TOWARDS A MORE ACCURATE EXTRACTION OF THE SPICE NETLIST FROM MAGIC BASED LAYOUTS

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**Introduction**

The extraction of the SPICE netlist form MAGIC based layouts is investigated. It is assumed that the layout is fully coherent with the corresponding mask representation.\(^1\)

The process of the extraction can be made in three steps:

1. **Step 1:** extraction of .EXT file from layout, through MAGIC command `extract`
2. **Step 2:** extraction of the netlist from .EXT file through `ext2spice` extractor
3. **Step 3:** correction of the netlist through `ext2spice.corr` program.

Each of these steps introduces some approximations, most of which can be optimized, and some errors, most of which can be corrected. Aim of this work is the description of each step, of the approximations and errors on each step, and of the corresponding optimizations and corrections to be made in order to improve the accuracy of the extraction.

The HP AMOS14TB 0.5µm process with linear capacitor and silicide block options and the corresponding SCN3MLC_SUBM.30.tech27 technology file will be used in the following examples.

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**Step 1 - From the Layout to the .EXT file**

The MAGIC command `extract` provides the extraction of the layout according to some information contained in the technology file. The technology file is composed of 17 sections (`tech, version, planes, types, contact, styles, compose, connect, cifoutput, cifinput, mzrouter, drc, extract, wiring, router, plowing, plot`) and the sections of interest for the extraction are the 8\(^{th}\) (`connect`) and the 13\(^{th}\) (`extract`).

The `connect` and the `extract` sections contained in the SCN3MLC_SUBM.30.tech27 Technology File are reported in Appendix A. See MAGIC Maintainer’s Manual #2 (Technology File Manual) for the formalism used in these sections.

**The information contained in the .EXT file**

The circuit is always extracted by using three elements: the `node` element, the `capacitance between nodes` element, and the `FET` element.

---

\(^1\) It is good practice to write into cif format (:cif MAGIC command) and to read from the written cif format (:cif read MAGIC command) the layout before the extraction.
a) Node Element

Each $\lambda \times \lambda$ tile of the layer is part of a node. The node is determined through the information contained in the `connect` section of the technology file. If each tile is connected to the adjacent one, both tiles are part of the same node. To each node element corresponds a line in the .EXT file which looks as follows:

node nodename R C X Y layer a1 p1 a2 p2 a3 p3 ........

where:

nodename is the label of that node. MAGIC assigns an arbitrary name if there is no label on that node.

$R$ is the resistance of the node and it is calculated according to the formula:

$$R = \sum_i R_i$$

$$R_i = \frac{RSQ_i}{2} \left[ \frac{P_i^2}{4A_i} - 2 + \sqrt{\left( \frac{P_i^2}{4A_i} - 2 \right) - 4} \right]$$

(1)

where $RSQ_i$ is the resistance per square of the $i^{th}$ layer, and its value in mOhm/square (default=0) is given in the `extract` section under the corresponding keyword `resist` (giving the precedence to the upper plane in case a layer is represented in more than one plane, as it is for the contacts), $A_i$ and $P_i$ are respectively the total area and the total perimeter of the $i^{th}$ layer. The value of $R$ is expressed in Ohm and it is approximated to the nearest integer. This formula gives an exact value only for rectangular sections composed of a single layer. The resistance is always calculated along the longest direction of the rectangle. In all other cases $R$ represents a bad approximation and for this reason $R$ is ignored during Step 2 of the extraction. As a consequence, the values reported in the `extract` section under the keyword `resist` are not determinant for the extraction.

$C$ is the capacitance between the node and the substrate and it is calculated according to the formula:

$$C = \sum_i C_i$$

$$C_i = ACAP_i \times A_i + PCAP_i \times P_i$$

(2)

where $ACAP_i$ and $PCAP_i$ are respectively the capacitance per $\lambda^2$ associated to the area and the capacitance per $\lambda$ associated to the perimeter (fringing) of the $i^{th}$ layer, and their values in aF (default=0) are given in the `extract` section respectively under the keywords `areacap` and `perimcap` (see the Manual for the formalism), $A_i$ and $P_i$ are respectively the total area and the total perimeter of the $i^{th}$ layer. The value of $C$ is expressed in aF and it is approximated to the nearest integer. In most cases $C$ represents a good approximation and for this reason it is taken into account dur-
ing Step 2 of the extraction. As a consequence, the values reported in the extract section under the keywords areacap and perimcap are determinant for the final extraction and their value must be accurately verified and updated through the General Process Specifications and the Beta Technology Files and the MOSIS Parametric Test Results.

\( X Y \) are the coordinates of the node in units of \( \lambda \). For single layer nodes, the low corner of the most left side is assumed as origin. For multiple layer nodes, the low left corner of the most down among the most left contacts is assumed as origin.

**layer** is the name of one of the layers of the node. For single layer nodes, it is the name of that layer. For multiple layer nodes, it is the name of the contact the corner of which is assumed as origin.

\( a_1 p_1 a_2 p_2 a_3 p_3 \ldots \) are respectively the area and the perimeter of the layers of the node which belong to the resist-classes 1, 2, 3 \ldots where resist-class 1 is the first line with keyword resist in the extract section, class 2 is the second line with keyword resist and so on. In the technology file used here for the examples, there are 6 resist-classes. The values \( a_1, p_1, a_2, p_2, \ldots \) are used for the evaluation of the resistance of the node but not for the evaluation of its capacitance. As a consequence these values are not determinant for the final extraction.

**b) Capacitance Element**

If two or more nodes are present in the layout, the capacitance between each couple of nodes (if \( > 0 \)) is extracted, and reported through the capacitance element. The corresponding line in the .EXT file looks as follows:

```
cap nodename1 nodename2 C
```

where:

**nodename1 nodename2** are the labels of the two nodes; MAGIC uses an the arbitrary names if there is no label.

\( C \) is the capacitance between the two nodes, in aF.

The capacitance between two nodes is evaluated only in the following cases:

- a layer of one node overlaps a layer of the other node and the two layers belong to different planes; the corresponding keyword in the extract section is overlap \( (aF/\lambda^2) \), and includes the possibility of a shield layer between the two layers (see Manual); if this capacitance is extracted for a given area, an equal area is subtracted in the evaluation of the node capacitance to substrate for the layer belonging to the upper plane, which is correct;

- a layer of one node is adjacent to a layer of the other node and the two layers belong to different planes; the corresponding keyword in the extract section is sideoverlap \( (aF/\lambda) \) and in-
cludes the possibility of a shield between the two layers; if this capacitance is extracted for a
given perimeter, an equal perimeter is subtracted in the evaluation of the node capacitance to
substrate for the layer belonging to the upper plane, which is correct;

• a layer of one node is close, but not adjacent, to a layer of the other node and the two layers
belong to the same plane; the corresponding keyword in the extract section is sidewall (aF,
and must be calculated by using the factor $2x\lambda_d/\lambda_w$ where $\lambda_d$ is the distance and $\lambda_w$ the
width); the capacitance is extracted only if the distance between the adjacent layers is below
the value reported under the sidehalo keyword at the beginning of the extract section.

c) FET Element

If two nodes are connected through a specific layer, and it exists a corresponding fet line with the
specified layer indicated in the 2nd element under the corresponding keyword fet in the extract
section (see Manual), a FET is extracted, the gate of which is the specific layer. The corresponding
line in the .EXT file looks as follows:

fet model X Y X+1 Y+1 a p snode gatenodename gatewidth 0 nodename1 width1 jlabel1 noden-
name2 width2 jlabel2

where:

model is the name of the model of the FET, as specified in the 5th element under the correspond-
ing keyword fet in the extract section.

X Y X+1 Y+1 are the Cartesian coordinates and the Cartesian coordinates plus 1 of the low left
corner gate layer in units of $\lambda$.

a p are respectively the area and the perimeter of the gate

snode is the label of the node at which the substrate must be considered connected, as specified
in the 6th element under the corresponding keyword fet in the extract section. The [7th] facultative
element under the corresponding keyword fet in the extract section considers the connection of
the body to a well, when the well is superposed to the channel of the FET (see Manual).

gatenode is the label of the gate node; gatewidth is the width of the gate contact (i.e. the length
of the channel of the FET) in units of $\lambda$.

nodename1, nodename2 are respectively the labels of the drain and of the source, the layer
(and number) of which are specified in the 3rd (and 4th) element under the corresponding keyword
fet in the extract section (see the Manual), width1, width2 are respectively the widths of the drain
and of the source (i.e. the width of the channel of the FET) in units of $\lambda$ and jlabel1, jlabel2 are
respectively the labels at the drain and at the source junction (in the layout these labels must be
followed by the "$" symbol). According to the Magic extraction criteria, during Step 1 the drain
junction of each finger is the one at the left or below the gate. This criteria for the choice of the
drain is not determinant for the final extraction because, as it will be discussed, during Step 2 the side with the \textit{jlabel S$ is assumed as source and the one with the \textit{jlabel D$ is assumed as drain, disregarding the orientation of the gate.

\textbf{The errors contained in the .EXT file}

Three \textbf{errors} are made during Step 1 in the evaluation of the capacitance for the node and capacitance element.

\textbf{Type 1 Error.} An over-estimation is made by the extractor when two or more layers of the same node and belonging to different planes are overlapped. The error is due to the fact that the capacitance of the upper layers to the substrate is in most cases added, but its value should be zero because of the shield of the lower layer.

This is the case for example of all the contact layers (pc, ndc, pdc, nsc, psc, cwc), which are characterized by the fact that lie both in the \textit{active} plane and in the \textit{ml} plane (see Manual). In the evaluation of the capacitance of the area to the substrate, both the planes are considered, but only the \textit{active} plane should be considered because the \textit{ml} plane is shielded by the \textit{active} plane. In the evaluation of the capacitance of the perimeter to the substrate, both the planes are considered, but only the \textit{active} plane should be considered because the \textit{ml} plane is shielded, except when the contact layers are along the perimeter of the node.

Both problems could be corrected by suitably modifying few lines of the \textit{extract} section, as shown. The line $^{2}$

\begin{verbatim}
  areacap (m1,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1
  2.700
\end{verbatim}

must be replaced with the line

\begin{verbatim}
  areacap (m1,m2c/m1)/metal1  2.700
\end{verbatim}

which means that no capacitance to substrate is calculated for the part of the contact lying in the metal1 plane.

