light source.
Tr	number	1e-15	S	The rise time of the intensity of the pulse-type light source.
Tf	number	1e-15	S	The fall time of the intensity of the pulse type light source.
Pw	number	0	s	The pulse width of the intensity of the pulse type light source.
\Pr	number	0	S	The repetition period of the intensity of the pulse type light source.
Power	number	1.0	-	The multiply factor to photon generated carrier density.
Powerhi	number	1.0	-	The higher multiply factor to photon generated carrier density.
Powerlo	number	0	-	The lower multiply factor to photon generated carrier density.

```
DLL string - - The name of dynamic library file.

Func string - - The name of the function loaded from dynamic library file which calculates power coefficient.
```

Hint

User need to build a vacuum region surrounding device and a PML region surrounding vacuum region. These two region should have a thickness of no less than one wave length.

The work flow of EMFEM solver shows as follows. GSS set its internal solver to EMFEM when meets **METHOD** command with **EMFEM** type. The actual solving action takes place when meets the next **SOLVE** command. GSS will search the first **PHOTOGEN** command in the input list, using the parameters in this command during the solve procedure. This **PHOTOGEN** command will be removed from input list after solving action. As a result, user can set multi **PHOTOGEN** statements and repeat **SOLVE** command for corresponding times to calculate several beams of monochrome wave, during which the photon generated carrier density will be added to previous result.

The iterative method such as GMRES usually leads to divergence when solving FEM problem. LU factorization is highly recommend.

EMFEM only gets the photon generated carrier density. User should set **one LSOURCE** to describe the time evolution of the light source. The actual photon generated carrier density used in semiconductor simulation is the original value multiplied with **power** coefficient specified within **LSOURCE**.

When DDML1E or DDML2E solver is loaded for further simulation, the photon generated carrier will be considered.

User can define their own light source by dynamic loaded library as voltage or current source. Here is a template.

```
foo.c:
    double lsrc_power(double time) /* in the unit of s */
    {
        double power;
        /* calculate the power of light source */
        return power;
    }
```

8.6 IV File Format

GSS can generate IV record file for DC sweep, transient and AC sweep calculations. Here is the file format for the three situations.

The file for DC sweep: The first line is begin with '#', followed by the name of each electrode. The remain part is the potential and current for each electrode, each takes one column. The unit of potential is volt and the unit of current is mA.

The file for transient calculation is nearly the same as the file for DC sweep, besides that the first column is the time with the unit of ps.

The file for AC sweep has the same head as above. The remaining part is organized as follows: The first column is the frequency with the unit of MHz. Then the IV properties of each electrode. Each electrode takes six columns, the real, image and amplitude of potential, followed by three columns for current.

Note: the electrode potential may not equal to the application voltage if lumped elements take place.

9 FILE I/O 34

9 File I/O

9.1 Introduction

The IMPORT and EXPORT statements are used to read and write solutions from a CGNS or TIF file. A model CGNS file only contains semiconductor device structure while a core CGNS file has previous solution data besides device structure. The TIF(Technology Interchange Format) file is an ASCII file used by Synopsys Medici software which equivalence to core CGNS file. We offer a small code TIFTool which can open TIF file, view the mesh and solution data and convert it to CGNS file.

9.2 IMPORT and EXPORT

Syntax

IMPORT	CoreFile= <s></s>	ModelFile= <s></s>	
EXPORT	CoreFile= <s></s>	[AscFile= <s>]</s>	[VTKFile= <s>]</s>

parameter	$_{\mathrm{type}}$	default	\mathbf{unit}	description
CoreFile	string	-	-	Write/read device structure and solution data
				to a CGNS file.
ModelFile	string	-	-	Read device structure from a CGNS file which
				probably crated by SGframework or converted
				from Medici TIF file by TIFTool.
AscFile	string	-	-	Write device structure and solution data to a
				TIF file. At present, we can't make our TIF
				file be accepted by Medici.
VTKFile	string	-	-	Write mesh and solution data to VTK file.