The line

\begin{verbatim}
  perimc (ndc/a,nsc/a,pdc/a,psc/a,pc/a,cwc/a)/active (ndiff,nsd,pdiff,psd,poly,cwnsd)/active
  -9.300
\end{verbatim}

(note the minus sign) must be added, which means that, if the contact is not along the perimeter of the node, the capacitance of that perimeter to the substrate is adjusted to be zero.

\footnote{Don’t be surprised if sometimes the name of a layer is reported more than once in the same group of a line in the Technology File (for example cwc/ml appears twice in this case). The origin of this redundancy is at the Technology File generation level (ask Anand Kandasamy for details), but it is without consequence for the extraction.}
Example 1

From .EXT file:

node "a_82_n45#" 7 136410 82 -45 p 0 0 0 0 15000 600 2500 200 0 0 0 0

resistance from Eq.(1):

\[
\left( \frac{P_1^2}{4A1} - 2 \right) + \sqrt{\left( \frac{P_1^2}{4A1} - 2 \right)^2 - 4} - 1.8 + \left( \frac{P_2^2}{4A2} - 2 \right) + \sqrt{\left( \frac{P_2^2}{4A2} - 2 \right)^2 - 4} - 0.07 = 6.788
\]

capacitance from Eq.(2): \( A1 \cdot 7.92 + P1 \cdot 15 + A2 \cdot 2.7 + P2 \cdot 9.3 = 136410 \)

which is over-estimated. After the modification of the technology file, it results from .EXT file:

node "a_82_n45#" 7 128265 82 -45 p 0 0 0 0 15000 600 2500 200 0 0 0 0

capacitance from Eq.(2): \( A1 \cdot 7.92 + P1 \cdot 15 + (P2 - 50 - 50 - 50) \cdot 9.3 = 128265 \)

and it is a better approximation.

It is worth noting that this is only a partial solution of the problem, and it is subject to errors. For example when a \( ml \) line connected to the contact flows above the active layer of that contact, its capacitance to the substrate is erroneously added. An alternative solution not subject to errors will be proposed in the next section.

**Type 2 Error.** In the case of contacts, metals and poly which flow over a well and are connected to that well (i.e. are parts of the same node), the area and perimeter capacitances to substrate of the metal and poly as well as the part of the perimeter capacitance to substrate of the contact adjacent to the well are added (no overlap or sideoverlap to the well and consequent subtraction occurs, because the well is part of the same node). This is an error because all these layers are shielded by the well.
To give a simple example, the linear capacitors can be considered, where the perimcap capacitance to substrate of \( cwc \) (the well contact) along the perimeter which is not adjacent to the \( cwnsd \) (the n-diffusion) as well as the area and perimeter capacitance to substrate of a metal connected to the \( cwc \) are erroneously added.

In the case of linear capacitors, a partial solution of the problem can be obtained by introducing the following additional changes in the extract section.

The line:

\[
\text{perimc (m1,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metall-}
\begin{align*}
\text{~(m1,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metall & 9.300}
\end{align*}
\]

must be replaced with the line (the layer \( cwc/m1 \) is deleted):

\[
\text{perimc (m1,ndc/m1,nsc/m1,pdc/m1,psc/m1,pc/m1,m2c/m1)/metall}
\begin{align*}
\text{~(m1,ndc/m1,nsc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metall & 9.300.}
\end{align*}
\]

The line added to correct the Error 1

\[
\text{perimc (ndc/a,nsc/a,pdc/a,psc/a,pc/a,cwc/a)/active (ndiff,nsd,pdiff,psd,poly,cwnsd)/active}
\begin{align*}
-9.300
\end{align*}
\]

must be replaced with the line:

\[
\text{perimc (ndc/a,nsc/a,pdc/a,psc/a,pc/a)/active (ndiff,nsd,pdiff,psd,poly)/active}
\begin{align*}
-9.300
\end{align*}
\]

**Example 2**

From .EXT file:

```
node "w_246_n38#" 0 2606.88 246 -38 cw 0 0 0 0 0 0 24 20 0 0 0 0
capacitance from Eq.(2): A1·8.46+ P1·0+ A2·2.7+ P2·9.3 = 2606.88
```
which is over-estimated. After the further modification of the technology file, it results from .EXT file:

```
node "w_246_n38#" 0 2532.48 246 -38 cw 0 0 0 0 0 24 20 0 0 0 0
```
capacitance from Eq.(2): \( A1 \cdot 8.46 + p1 \cdot 0 + A2 \cdot 2.7 + (P2 - 4 - 4) \cdot 9.3 = 2532.48 \)

and it is a better approximation.

It is worth noting that an over-estimating error due to the capacitance of the metal to substrate is still present. The solution proposed in the next section will solve also this problem.

**Type 3 Error.** In the case of linear capacitors, the sideoverlap capacitance between \( wcap \) (the active area of the capacitor) and \( cwnsd \) (the diffusion) and the sideoverlap capacitance between \( poly \) and \( capwell \) (the well of the capacitor) are considered under the same sideoverlap keyword. This is an error because they are characterized by different thickness of the oxide and then subject to two different values of \( aF/\lambda \). To solve this problem the following changes must be made.

The line:

```
sideoverlap (poly,pres,wcap,pc/a,wcap,pc/a)/active ~ (poly,pres,wcap,pc/a,wcap,pc/a)/active ~ space/w 15
```

must be replaced with the line:

```
sideoverlap (poly,pres,wcap,pc/a,wcap,pc/a)/active ~ (poly,pres,wcap,pc/a,wcap,pc/a,cwnsd)/active ~ space/w 15
```

and the line

```
sideoverlap (wcap)/active (cwnsd)/active ~ space/w 40
```

must be added, where the value 40 \( aF/\lambda \) can be deduced from a simulation, from a suitably equation or from the MOSIS Parametric Test Results (see Appendix B).
Example 3

From .EXT file:

node "A" 3 5336.28 141 -18 pc 0 0 0 0 11772 434 540 226 0 0 0 0
node "B" 1 115597 144 -20 cw 0 0 0 0 0 500 210 0 0 0 0
-cap "A" "B" 2.09531e+06

Capacitances from Eq.(2):
A1 = 7.92  + P1  + P1 9.3 = 5336.28i
A2 = 8.46 = 115597.44i

which are correct.

Internodal capacitance: \[ A3\cdot207.9^+ (A4-A3)^+ 7.92^+ P4 15^+ P5 9.3^+ 10040 = 2095305.36i \]

which is under-estimated due to Error 3.

After the further modification of the technology file, it results from the .EXT file:

node "A" 3 5336.28 141 -18 pc 0 0 0 0 11772 434 540 226 0 0 0 0
node "B" 1 115597 144 -20 cw 0 0 0 0 0 500 210 0 0 0 0
-cap "A" "B" 2.09781e+06

Internodal capacitance:
\[ A3\cdot207.9^+ (A4-A3)^+ 7.92^+ (P4-100)^+ P5 9.3^+ 10040 = 2097805.36i \]
which is a better (indeed with negligible difference in this case) approximation.

It is worth noting that, in this case, the solution adopted is effective in the correction of the error in the evaluation of the internodal capacitance. On the other hand, the absence of errors in the evaluation of the capacitance to the substrate is due to the modifications introduced in the extract section of the technology file to correct the errors of type 1 and 2.

A solution which globally corrects all the type 1 and type 2 errors

It has been shown that the three types of errors introduced during Step 1 can be partially corrected by modifying the extract section of the technology file. Apart from the correction of the error of type 3, which must be introduced in any case and it represents a global correction, the other modifications correct only partially the errors of type 1 and 2.

A different approach to globally correct all the errors of type 1 and 2 is now discussed. This approach is based on the introduction of an additional plane (we will call it “substrate”) and of an additional layer (we will call it “substrate” or “ss”) which lies in the additional substrate plane. By using this additional layer the capacitances to substrate are now evaluated as overlap and sideoverlap capacitances, instead of areacap and perimc capacitances. The main advantage is that the shielding layers (available only with the overlap and sideoverlap keywords) can now be used for the capacitances to substrate. Another advantage is that the new layer, which must be present in all the layout under extraction, can be divided into areas thus allowing the simulation of the resistance of the substrate from one point to another of the layout. This additional layer must be also invisible at the CIF extraction level. The modifications which must be made in the technology file to introduce and use the additional plane and layer are discussed in the Appendix C. A label SS can be given to the additional node the layer of which is ss.

Examples

By applying the modified technology to Example 1 it follows from the .EXT file:

without the ss layer (note the zero value capacitance)

node "a_82_n172#" 7 0 82 -172 p 0 0 0 0 15000 600 2500 200 0 0 0 0

With the ss layer (note that the capacitance to substrate is now represented by the capacitance element, while the node elements are ignored during the step 2 of the extraction because their capacitance is zero)

node "a_82_n172#" 7 0 82 -172 p 0 0 0 0 15000 600 2500 200 0 0 0 0
node "SS" 0 0 74 -184 ss 0 0 0 0 0 0 0 0 0
cap "a_82_n172#" "SS" 128265
By applying the modified technology to **Example 2** it follows from the .EXT file:

Without the ss layer (note the zero value capacitance)

node "w_246_n38#" 0 246 -38 cw 0 0 0 0 0 24 20 0 0 0 0

With the ss layer (again note that the capacitance to substrate is now represented by the capacitance element)

node "w_246_n38#" 0 246 -38 cw 0 0 0 0 0 24 20 0 0 0 0
node "SS" 0 245 -39 ss 0 0 0 0 0 0 0 0 0 0 0 0
cap "w_246_n38#" "SS" 2436.48

and now the result is now correct: \( A_{1.846} = 2436.48 \).

By applying the modified technology to **Example 3** it follows from the .EXT file:

Without the ss layer (note the zero value capacitance)

node "A" 3 0 141 -18 pc 0 0 0 0 11772 434 540 226 0 0 0 0
node "B" 1 0 144 -20 cw 0 0 0 0 0 500 210 0 0 0 0
cap "A" "B" 2.09781e+06

With the ss layer (again note that the capacitances to substrate is now represented by the capacitance element)

node "A" 3 0 141 -18 pc 0 0 0 0 11772 434 540 226 0 0 0 0
node "B" 1 0 144 -20 cw 0 0 0 0 0 500 210 0 0 0 0
node "SS" 0 0 134 -25 ss 0 0 0 0 0 0 0 0 0 0 0 0
cap "A" "SS" 5336.28
cap "A" "B" 2.09781e+06
cap "B" "SS" 115597
**Step 2 - From the .EXT file to the .CIR file (Ext2spice)**

The Ext2spice program provides the extraction of the Spice netlist from the Magic.EXT file. The Ext2spice program here considered was released by Lloyd Clonts (Oak Ridge National Laboratory) in June 23rd 1998.

The Ext2spice program runs with the command line:

```bash
ext2spice [options] nomefile.ext
```

and the output (-g option) are two files:

- `nomefile.cir`: Spice netlist containing options, sources, library lines and the line `.include nomefile.spice`
- `nomefile.spice`: Spice netlist containing the elements extracted from the .EXT file

The netlist containing the elements extracted from the .EXT file will be discussed, element by element, in the next sections. Information about the options can be found in the Ext2spice manual (Appendix D). Ext2spice uses a support file called Ext.defs (Appendix E) the role of which will be discussed in the next sections. In the Ext.defs file are also contained the Spice command lines which will appear in the .cir file.

**Capacitor Extraction**

The capacitors from the node element and from the capacitance element of the .EXT file are reported in the .spice file.

In particular, to each node element in the .EXT file

```plaintext
node nodename R C X Y layer a1 p1 a2 p2 a3 p3 ........
```

it corresponds in the .spice file the element

```plaintext
Cx newnodename $G_Vsub! C
```

---

3 In the original release (1QT 1998) some errors were found, which have been corrected by Lloyd under our stimulus, generating the new release here discussed.
where Cx is an arbitrary name and newnodename is equal to nodename if the node has a label or an arbitrary node number (indeed much shorter than the arbitrary node names used by Magic in the .EXT file) if there is no label on that node. The line "Vsubstr $G\_Vsub! 0 DC 0V" is added by Ext2spice (-D option) to insure the connection of the global substrate node $G\_Vsub!.

To each capacitance element in the .EXT

\[
\text{cap nodename1 nodename2 C}
\]

it corresponds, in the .spice file, the element

\[
\text{Cx newnodename1 newnodename2 C}
\]

where Cx is the name of the element according to the general device labeling (see Ext2spice Manual) or an arbitrary name if there is no label on that element; newnodename1 (newnodename2) is equal to nodename1 (nodename2) if the node has a label or an arbitrary node numbers (much shorter than the arbitrary node name used in the .EXT file) if there is no label on that node.

It is worth pointing out that a capacitance is reported only if it is not floating and if its value is higher that the value indicated with the option -c cmin (default minimum: 1fF). All the values of these capacitances (linear, parasitics, ...) are reported as well. The accuracy in their value is related to the accuracy of their extraction during Step 1.

FET Extraction

All the FETs from the fet element of the .EXT file are reported in the .spice file.

In particular, to each fet element in the .EXT file (with some exceptions discussed in the next sections, such as in the case of the extraction of resistors)

\[
\text{fet model X Y X+1 Y+1 a p snode gatenodename gatewidth 0 nodename1 width1 jlabel1 nodename2 width2 jlabel2}
\]

it corresponds in the .spice file the element

\[
\text{Mx newdrainnodename newgatenodename newsourcenodename substratenodename newmodel [M=m] W=w L=l AD=ad PD=pd AS=as PS=ps [NRG=nrg NRD=nrd NRS=nrs]}
\]

If the -mm option is used, multifinger FETs are merged in a single FET by adding the M=m term, where m is the number of fingers.
Mx is the name of the element according to the general device labeling (see Ext2spice Manual) or an arbitrary name if there is no label on that element;

\textit{newgatenodename} is equal to \textit{gatenodename} if the node has a label or an arbitrary node numbers (much shorter than the arbitrary node name used by Magic in the .EXT file) if there is no label on that node.

\textit{newdrainnodename newsourcenodename} respectively the drain (to which the area \textit{AD} and the perimeter \textit{PD} are associated) and the source name (to which the area \textit{AD} and the perimeter \textit{PD} are associated), are equal to \textit{nodename1} and \textit{nodename2} with a order which, if \textit{jlabel1} and \textit{jlabel2} are zero, depends on the orientation of the FET. The drain node is chosen according to Magic extraction criteria where the drain junction of each finger is the one at the left or below the gate. On the contrary, if one \textit{jlabel1} and/or \textit{jlabel2} is labeled as D (or S) (remember that in the layout these labels must be followed by the "$" symbol), the corresponding junction will be assumed as drain (or source). Attention must be paid to the fact that, for multifinger FETs, the junction label \textit{jlabel} must lie in the same finger where is placed the label of the device, otherwise it will be ignored!

\textit{substratenodename} is equal to \textit{snode}

In order to recognize the substrate node used in Ext2spice ($G_{-}\text{Vsub}$!) and to use the additional ss layer, the following changes must be made for n-channel FETs in the \textit{fet} lines of the \textit{extract} section:

```
# REPLACED fet nfet ndiff,ndc 2 nfet Gnd! pwell 15 324
fet nfet ndiff,ndc 2 nfet $$G_{-}\text{Vsub}$! pwell,ss 15 324
# REPLACED fet nfet ndiff,ndc 1 nfet Gnd! pwell 15 324
fet nfet ndiff,ndc 1 nfet $$G_{-}\text{Vsub}$! pwell,ss 15 324
```

A problem arises when the body must be connected to pwell. When the MAGIC extractor must choose between ss and pwell, it chooses ss (i.e. the layer with the lower planeorder), which is an error. A way to solve this problem is to delete the layer ss from the areas where \textit{nfet} and \textit{pwell} are simultaneously present. On the other hand, for most of the technologies that we use, the connection of the body of an n-channel FET to a pwell different from the p substrate is not allowed (i.e. the body of n-channel FETs is always connected to the substrate).

\textit{newmodel} is equal to the model defined in the Ext.defs file with the lines:

```
def model newmodel M n
def model newmodel M p
```

where \textit{model} is equal to \textit{name} of the \textit{fet} element in the .EXT file (see Technology File Manual) and it is different for n-channel and p-channel devices

\textit{w} is the gate width, and it is equal to \textit{width1} (=\textit{width2})
$l$ is the gate length, and it is equal to $\text{gatewidth}$

$ad$, $pd$, $as$ and $ps$ are the areas and perimeters of drain and source respectively, and the layer to be used in this evaluation is the one indicated under the $\text{resistclass}$ keyword in the extract section of the technology file. In particular:

the drain/source layer for n-channel devices is defined by using the Ext.defs line $\text{nrclass n}$, where $n$ is the $n^{th}$ $\text{resistclass}$ line in the extract section;

the drain/source layer for p-channel devices is defined by using the Ext.defs line $\text{prclass n}$, where $n$ is the $n^{th}$ $\text{resistclass}$ line in the extract section.

Note that, in order to correctly extract the areas and perimeters in our case, the original lines in the Ext.defs file:

```
prclass 1
nrclass 2
nwclass 3
pwclass 4
plrclass 5
```

must be replaced with the lines (we are not interested in the $\text{nwclass}$, $\text{pwclass}$ and $\text{plrclass}$):

```
prclass 2
nrclass 1
nwclass 9
pwclass 9
plrclass 9
```

Note that, because the area and perimeter capacitances of the diffusions ($\text{ndiff}$, $\text{pdiff}$) and of the corresponding contacts ($\text{ndc}$, $\text{pdc}$) are included in the FET model, the corresponding parasitics in the node and capacitance elements of the .EXT file must be set to zero. In order to achieve this result, the layers $\text{ndiff}$, $\text{pdiff}$, $\text{ndc/a}$, $\text{pdc/a}$, $\text{ndc/ml}$ and $\text{pdc/ml}$ must be deleted from all the $\text{areacap}$, $\text{perimcap}$, $\text{overlap}$, $\text{sideoverlap}$ (not $\text{sidewall}$) lines in the $\text{extract}$ section which contain these layers.

Note also that, according to the $\text{extract}$ section lines:

```
resist (ndiff,anres,ndc/a,nsd,nsc/a)/active 1900
resist (pdiff,apres,pdc/a,psd,psc/a)/active 2000
```

the layers $\text{anres}$, $\text{nsd}$, $\text{nsc/a}$, $\text{apres}$, $\text{psd}$, $\text{psc/a}$ take part in the calculus of the areas and perimeters. In order to avoid this error (consider for example the case of a FET diffusion connected to a n-well with a nsc/a all around it: the area and perimeter calculated will be strongly over-estimated!), the two lines must be replaced with the lines:
Ext2spice calculates correctly the areas also for multifinger FETs and for FETs which share a diffusion. For equal width FETs sharing a diffusion, it calculates the total area and divides it by the total number of fingers. For different FETs having different width and sharing a diffusion, the calculus is made in order to assign to each finger the same area per unit of width. For example, in the case of two FETs with widths $W_1$ and $W_2$ and number of fingers $N_{F1}$ and $N_{F2}$, it results for the area per finger $A_{F2}$ of the second FET:

$$A_{F2} = \frac{A_{tot} \frac{W_2}{W_1 + W_2}}{N_{F1} \frac{W_1}{W_1 + W_2} + N_{F2} \frac{W_2}{W_1 + W_2}}$$

(3)

where $A_{tot}$ is the total diffusion area. With an analog procedure, the perimeter is correctly calculated for multifinger FETs and for FETs which share a diffusion:

$$P_{F2} = \frac{P_{tot} \frac{W_2}{W_1 + W_2}}{N_{F1} \frac{W_1}{W_1 + W_2} + N_{F2} \frac{W_2}{W_1 + W_2}}$$

(4)

If the -mn option is used, the number of squares of the gate $N_{RG}$, drain $N_{RD}$ and source $N_{RS}$ of the FET are calculated for the evaluation of the corresponding parasitic resistances.

The parameter $N_{RG}$ is calculated only for the FET active area ($n_{fet}$, $p_{fet}$) and in the drain to source direction, which can be considered a bad approximation. Moreover, in the case of multifinger FETs, the $N_{RG}$ of a finger is equal to the number of squares of the active area of that finger divided by $M$, which is an error. Finally, it must be considered that SPICE uses the same resistance per square of the diffusion to calculate $R_G$, $R_D$ and $R_S$, which is another error.