Example

EXPURT	CoreFile=init.cgns	AscFile=init.tif
IMPORT	ModelFile=pn.cgns	
IMPORT	CoreFile=pn.cgns	

Hint

VTK file is intended to be used for post process. User can use Paraview¹, MayaVi or VisIt² to open and view VTK file. Further more, CGNS file is also supported by VisIt.

 $^{^{1}}$ http://www.paraview.org

²http://www.llnl.gov/visit

10 POST PROCESS 35

10 Post Process

10.1 Plot

The **PLOT** statement initializes the graphical display device for two and three dimensional plots of device characteristics(3D) and device meshes(2D).

Syntax

parameter	\mathbf{type}	$\operatorname{default}$	\mathbf{unit}	description
Variable	string	-	-	This parameter specifies plot context.
PS.OUT	string	-	-	Specifies the postscript file name. The plot window will be saved to it.
TIFF.OUT	string	-	-	Specifies the TIFF file name. The plot window will be saved to it. Only available for X11 system.
Resolution Measure	string string	RES.Middle Linear	-	The resolution of plot window. Specifies the data axis to be linear or logarithmic.
AzAngle	number	240	degree	The initial azimuthal rotation angle, $0 \le \mathbf{AzAngle} < 360$.
ElAngle	number	60	degree	The initial elevation rotation angle. $0 \le ElAngle < 70$.
Style	string	Color	-	The plot style.

Example

```
PLOT Variable=Mesh PS.OUT=mesh.ps
PLOT Variable=Nd Resolution=RES.High AzAngle=240 ElAngle=40 Style=Scale
PLOT Variable=Potential Resolution=RES.Middle TIFF.out=potential.tiff\
AzAngle=240 ElAngle=40 Style=Color
```

Hint

PlotMesh is an interactive GUI for mesh display only exist for X11 system.

The **PLOT** command can be used on both X11 and Win32 systems. The 3D plot can be rotated by mouse and terminated by ESC key press. If PS.OUT or TIFF.OUT argument is specified, the latest window image will be saved.

10 POST PROCESS 36

10.2 Probe

The **PROBE** statement is used to extract field data along a user defined segment. The segment can be a boundary or a segment pre-defined in the region. For the in-region segment, GSS will set it as Neumann boundary with no heat flux which takes not effect to simulation result.

Syntax

Probe Variable=(Na|Nd|ElecDensity|HoleDensity|Potential|EFieldX|EFieldy|Temperature)
Region=<s> Segment=<s> ProbeFile=<s> Append=

parameter	\mathbf{type}	default	\mathbf{unit}	description
Variable	string	-	-	This parameter specifies probe context.
Region	string	-	-	Specifies the region name which the segment belongs to.
Segment	string	-	-	Specifies the segment name for probing the data.
ProbeFile	string	-	-	The file name for recording the data.
Append	bool	OFF	-	Specifies the data should be appended to the file.

Example

```
PROBE Variable=Nd Region=Si Segment=PB1 ProbeFile=Nd.txt Append=Off
PROBE Variable=Potential Region=Si Segment=PB2 ProbeFile=P.txt Append=On
```

Hint

Each **PROBE** statement records one variable for the whole segment to a user-specified file. GSS pushes **PROBE** statement as the sequence of input text into a stack until a **SOLVE** statement is met. These **PROBE** statements record the data during solve process. After that, GSS will clear the stack. In short, **PROBE** only operates for next **SOLVE** process.

The file format for probe is show as follows:

The head of file is the segment information, including region name, segment name, total node number and the location of each node. The last line of head shows the solve type and variable name. The solve type can be "EQUILIBRIUM", "STEADYSTATE", "DC-SWEEP_VSCAN", "DCSWEEP_ISCAN" and "TRANSIENT". For the last three types, GSS will record V/I/Time in the first column, respectively. The variable value for all the nodes are listed in the same line.

11 Convergence Problem

The core arithmetic of GSS is solving the large scale nonlinear equations arisen from semiconductor drift-diffusion model by Newton's Iterative method. There are three factors which affect the convergence of nonlinear solvers: the initial value, the Jacobian Matrix and the inner linear solver. One must ensure that the initial value is sufficiently near the real solution, the Jacobian Matrix is exact or at least nearly exact and the inner linear solver can give a suitable solution. When one of the three demands is not satisfied, the convergence problem may raise. However, several skills can help convergence.