The parameters $N_{RD}$ and $N_{RS}$ are calculated in the drain to source direction by using the drain and source areas and perimeters respectively. This approach introduces an approximation which is strongly dependent on the geometry of the diffusion areas. It introduces also an over-estimation because also the areas of the diffusion contacts are included in the calculus. Moreover, in the case of multifinger FETs, the $N_{RD}$ ($N_{RS}$) of a finger is equal to the sum of the number of squares of the drain (source) divided by $M^2$, which is an error.

It is worth noting that, during the simulation, SPICE considers the $R_S$, $R_G$, $R_D$ as values per finger, and it divides these values by $M$ for the simulation of the overall FET, which is the correct approach.
The Step 3 will provide a more accurate extraction of the parasitic resistances $R_G$, $R_D$ and $R_S$, by correcting the errors and by including the resistance of the contact.

**Example 4**

![Diagram](image)

From the .EXT file (we use the technology file modified as in Step 1, in its version with the ss layer) it results:

```plaintext
node "A" 3 0 59 31 ndc 70 34 0 0 0 0 40 28 0 0 0 0 0
node "C" 9 0 66 26 pc 0 0 0 0 80 48 32 32 0 0 0 0 0
node "B" 12 0 83 28 nw 130 66 0 0 0 0 136 70 0 0 0 0 0
node "SS" 0 0 55 23 ss 0 0 0 0 0 0 0 0 0 0 0 0
cap "C" "SS" 1080
cap "A" "SS" 167.4
cap "B" "SS" 1930.92
fet nfet 66 31 67 32 40 28 "Gnd!" "C" 8 "Mx" "A" 10 0 "B" 10 "D"
```

The capacitances of A and B nodes to substrate are over-estimated (for example the capacitance of the A node must be zero, because included in the FET model through the source area and perimeter) and the body of the FET is connected to Gnd! while should be connected to the substrate SS. After the further modifications of the technology file, it results from the .EXT file:

```plaintext
node "A" 3 0 59 31 ndc 70 34 0 0 0 0 40 28 0 0 0 0 0
node "C" 9 0 66 26 pc 0 0 0 0 80 48 32 32 0 0 0 0 0
node "B" 3 0 83 28 nw 70 34 0 0 0 0 136 70 0 0 0 0 0
node "SS" 0 0 55 23 ss 0 0 0 0 0 0 0 0 0 0 0 0
cap "C" "SS" 1080
cap "B" "SS" 1800.72
fet nfet 66 31 67 32 40 28 "SS" "C" 8 "Mx" "A" 10 0 "B" 10 "D"
```
which is correct both in terms of capacitances and of connections.

Now, from the .spice file (-mn option) it results:

```
Mx B C A SS CMOSN W=3.00U L=1.20U AD=6.30P PD=10.20U AS=6.30P PS=10.20U
+ NRG=0.40 NRD=0.70 NRS=0.70
C1 B SS 1.8F
C2 C SS 1.1F
```

and the capacitances, areas and perimeters are correctly extracted, while the NRG, NRD and NRS parameters are affected by the previously discussed errors.

**Example 5**

In this example two n-channel FETs (M1, gate 10\(\lambda\times2\lambda\), 3fingers and M2, gate 20\(\lambda\times2\lambda\), 2fingers) share the source.

From the .EXT file (we use the technology file modified as in Step 1 and in Step 2, in its version with the ss layer) it results:

```
node "A2" 7 0 107 31 ndif 120 52 0 0 0 0 80 48 16 16 0 0
node "A" 13 0 61 31 ndc 110 62 0 0 0 0 80 56 88 52 0 0
node "B" 26 0 68 31 ndif 470 164 0 0 0 0 240 152 168 100 0 0
node "C" 61 0 61 24 pc 0 0 0 0 320 212 224 128 0 0 0 0
node "C2" 60 0 100 24 pc 0 0 0 0 264 192 160 96 0 0 0 0
node "SS" 0 0 59 21 ss 0 0 0 0 0 0 0 0 0 0 0 0
  cap "A" "SS" 243.96
  cap "B" "C2" 248.48
  cap "C" "SS" 5598
  cap "A" "B" 627.33
  cap "A2" "SS" 57.6
  cap "B" "C" 248.48
```
Now, from the .spice file (\textit{-mm} and \textit{-mn} options) it results:

\begin{verbatim}
M1 A C B SS CMOSN M=3 W=3.00U L=0.60U AD=3.30P PD=6.20U AS=6.04P PS=7.03U + NRG=0.07 NRD=0.12 NRS=0.22
M2 A2 C2 B SS CMOSN M=2 W=6.00U L=0.60U AD=5.40P PD=7.80U AS=12.09P PS=14.06U + NRG=0.05 NRD=0.07 NRS=0.17
C1 A C 0.2F
C2 A2 C2 0.0F
C3 A2 B 0.4F
C4 C2 SS 4.1F
C5 B SS 0.4F
C6 B C 0.2F
C7 A2 SS 0.1F
C8 A B 0.6F
C9 C SS 5.6F
C10 B C2 0.2F
C11 A SS 0.2F
\end{verbatim}

and the capacitances, areas and perimeters are correctly extracted, while the NRG, NRD and NRS parameters are again affected by errors. Note that in the perimeter calculation, the parts adjacent to the channel are included, which is a reasonable approach.

From Eq.(3) it results for the shared area:

\[
\begin{align*}
(10 \cdot 6 + 10 \cdot 11 + 20 \cdot 10 + 20 \cdot 5) \cdot \frac{10}{20 + 10} &= 6.043, \\
3 \cdot \frac{10}{10 + 20} + 2 \cdot \frac{20}{10 + 20} &= 6.043, \\
(10 \cdot 6 + 10 \cdot 11 + 20 \cdot 10 + 20 \cdot 5) \cdot \frac{20}{20 + 10} &= 12.086, \\
3 \cdot \frac{10}{10 + 20} + 2 \cdot \frac{20}{10 + 20} &= 12.086.
\end{align*}
\]
From Eq.(4) it results for the shared perimeter:

\[
(10 + 5 + 10 + 5 + 10 + 11 + 11 + 10 + 10 + 20 + 20 + 20 + 6 + 6) \cdot \frac{10}{20 + 10} \\
\frac{3 \cdot \frac{10}{10 + 20} + 2 \cdot \frac{20}{10 + 20}}{3 + 14.057}
\]

Resistor Extraction

The Magic extractor doesn't have any additional element for the extraction of the resistors. As a consequence, the three basic elements (the *node* element, the *capacitance* element and the *fet* element) must be used for this purpose. A very simple technique for the extraction of the resistors is to extract these devices as particular FETs. This technique will be discussed here and applied to the case of a poly resistors with silicide block (sb).

A poly resistor is composed of a polysilicon layer (*poly*) (the great part of which is subject to the silicide block *sb* (*pres*)), and by two polycontacts (*pc*). Till now, when a poly resistor is extracted, the two contacts and the *poly* appears as parts of the same node. The reason is the line:

```
poly,pres,wcap,nfet,wcap,pfet,pc/a poly,pres,wcap,nfet,wcap,pfet,pc/a
```

which is present in the *connect* section of the technology file and which "tells" the extractor that, when *poly*, *pres* and *pc* are in contact, they it must be considered connected each other (i.e. parts of the same node).

In order to extract the *pres* layer as a FET, this line must be replaced with the line:

```
poly, wcap,nfet,wcap,pfet,pc/a poly, wcap,nfet,wcap,pfet,pc/a
```

and the line:

```
fet pres poly 2 polyres $$G_{\text{Vsub}}$$ ss,nwell,pwell 0 0
```

must be added in the *extract* section.

Ext2spice recognizes this "fet" as a resistance if the Ext.defs file contains the lines:
where RSQ is the resistance per square of pres (90Ω in this technology) and CPAR is the parasitic capacitance (in fF/μ^2) used in the distributed model (-d option, see Ext2spice Manual). The number of squares SQ of the extracted resistance is then calculated by Ext2spice, from the corresponding fet line in the .EXT file, by using the "channel" (pres) area and the "channel" width, according to the equation:

\[
SQ = \frac{A}{W^2}
\]  

where \( W \) is the minimum between the widths of the "drain" and "source" connections (the poly adjacent to pres in this case). It turns out that the width of these connections is determinant for the correct extraction of the resistance.

In most cases the extracted SQ represents a good approximation of the value of the resistor. On the other hand, for a more accurate approximation:

- each corner along pres must be weighted with a factor 0.55 \(^4\);
- the real width of pres should be used, which is in general smaller than the drawn width;
- in the distributed case, the perimeter component of the parasitic capacitance is not included;
- the resistance of the poly near the pc should be added;
- the resistance of the pc should be added.

The Step 3 will provide a more accurate extraction, by including all these additional terms

Example 6

---

From the .EXT file (we use the technology file modified as in Step 1, in its version with the ss layer) it results:

node "a_n21_31#" 0 -21 31 sb 0 0 0 0 0 0 0 0 0 0 0 0 0
node "A" 26 0 -27 33 pc 0 0 0 0 360 154 50 40 0 0 0 0 0
equiv "A" "B"
node "a_n21_38#" 0 -21 38 sb 0 0 0 0 0 0 0 0 0 0 0 0 0
node "SS" 0 -35 13 ss 0 0 0 0 0 0 0 0 0 0 0 0 0
cap "A" "SS" 5440.2

and the two nodes A and B are "short circuited" (equiv "A" "B").

After the further modifications of the technology file, it results from the .EXT file:

node "B" 20 24 17 pc 0 0 0 0 30 17 25 20 0 0 0 0 0
node "a_n21_31#" 0 -21 31 sb 0 0 0 0 0 0 0 0 0 0 0 0 0
node "a_n21_33#" 18 0 -21 33 pres 0 0 0 300 120 0 0 0 0 0 0
node "A" 20 -27 33 pc 0 0 0 0 30 17 25 20 0 0 0 0 0
node "a_n21_38#" 0 -21 38 sb 0 0 0 0 0 0 0 0 0 0 0 0 0
node "SS" 0 -35 13 ss 0 0 0 0 0 0 0 0 0 0 0 0 0
cap "A" "SS" 632.1
cap "B" "SS" 632.1
cap "a_n21_33#" "SS" 4176
fet polyres -21 33 -20 34 300 130 "SS" "a_n21_33#" 0 "R1" "A" 5 0 "B" 5 0

and the resistor is extracted as a FET while the parasitic capacitances of the two contacts (from pc and poly) are correctly extracted.

From the .spice file (-mm and -mn options) it results:

```
R1 A B RPOLY 12.00
* R1=1080.0 (width=1.50U)
C1 B SS 0.6F
C2 A SS 0.6F
******************************************************************************
** Model Definitions for PSPICE
******************************************************************************
.model RPOLY RES (R=90.000)
```

and the number of squares SQ is correctly extracted, apart the factor 0.55 which must be attributed to the corner.

Or, in the distributed case (-mm, -mn and -d options)

```
XR1 A B SS D_RPOLY params: SQ=12.00 A=27.00
* R1=1080.0 (width=1.50U)
C1 B SS 0.6F
C2 A SS 0.6F
.subckt D_RPOLY 1 2 3 params: SQ={SQ} A={A}
```
and also the area A of \textit{pres} for the evaluation of the parasitic capacitance associated to \textit{pres} is correctly extracted.

In this example, the limits in the accuracy of the extraction are the ones previously discussed.
Step 3 - Corrections and Improvements (ext2spice.corr)

The Ext2spice.corr program provides the correction and improvement of the spice netlist generated by the Ext2spice program. In particular:

- eliminates, if present, all the capacitors the value of which is $0,0F^5$;
- calculates the value of resistors, both in the non-distributed and in the distributed case, by including in the evaluation the corners, the resistance of the poly adiacent the contact, the resistance of the contact, the perimeter component of the parasitic capacitance and the difference between the real and the drawn width; for each resistor the result of these calculations is also reported as a comments;
- converts the NRG, NRD and NRS parameters of the FETs into RG, RD and RS parameters, providing also a more accurate approximation, by including the resistance of the contacts and a separate coefficient for the resistance per square of polysilicon, ndiffusion and pdiffusion;
- locates the linear capacitors and provides these capacitors with a model that includes their dependence on the voltage and on the temperature.

Finally:

- by the introduction of an additional layer for each metal and by some further modification of the Technology file and of the Ext.defs file, it provide the extraction of the metal lines as resistors, including all the related parasitic capacitances.

Ext2spice.corr

The code of the Ext2spice.corr program is reported in the Appendix F. The program can be invoked in cascade to Ext2spice by using the following script:

```
#!/bin/csh
rm -f $1.cir
~ ext2spice -mm -mn -c .001 -g -d -D -S $1
cp $1.spice $1.spice.old
~ ext2spice.corr $1.spice.old > $1.spice
```

The old .spice file is saved as *nomefile.spice.old*

---

5 This function is necessary because in some cases the 0 value capacitors are not filtered by Ext2spice.
Improvements in the extraction of the Resistors

(a) - Corners
In order to take into account in the calculus of the value of the resistor all the corners present along pres, the number of corners of each resistor must be known. This number can be included in the label of each resistor. When Ext2spice.corr finds a resistor with a label Rxx_Ln (the label Rxx_Ln must be attached to pres), extracts n as the number of corners for that resistor. If the term _Ln is omitted from the label, the number of corners of that resistor is assumed to be zero. Ext2spice.corr provides also the necessary modifications in the .spice file in order to take into account this new parameter, both in the case of the non-distributed and distributed model.

(b) - Resistance of the contact and of the poly adjacent to the contact
The resistance of the contact pc and of the poly adjacent to the contact are included in the calculus. For the adjacent poly, the minimum length 1λ is assumed (plus 1λ for the distance between the real contact and the poly). The value of the resistance of the poly contact PC [Ohm] and of the resistance per square of the poly RSQPOLY [Ohm/square] must be given at the beginning of the Ext2spice.corr code. These values can be found in the General Process Specifications, in the Technology File of the Process or in the Mosis Parametric Test Results.

(c) - Perimeter component of the parasitic capacitance
In the case of the distributed model, the parasitic capacitance associated to the resistor is extracted. But Ext2spice provides only the area component of this capacitance. In order to extract also the perimeter component, the .MODEL lines are replaced with .PARAM lines and the capacitance is calculated according to the values of the parameters. The value of the areacap RACAP [F/m²] and perimcap RPCAP [F/m] must be given at the beginning of the Ext2spice.corr code. These values can be found in the General Process Specifications, in the Technology File of the Process or in the Mosis Parametric Test Results. Attention must be paid when the additional ss layer is used and pres lies is in a well, because the parasitic capacitance is calculated from pres to the layer ss and not from pres to the well, which is an error. The problem is that when the MAGIC extractor must choose between ss and well, it chooses ss (i.e. the layer with the lower planeorder). The solution is to remove the ss layer from all the areas where pres is overlapped to a well.

(d) - Real and drawn width
Ext2spice.corr provides the correction through the parameter RDW, which is the delta-width. The value of RDW [m] must be given at the beginning of the Ext2spice.corr code. This value can be found in the General Process Specifications, in the Technology File of the Process or in the Mosis Parametric Test Results.

(e) - Calculus of the value of the resistance
The value of the resistance is finally calculated through the equation:
\[
R = 2PC + 2RSQPOLY \frac{2LAMBDA}{W} + \frac{RSQ(SQ - 0.45NL)}{1 - \frac{RDW}{W}}
\]  

where \(W\) is the width of \(pres\) and \(NL\) is the number of corners. In the non-distributed case \(W\) is not available and a minimum width \(WMIN\) is assumed. The values of \(LAMBDA\) \([m]\), \(RSQ\) \([\text{Ohm/square}]\) and \(WMIN\) \([m]\) must be given at the beginning of the Ext2spice.corr code. These values can be found in the General Process Specifications, in the Technology File of the Process or in the Mosis Parametric Test Results.

**Example 7**

After the relabeling of the resistance of the Example 6 as \(R1\_L1^\wedge\), it follows from Ext2spice.corr in the non-distributed case:

\[
\begin{align*}
R100 & \ A \ B \ 1056.3 \\
C1 & \ B \ 1 \ 0.6F \\
C2 & \ A \ 1 \ 0.6F
\end{align*}
\]

and in the distributed case:

\[
\begin{align*}
XR1 & \ A \ B \ 1 \ D\_RPOLY \ \text{params:} \ SQ=12.00 \ A=27.00 \ NL=1 \\
& \ * \ XR1=1056.3 \ (W=1.5e-06, \ Weff=1.5e-06, \ Cpar=6.687e-15) \\
C1 & \ B \ 1 \ 0.6F \\
C2 & \ A \ 1 \ 0.6F
\end{align*}
\]

**Global Node Definitions**

**Distributed Model Definitions**

**Multiple element model for RPOLY**

**Unit element model for RPOLY**

```
**SUBCKT D\_RPOLY 1 2 3 params: SQ=(SQ) A=(A) NL=(NL)
XR1 4 5 E\_RPOLY params: SQ = (SQ/5) A = (A/5)
XR2 5 6 E\_RPOLY params: SQ = (SQ/5) A = (A/5)
XR3 6 7 E\_RPOLY params: SQ = (SQ/5) A = (A/5)
XR4 7 8 E\_RPOLY params: SQ = (SQ/5) A = (A/5)
XR5 8 9 E\_RPOLY params: SQ = (SQ/5) A = (A/5)
.ENDS
**SUBCKT E\_RPOLY 1 2 3 params: SQ=(SQ) A=(A) NL=(NL)
R1 2 \ (2*PC+2*RSQPOLY*2*LAMBDA/SQRT(A*1p/SQ)+RSQ*(SQ-NL*0.45)/(1-RDW/SQRT(A*1p/SQ)))
C1 3 \ (RACAP*A*1p/2+RPCAP/2*2*SQRT(A*1p*SQ))
C2 2 3 \ (RACAP*A*1p/2+RPCAP/2*2*SQRT(A*1p*SQ))
.ENDS
```
Improvements in the extraction of the FETs

In order to give a better approximation of the parasitic resistances, their values are directly calculated by Ext2spice.corr. On each FET line, the new expressions for RG, RD and RS replaces the old expressions for NRG, NRD, NRS.

(a) - Parasitic resistance RG
The following equation is used for the evaluation of RG:

\[
R_G = RSQPOLY \left( \frac{1}{N_{RG}} + \frac{4LAMBDA}{L} \right) + \frac{PC}{2} \frac{CLT \cdot LAMBDA}{L}
\]

(b) - Parasitic resistances RD and RS
The following equations are used to approximate the values of RD and RS.

For single finger n-channel FETs:

\[
R_D = NRSQ \left( N_{RD} - \frac{3 \cdot LAMBDA}{W} \right) + NDC \frac{2 \cdot CLT \cdot LAMBDA}{W}
\]

\[
R_S = NRSQ \left( N_{RS} - \frac{3 \cdot LAMBDA}{W} \right) + NDC \frac{2 \cdot CLT \cdot LAMBDA}{W}
\]

and for single finger p-channel FETs:

\[
R_D = PRSQ \left( N_{RS} - \frac{3 \cdot LAMBDA}{W} \right) + PDC \frac{2 \cdot CLT \cdot LAMBDA}{W}
\]
\[ RS = \text{PRSQ}\left( \text{NRS} - \frac{3 \cdot \text{LAMBDA}}{W} \right) + \text{PDC} \frac{2 \cdot \text{CLT} \cdot \text{LAMBDA}}{W} \]  

(8d)

For multifinger n-channel FETs:

\[ \text{RD} = \text{NRSQ}\left( \text{NRD} \cdot M^2 - \frac{\text{M} \cdot \text{LAMBDA}}{W} - \frac{2 \cdot \text{LAMBDA}}{W} \right) \frac{1}{M} + \text{NDC} \frac{2 \cdot \text{CLT} \cdot \text{LAMBDA}}{W} \]  

(9a)

\[ RS = \text{NRSQ}\left( \text{NRS} \cdot M^2 - \frac{\text{M} \cdot \text{LAMBDA}}{W} - \frac{2 \cdot \text{LAMBDA}}{W} \right) \frac{1}{M} + \text{NDC} \frac{2 \cdot \text{CLT} \cdot \text{LAMBDA}}{W} \]  

(9b)

and for multifinger p-channel FETs:

\[ \text{RD} = \text{PRSQ}\left( \text{NRS} \cdot M^2 - \frac{\text{M} \cdot \text{LAMBDA}}{W} - \frac{2 \cdot \text{LAMBDA}}{W} \right) \frac{1}{M} + \text{PDC} \frac{2 \cdot \text{CLT} \cdot \text{LAMBDA}}{W} \]  

(9c)

\[ RS = \text{PRSQ}\left( \text{NRS} \cdot M^2 - \frac{\text{M} \cdot \text{LAMBDA}}{W} - \frac{2 \cdot \text{LAMBDA}}{W} \right) \frac{1}{M} + \text{PDC} \frac{2 \cdot \text{CLT} \cdot \text{LAMBDA}}{W} \]  

(9d)

The \( M^* \lambda/W - 2*\lambda/W \) terms take into account the resistance of the diffusion part of the real contact. The \( XDC/(W/\text{CLT}/\lambda) \) term approximates the contact resistance and \( XDC \) must be NDC for n-channel and PDC for p-channel. The calculus is subject to the approximation on the original NRS (or NRD). The minimum distance between the channel and \( ndc \) (or \( pdc \)) is assumed. The contact resistance results overestimated for small \( L \). Note that, during the simulation, SPICE considers the RS, RG, RD as values per finger, dividing these values by \( M \) for the simulation of the overall FET, which is the correct approach.