If the first time running failed due to bad initial value, one can employ a transient solver to do a time evolved solution. Set time step to a few ps, and the solution on every step may get convergence. After some certain steps, the initial shock is damped and physical variables are forced to get close to real quantities. Then the steady-state solver may work and you can get the equilibrium solution.

Since version 0.46, GSS use automatically differentiation to calculate Jacobian matrix. In most situations, author can guarantee that Jacobian matrix is exact, except some rigorous situations when round-off error is un-neglectable. However, GSS offers alternative choice, the Matrix-Free method to set Jacobian Matrix by finite difference approximation. This choice can be invoked with the command line option -snes_mf_operator. The Matrix-Free method works well when impact ionization takes place, but it runs much slower than original method.

Sometimes the LineSearch method may failed due to bad search direction. If one get divergence message during DC sweep and transient simulation when using LineSearch method, one can try TrustRegion or basic Newton method.

Newton damping is a powerful tool to help convergence. It can work with LineSearch and Basic Newton solvers. GSS has two damping method, BankRose and Potential. Usually, damping Potential is better than BankRose.

The most difficult problem is the failure of inner linear solver. When the Jacobian Matrix is singular, problems may happen. Especially one sets electrodes with lumped resistor or current sources. If one get a convergence failed message for these situations, please check the problem by adding command line option -ksp_monitor to exam the convergence history. For more information, one can use -ksp_singmonitor to get the condition number of matrix (this works only with GMRES method).

The author suggests some method to overcome the problem. First, one may improve the condition number by enlarging the DopingScale, but this will increase numerical error. Second, one should carefully choose the linear solver.

Here is the introduction of the main linear solvers GSS can use. GMRES is a robust method for non-symmetric matrices. It must retain all the previous vectors during iterative. The implemented code often uses a "restart" method to avoid large memory requirement. Sometimes the solution breaks when restart too often. One can increase this restart steps by $-ksp_gmres_restart < n >$ (n is the restart steps, default 150) BiCG and CGS often have irregular convergence behavior. The irregular result may get things worse. Bi-CGSTAB is the improved method to BiCG and CGS, which avoids the irregular convergence patterns of BiCG/CGS while maintaining about the same speed of convergence. TFQMR avoids the irregular convergence behavior of BiCG. Also it avoids some breakdown situations of BiCG. When BiCG temporarily stagnates or diverges, TFQMR may still works. At last, LU factor-

ization is the basic method for solving linear systems. Besides build-in LU solver, PETSC can be compiled with external LU factorization package such as SuperLU and UMFPACK. This method works slow but usually more stable than iteration solver.

In conclusion, LU factorization is recommend for conquering the singular problem. But user can try GMRES with large restart steps, Bi-CGSTAB and TFQMR methods for better efficiency.

12 Memory and CPU requirement

Thanks to C++'s dynamic memory manege system, GSS can solve problems with any scale (at least, theoretically). The memory requirement is not a serious bottleneck. A very large problem which contains 100K nodes only requires about 300MB memory. This requirement is easy to be satisfied with modern computers.

Because the core arithmetic of GSS is solving nonlinear equations, which involves lots of solutions of linear system, the CPU time is related with linear solvers, which is $O(n^3)$ with LU solver and $O(n^2)$ with krylov iterative solver, in which n is the problem scale.

Fig5 shows the CPU time vs node number with a PN diode simulation by BCGS method on a Xeon 3.6GHz workstation. The time is approximate the square of problem scale. It only requires serial seconds when total node number less than 5000. But CPU time raises to some minutes when node's number reach to 100K.

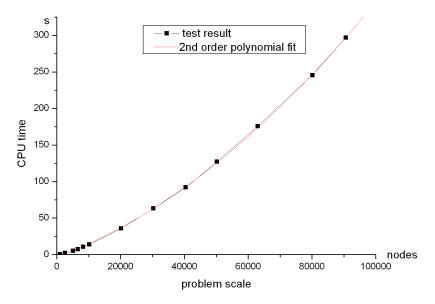


Figure 5: CPU time vs problem scale

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