The values of \( \text{NRSQ} \) and \( \text{PRSQ} \) [Ohm/square], respectively the resistance per square of the ndiffusion and of the pdiffusion, as well as the values of \( \text{NDC} \) and \( \text{PDC} \) [Ohm], respectively the resistance of the contacts \( ndc \) and \( pdc \), must be given at the beginning of the Ext2spice.corr code. These values can be found in the General Process Specifications, in the Technology File of the Process or in the Mosis Parametric Test Results.

**Example 8**

For the FET of **Example 4** it follows:

\[
\begin{align*}
\text{Mx B C A SS CMOSN W}=3.00U & \ 
\text{L}=1.20U \ 
\text{AD}=6.30P \ 
\text{PD}=10.20U \ 
\text{AS}=6.30P \ 
\text{PS}=10.20U \\
+ \ 
\text{RG}=7.4375 \ 
\text{RD}=3 \ 
\text{RS}=3 \\
\text{C1 B SS 1.8F} \\
\text{C2 C SS 1.1F}
\end{align*}
\]

and for the FET of **Example 5** it follows:

\[
\begin{align*}
\text{M1 A C B SS CMOSN M}=3 \ 
\text{W}=3.00U \ 
\text{L}=0.60U \ 
\text{AD}=3.30P \ 
\text{PD}=6.20U \ 
\text{AS}=6.04P \ 
\text{PS}=7.03U
\end{align*}
\]
Improvements in the extraction of the Linear Capacitors

The improvement in the extraction of the linear capacitors consists in the addition of the dependence of the capacitance on the voltage and on the temperature. In order to achieve this result, all the linear capacitors must be modeled by using a .MODEL statement in the .spice file. Till now it was not possible to distinguish between parasitic and linear capacitors. But adding the following line in the extract section of the technology file:

```
fet wcap cwnsd 1 wcap $$G_Vsub! 0 0
```

each linear capacitor will be represented both as \textit{cap} and as \textit{fet} element in the .EXT file.

If the following lines are added in the Ext.defs support file,

```
def wcap LCAP C c
distrib wcap 0.475 0.0 0
```

Ext2spice will extract, for each linear capacitor, a "duplicated" capacitor (apart the value) and models each duplicate with LCAP.

Ext2spice.corr deletes, by comparison, all these "duplicates" and models the corresponding original capacitors (the value of which is more accurate) through the model LCAP. The original and correct values of the capacitors are preserved and the .MODEL line for LCAP is added. The values of the voltage parameters \textit{VCI} and \textit{VC2} and of the temperature parameters \textit{TC1} and \textit{TC2} used in the .MODEL statement can be found in the General Process Specifications or in the Mosis Parametric Test Results.

\textbf{Example 9}

For the linear capacitor of \textbf{Example 3} it follows:

```
C2 A B LCAP 2097.8F
```

******************************************************************************

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* Global Node Definitions

.MODEL LCAP CAP (C=1 VC1=0.001 VC2=0.0005 TC1=0 TC2=0)

Extraction of the resistance of the Metal lines

The extraction of the resistance of the metal lines requires that, as in the case of poly resistors, a part or all the line becomes visible as a *fet* to the MAGIC extractor. A possible and effective approach is the introduction of an additional layer for each metal, which must be invisible at the .CIF extraction level and which represents the active area of this "fet". In order to preserve all the parasitic capacitances related to the lines, the active area of the "fet" must be as small as possible, i.e. must be a square along the line. The resistance of the line is then extracted by evaluating the number of squares of the "drain" and of "source" of this "fet" (which equals with good approximation the number of squares of the line). The use of more squares suitably distributed along the line allows a more accurate extraction.

In order to use the additional layers, the following changes must be made in the Technology File.

In the *types* section, three additional layers (*mlres*, *m2res* and *m3res*) for the three metals (*m1*, *m2* and *m3*) must be added:

```
metal1 mlres,m1r
metal2 m2res,m2r
metal3 m3res,m3r
```

In the *styles* section, to "see" the additional layers at the MAGIC layout level; the lines

```
m1res 20
m1res 12
m1res 13
m2res 21
m2res 12
m2res 13
m3res 22
m3res 12
m3res 13
```

must be added.

In the *connect* section, to set the connections, the lines:

```
m1,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1
m1,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1
m2,m2c/m2,m3c/m2,m3c/m2 m2,m2c/m2,m3c/m2,m3c/m2
m3,m3c/m3 m3,m3c/m3
```
must be replaced with the lines:

```
m1,m1res,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1
ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1
m2,m2res,m2c/m2,m3c/m2,m3c/m2,m2c/m2,m3c/m2,m3c/m2
m3,m3res,m3c/m3,m3c/m3
```

In the `cifoutput` section, to let the layer be invisible at the .CIF level (i.e to let it be “like” the original metal):

```
layer CMF m1,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1
labels m1,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1
layer CMS m2,m2c/m2,m3c/m2,m3c/m2
labels m2,m2c/m2,m3c/m2,m3c/m2
layer CMT m3,m3c/m3
labels m3,m3c/m3
```

must be replaced with the lines:

```
layer CMF m1,m1res,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1
labels m1,m1res,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1
layer CMS m2,m2res,m2c/m2,m3c/m2,m3c/m2
labels m2,m2res,m2c/m2,m3c/m2,m3c/m2
layer CMT m3,m3res,m3c/m3
labels m3,m3res,m3c/m3
```

In the `extract` section, to enable the extraction of the "fet" diffusion areas, the `resistclass` lines must be modified as follows (this change does not affect the calculus of the real FET areas, if the Ext.defs support file is modified as described later):

```
# REPLACED resist (ndiff,anres,ndc/a,nsd,nsc/a)/active 1900
resist ((ndiff,ndc/a)/active,(pdiff,pdc/a)/active) 0
# REPLACED resist (pdiff,apres,pdc/a,psd,psc/a)/active 2000
resist (m1/metal1,m2/metal2,m3/metal3) 0
```

while the old `resistclass` lines containing `m1`, `m2` and `m3` must be deleted.

Finally, to enable the "fet" extraction, the following lines

```
fet m1res m1,ndc/m1,nsc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1 2 m1res $$G_Vsub! 0 0
fet m2res m2,m2c/m2,m3c/m2,m3c/m2 2 m2res $$G_Vsub! 0 0
fet m3res m3,m3c/m3 2 m3res $$G_Vsub! 0 0
```

must be added. Note that all the contacts have been included in the evaluation of the areas of the fets for the extraction of the resistances.
In order to extract the new "fet" as a resistance, the following lines must be added in the Ext.defs support file. Basically the idea is to extract the areas of the new "fet" by considering these as p-channel FETs. The areas of the real p-channel FETs is extracted by considering these as n-channel FETs. This approach, relevant only at the Ext2spice level, has no consequence for the final correct extraction of the areas of all real FETs.

The line:
```plaintext
def pfet CMOSP M p
```
must be replaced with the line:
```plaintext
def pfet CMOSP M n
```
and the lines:
```plaintext
def mlres RM1 M p
def m2res RM2 M p
def m3res RM3 M p
```
must be added.

Now Ext2spice.corr provides the accurate extraction of the resistances associated to the metal lines from the "drain" and "source" areas of these "fets". The values of $RSQM1$, $RSQM2$, and $RSQM3$ [Ohm/square], respectively the resistance per square of $m1$, $m2$ and $m3$, must be given at the beginning of the Ext2spice.corr code. These values can be found in the General Process Specifications, in the Technology File of the Process or in the Mosis Parametric Test Results.

Drawback of this technique is that the line, at the MAGIC layout level, doesn't appear connected so that the check of the connections must be made before this modification, or the lines in the `connect` section must be suitably switched.

**Example 10**

![Diagram](https://example.com/diagram.png)

In this example we have three couples of minimum wide lines in the three different metals ($m1$, $m2$, $m3$). A small square of the correct type (respectively $m1res$, $m2res$, $m3res$) has been introduced in the middle of each line, thus dividing the line in two parts. The square is not visible here be-
cause the picture is extracted from the .CIF file, and, as required, the layers
\((m1res, m2res, m3res)\) are not visible at the .CIF level. From Step 2, with
regards to the fet elements, it follows:

M1 A1 4 B1 \$G\_Vsub! RM1 W=0.90U L=0.00U AD=21.87P PD=50.40U AS=22.14P PS=51.00U
+ NRG=1.00 NRD=27.00 NRS=27.33
M2 A2 7 B2 \$G\_Vsub! RM1 W=0.90U L=0.00U AD=21.87P PD=50.40U AS=22.14P PS=51.00U
+ NRG=1.00 NRD=27.00 NRS=27.33
M3 A3 10 B3 \$G\_Vsub! RM2 W=0.90U L=0.00U AD=21.87P PD=50.40U AS=22.14P PS=51.00U
+ NRG=1.00 NRD=27.00 NRS=27.33
M4 A4 13 B4 \$G\_Vsub! RM2 W=0.90U L=0.00U AD=21.87P PD=50.40U AS=22.14P PS=51.00U
+ NRG=1.00 NRD=27.00 NRS=27.33
M5 A5 16 B5 \$G\_Vsub! RM3 W=1.50U L=0.00U AD=34.65P PD=55.80U AS=40.50P PS=63.60U
+ NRG=1.00 NRD=15.40 NRS=18.00
M6 A6 19 B6 \$G\_Vsub! RM3 W=1.50U L=0.00U AD=34.65P PD=55.80U AS=40.50P PS=63.60U
+ NRG=1.00 NRD=15.40 NRS=18.00

And from Step 3:

RM1 A1 B1 3.87333
RM2 A2 B2 3.87333
RM3 A3 B3 3.87333
RM4 A4 B4 3.87333
RM5 A5 B5 1.72
RM6 A6 B6 1.72
C2 A6 SS 1.0F
C5 B5 SS 1.1F
C8 A5 SS 1.0F
C9 B2 SS 2.7F
C10 B6 B5 0.8F
C11 B2 B1 0.4F
C13 A2 SS 2.7F
C14 A6 A5 0.7F
C16 B4 SS 1.4F
C19 A2 A1 0.4F
C20 B4 B3 0.4F
C21 A4 SS 1.4F
C22 SS B1 2.2F
C23 B3 SS 1.4F
C24 B6 SS 1.1F
C25 A4 A3 0.4F
C26 A3 SS 1.4F
C28 SS A1 2.2F

The accuracy of this evaluation, which includes the parasitic capacitances
between the lines lying in the same plane, can be easily verified.

\textbf{Conclusions}

With a relatively simple modification of the Technology File (see Appendix G for a summary of
the modifications), and with the use of the Magic Extract function, of the Ext2spice program
(and its support file Ext.defs) and of the Ext2spice.corr program, it is possible to extract the SPICE netlist from a MAGIC based layout with an improved accuracy. The limit in this approach is that every technology requires the modification of the corresponding Technology File (together with some minor changes in the two programs). The extraction of the capacitances is strongly dependent on the accuracy of the parameters from the Technology File, which must be verified by the simulations (for example by using the Maxwell simulator) and updated by using the beta versions released by MOSIS.
Appendix A - Connect and extract sections

connect
    nwell,cowell,ns,cwc,nsd,cwnsd,cwc/a nwell,cowell,ns,cwc,nsd,cwnsd,cwc/a
    pwell,psc,psd pwell,psc,psd
    m1,nndc/m1,ns/cwc/m1,cwc/m1,pdc/m1,psc/m1,nsd/m1,cwc/m1,pc/m1,m2/m1
        m3,m2/m1,ns/cwc/m1,cwc/m1,pdc/m1,psc/m1,nsd/m1,cwc/m1,pc/m1,m2/m1
    m3,m3c/m3 m3,m3c/m3
    ndiff,anres,ndc/a ndiff,anres,ndc/a
    pdiff,apres,pdc/a pdiff,apres,pdc/a
    ndiff,anres,nsd ndc/a,nsd/a,psc/a
    pdiff,apres,psd pdc/a,nsd/a
    poly,pres,wcap,nfet,wcap,pfet,pc/a poly,pres,wcap,nfet,wcap,pfet,pc/a
    gc poly,pres,wcap
    gv1 m1,m2
    gc ndiff,anres,pdiff,apres,nsd,psd,m1
    m3,m3c/m3 m3,m3c/m3
    gv2 m2,m3
    pad m1,m2,m3
end

extract
    style HP0.5um(hpcmos14tb)from:N81D
    cscale 1
    lambda 30
    step 100
    sidehalo 8
    planeorder well 0
    planeorder implant 1
    planeorder select 2
    planeorder active 3
    planeorder metal1 4
    planeorder metal2 5
    planeorder metal3 6
    planeorder oxide 7
    planeorder xp 8
    planeorder comment 9
    planeorder contact 10
    planeorder via1 11
    planeorder via2 12
    resist (ndiff,anres,ndc/a,nsd,nsc/a)/active 1900
    resist (pdiff,apres,pdc/a,psd,psc/a)/active 2000
    resist (poly,pres,wcap,pc/a,wcap,pc/a,nfet,pfet)/active 1800
    resist (m1,nndc/m1,ns/cwc/m1,cwc/m1,pdc/m1,psc/m1,nsd/m1,cwc/m1,pc/m1,m2/m1,m2c/m1)/metal1 70
    resist (m2,m2c/m2,m3c/m2,m3c/m2)/metal2 70
    resist (m3,m3c/m3,pad)/metal3 50
    contact ndc 4 2000
contact pdc 4 2000
contact pc 4 1700
contact m2c 4 700
contact m3c 4 360

# wellcap
overlap wcap capwell 207.900

# nwell, cwell, pwell
areacap (nwell, cwell)/well 8.460
areacap (cwell)/well 8.460

# ndiff
areacap (anres)/active 48.330
overlap (anres)/active ~space/w 48.330
perimc (anres)/active ~(anres)/active 67.800
sideoverlap (anres)/active ~(anres)/active ~space/w 67.800
# MODEL HANDLES THIS: areacap (ndiff, anres, ndc/a)/active 48.330
# MODEL HANDLES THIS: overlap (ndiff, anres, ndc/a)/active ~space/w 48.330
# MODEL HANDLES THIS: perimc (ndiff, anres, ndc/a)/active ~(ndiff, anres, ndc/a, nfet, pfet)/active 67.800
# MODEL HANDLES THIS: sideoverlap (ndiff, anres, ndc/a)/active ~(ndiff, anres, ndc/a, nfet, pfet)/active ~space/w 67.800

# pdiff
areacap (apres)/active 84.600
overlap (apres)/active ~space/w 84.600
perimc (apres)/active ~(apres)/active 67.500
sideoverlap (apres)/active ~(apres)/active ~space/w 67.500
# MODEL HANDLES THIS: areacap (pdiff, apres, pdc/a)/active 84.600
# MODEL HANDLES THIS: overlap (pdiff, apres, pdc/a)/active ~space/w 84.600
# MODEL HANDLES THIS: perimc (pdiff, apres, pdc/a)/active ~(pdiff, apres, pdc/a, nfet, pfet)/active 67.500
# MODEL HANDLES THIS: sideoverlap (pdiff, apres, pdc/a)/active ~(pdiff, apres, pdc/a, nfet, pfet)/active ~space/w 67.500

# polycap

# poly
sidewall (poly, pres, wcap, pc/a, wcap, pc/a)/active ~(poly, pres, wcap, pc/a, wcap, pc/a)/active ~poly, pres, wcap, pc/a, wcap, pc/a)/active (poly, pres, wcap, pc/a, wcap, pc/a)/active 2.136
areacap (poly, pres, wcap, pc/a, wcap, pc/a)/active 7.920
overlap (poly, pres, pc/a, pc/a)/active ~space/w 7.920
perimc (poly, pres, wcap, pc/a, wcap, pc/a)/active ~(poly, pres, wcap, pc/a, wcap, pc/a)/active 15
sideoverlap (poly, pres, wcap, pc/a, wcap, pc/a)/active ~(poly, pres, wcap, pc/a, wcap, pc/a)/active ~space/w 15
# MODEL HANDLES THIS: overlap (nfet)/active (ndiff, anres, ndc/a)/active 324.360
# MODEL HANDLES THIS: sideoverlap (nfet)/active ~(nfet)/active (ndiff, anres, ndc/a)/active 15
# MODEL HANDLES THIS: overlap (pfet)/active (pdiff, apres, pdc/a)/active 308.160
# MODEL HANDLES THIS: sideoverlap (pfet)/active ~(pfet)/active (pdiff, apres, pdc/a)/active 15

# poly2

# metal1
sidewall (m1, ndc/m1, nsc/m1, cwc/m1, pdc/m1, psc/m1, cwc/m1, pc/m1, m2c/m1)/metal1 ~(m1, ndc/m1, nsc/m1, cwc/m1, pdc/m1, psc/m1, cwc/m1, pc/m1, m2c/m1)/metal1 ~~~(m1, ndc/m1, nsc/m1, cwc/m1, pdc/m1, psc/m1, cwc/m1, pc/m1, m2c/m1)/metal1 ~~~(m1, ndc/m1, nsc/m1, cwc/m1, pdc/m1, psc/m1, cwc/m1, pc/m1, m2c/m1)/metal1 (m1, ndc/m1, nsc/m1, cwc/m1, pdc/m1, psc/m1, cwc/m1, pc/m1, m2c/m1)/metal1 6.690
areacap (m1,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1 2.700

#metal-sub blocked by --space/a,--space/c
overlap (m1,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1 --space/w 2.700 --space/a
perimc (m1,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1
--(m1,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1 9.300
sideoverlap (m1,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1
--(m1,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1 --space/w 9.300 --space/a

#metal-diff blocked by --space/c
overlap (m1,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1
(ndiff,anres,ndc/a)/active 2.790
sideoverlap (m1,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1
--(m1,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1
(ndiff,apres,ndc/a)/active 9.300
overlap (m1,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1
(pdiff,apres,ndc/a)/active 2.790
sideoverlap (m1,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1
--(m1,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1
(pdiff,apres,ndc/a)/active 9.300

#metal-polycap blocked by poly

#metal-polycap blocked by poly2
overlap (m1,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1
(poly,pres,wcap,pc/a,wcap,pc/a,nfet,pfet)/active 5.400
sideoverlap (m1,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1
--(m1,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1
(poly,pres,wcap,pc/a,wcap,pc/a,nfet,pfet)/active 15
sideoverlap (poly,pres,wcap,pc/a,wcap,pc/a,nfet,pfet)/active
--(poly,pres,wcap,pc/a,wcap,pc/a,nfet,pfet)/active
(m1,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1 15

#metal-polycap blocked by poly2 not blocked

#metal2
sidewall (m2,m2c/m2,m3c/m2)/metal2 --(m2,m2c/m2,m3c/m2)/metal2 --(m2,m2c/m2,m3c/m2)/metal2
(m2,m2c/m2,m3c/m2)/metal2 9.090
areacap (m2,m3c/m2)/metal2 1.260

#metal2-sub blocked by --space/a,--space/m1,poly2/polycape
overlap (m2,m3c/m2)/metal2 --space/w 1.260 --space/a,--space/m1
perimc (m2,m2c/m2,m3c/m2)/metal2 --(m2,m2c/m2,m3c/m2)/metal2 5.100
sideoverlap (m2,m2c/m2,m3c/m2)/metal2 --(m2,m2c/m2,m3c/m2)/metal2 --space/w 5.100
--space/a,--space/m1
overlap (m2,m3c/m2)/metal2 (ndiff,anres,ndc/a)/active 1.260 --space/m1
sideoverlap (m2,m2c/m2,m3c/m2)/metal2 --(m2,m2c/m2,m3c/m2)/metal2 (ndiff,anres,ndc/a)/active
5.100 --space/m1
overlap (m2,m2c/m2,m3c/m2)/metal2 (pdiff,apres,pdc/a)/active 1.260 --space/m1
sideoverlap (m2,m2c/m2,m3c/m2)/metal2 --(m2,m2c/m2,m3c/m2)/metal2 (pdiff,apres,pdc/a)/active
5.100 --space/m1

#metal2-polycap by --space/m1,poly
sideoverlap (m3,m3c/m3,pad)/metal3 ~m3,3c/m3,pad)/metal3
  (m1,ndc/m1,nse/m1,cwc/m1,pdc/m1,pse/m1,cwc/m1,p2c/m1,p2c/m1)/metal1  5.400
  -space/m2
sideoverlap (m1,ndc/m1,nse/m1,cwc/m1,pdc/m1,pse/m1,cwc/m1,p2c/m1,p2c/m1)/metal1
  (m1,ndc/m1,nse/m1,cwc/m1,pdc/m1,pse/m1,cwc/m1,p2c/m1,p2c/m1)/metal1
  (m3,m3c/m3,pad)/metal3  5.400 -space/m2

#M3->M2
overlap (m3,pad)/metal3 (m2,m2c/m2,m3c/m2)/metal2  4.230
sideoverlap (m3,m3c/m3,pad)/metal3 ~m3,m3c/m3,pad)/metal3 (m2,m2c/m2,m3c/m2)/metal2 10.800
sideoverlap (m2,m2c/m2,m3c/m2)/metal2 ~m2,m2c/m2,m3c/m2)/metal2 (m3,m3c/m3,pad)/metal3
  10.800

#metal4

#fets
fet  pfet pdiff,pdc 2 pfet Vdd! nwell 15 308
fet  pfet pdiff,pdc 1 pfet Vdd! nwell 15 308

fet  nfet ndiff,ndc 2 nfet Gnd! pwell 15 324
fet  nfet ndiff,ndc 1 nfet Gnd! pwell 15 324

fetresis pfet linear 15658
fetresis pfet saturation 15658
fetresis nfet linear 4267
fetresis nfet saturation 4267

end
Appendix B - Linear capacitor fringing capacitance

The first step to determine the fringing component for the capacitance of interest is to check the MOSIS General Process Specifications and the MOSIS Parametric Test Results relative to the technology used. In our case both the General Process Specifications and the Parametric Test Results don’t report the value for the required fringing component. The Parametric Test Results report only the fringing component from poly to $N^*$, as shown in Fig. B1 but without specifying if this value is relative to the gate oxide thickness (90Å for this process), to the linear capacitor oxide thickness (150Å for this process) or, with less probability, to the field oxide thickness (3500Å for this process). It can also be observed that the value is subject to a strong dispersion and that its average value 164.53 aF/µm is much higher than the value 15/03.=50 aF/µm reported in the technology file for the fringing component from poly to well. In order to get a more reliable value for the required fringing component, an investigation based on the geometrical parameters given by the General Process Specifications (i.e. poly thickness T and oxide thickness D in the case of interest) must be carried out.

The following equations for the evaluation of the fringing component from the two sides of a transmission line in the oxide can be found in the literature.

\[ \varepsilon_{\text{ox}} \frac{2\pi}{\ln \left[ 1 + \frac{2D}{T} \left( 1 + \sqrt{1 + \frac{T}{D}} \right) \right]} - \frac{T}{2D} \] (B1)


\[ \varepsilon_{\text{ox}} \frac{2\pi}{\ln \left[ 1 + \frac{2D}{T} \left( 1 + \sqrt{1 + \frac{T}{D}} \right) \right]} - \frac{T}{4D} \] (B2)


Footnote 6: For the dielectric constant of the oxide, the value $\varepsilon_{\text{ox}} = 3.453133 \times 10^{-11} \text{F/m}$ is assumed.
\[
\varepsilon_{\text{ox}} \left\{ 0.398 \frac{T}{D} \left( 1 + \ln \frac{2D}{T} \right) + 1.393D + 0.667 \ln \left[ \frac{W}{D} + 0.398 \frac{T}{D} \left( 1 + \ln \frac{2D}{T} \right) + 1.444 \right] \right\}
\] (B3)


\[
\varepsilon_{\text{ox}} \left[ 0.15 \frac{W}{D} + 2.8 \left( \frac{T}{D} \right)^{0.222} \right]^{0.728}
\] (B4)


\[
\varepsilon_{\text{ox}} \left[ 0.15 \frac{W}{D} + 2.8 \left( \frac{T}{D} \right)^{0.222} + 4.12 \frac{L}{D} \left( \frac{T}{D} \right)^{0.728} \right]
\] (B5)


\[
\varepsilon_{\text{ox}} \left[ 0.15 \frac{W}{D} + 2.8 \left( \frac{T}{D} \right)^{0.222} + 4.12 \frac{L}{D} \left( \frac{T}{D} \right)^{0.728} \right]
\] (B6)

In order to evaluate the accuracy of these equations, each one characterized by a limited range of application, a simulation has been carried out by using the recently purchased Maxwell 2D software from Ansoft. A comparison in the case of a \(W=1\mu m\) and \(W=10\mu m\) width line with a \(D=1000\AA\) oxide thickness versus the line thickness \(T\) is shown in Fig.s B2. In the case of the Maxwell simulation, a term \(\varepsilon_{\text{ox}}W/D\) (i.e. the area component) has been subtracted from the simulation result.

![Comparison of the fringing component as results from the Eq.s B1 to B6 and from the Maxwell simulator at equal geometry: (a) \(W=1\mu m\); (b) \(W=10\mu m\).](image)

Fig. B2 - Comparison of the fringing component as results from the Eq.s B1 to B6 and from the Maxwell simulator at equal geometry: (a) \(W=1\mu m\); (b) \(W=10\mu m\).
From Fig. B2(a) (W=1\mu m) it can be observed that a good prediction can be expected only from the first (B1), the third (B3) and the sixth (B6) equation, depending in any case on the range of the thickness T. From Fig. B2(b) (W=10\mu m) the prediction is much worse and only the third (B3) equation seems to give a good prediction, limited to the first range of the thickness T.

An equation which can give a more accurate prediction in the range W=0.05\mu m+100\mu m is proposed:

\[
\varepsilon' \left( \frac{W}{D} \right) = \frac{2.91}{0.83 + \left( \frac{0.25 D}{W} \right)^{0.25}} + \frac{3.3}{1 + \left( \frac{3 D}{W} \right)^{0.25}} + 3 \left( \frac{D}{W} \right)^{0.4} \ln \left[ 1 + \left( \frac{D}{2 W} \right)^{1.05} \right] \ln \left[ 1 + \left( \frac{T}{D} \right)^{0.72} \right]
\] (B7)

The accuracy of this equation in the evaluation of the fringing component can be observed from the comparisons shown in Fig. B3.

![Graphs showing comparisons of fringing components](image)

**Fig. B3** - Comparison of the fringing component as results from the Eq. B7 and from the Maxwell simulator at equal geometry: (a) W=0.05\mu m; (b) W=1\mu m; (c) W=10\mu m; (d) W=100\mu m.
In our case, the *poly* thickness T and *oxide* thickness D for the linear capacitor can be determined from the General Process Specifications. Introducing these values (T=250nm, D=15nm) in the Eq.B7 and by multiplying for a factor 0.5 (fringing of a single side), the coefficient for the fringing capacitance can be determined and it depends on the "width" as shown in Fig.B4. It is worth noting that this coefficient increases as W increases, and the reason is that the fringing component includes the upper plane of the *poly*, which increases with the width W. By considering that the typical width(s) of the linear capacitors lies between 10μm and 50μm, a value close to 40 aF/λ can be chosen.

![Fig. B4 - Fringing component of poly to cwnsd according to Eq.B7, expressed in aF/λ.](image)
Appendix C - Introduction of the additional plane and layer

The following line must be introduced at the beginning of the *planes* section of the technology file:

```
substrate,ss
```

The following line must be introduced at the beginning of the *types* section of the technology file:

```
ss  ss
```

No line is added in the *styles* section of the technology file, if we don't want to see the new layer.

No line is added or modified in the *cifoutput* section of the technology file, if we don't want to see the new layer during the CIF extraction.

Some lines must be modified in the extract section:

```
planeorder well 0
planeorder implant 1
planeorder select 2
planeorder active 3
planeorder metal1 4
planeorder metal2 5
planeorder metal3 6
planeorder oxide 7
planeorder xp 8
planeorder comment 9
planeorder contact 10
planeorder via1 11
planeorder via2 12
```

becomes:

```
planeorder ss 0
planeorder well 1
planeorder implant 2
planeorder select 3
planeorder active 4
planeorder metal1 5
planeorder metal2 6
planeorder metal3 7
planeorder oxide 8
planeorder xp 9
planeorder comment 10
planeorder contact 11
planeorder via1 12
planeorder via2 13
```
Each *areacap* and *perimcap* line in the *extract* section must be replaced with a corresponding *overlap* and *sideoverlap* line as follows:

```plaintext
# REPLACED areacap (cwell)/well  8.460
overlap (nwell,cwell)/well ss/ss  8.460

# REPLACED areacap (poly,pres,wcap,pc/a,wcap,pc/a)/active  7.920
overlap (poly,pres,wcap,pc/a,wcap,pc/a)/active ss/ss  7.920 ~space/w

# REPLACED perimc (poly,pres,wcap,pc/a,wcap,pc/a)/active ~(poly,pres,wcap,pc/a,wcap,pc/a)/active 15
sideoverlap (poly,pres,wcap,pc/a,wcap,pc/a)/active ~(poly,pres,wcap,pc/a,wcap,pc/a)/active ss/ss 15 ~space/w

# REPLACED areacap (m1,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1)/metal1  2.700
overlap (m1,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,m1,m2c/m1)/metal1 ss/ss  2.700 ~space/w,~space/a

# REPLACED perimc (m1,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1)/metal1 ~(m1,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,m1,m2c/m1)/metal1  9.300
sideoverlap (m1,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,m2c/m1)/metal1 ~(m1,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,m1,m2c/m1)/metal1 ss/ss  9.300 ~space/w,~space/a

# REPLACED areacap (m2,m3c/m2)/metal2  1.260
overlap (m2,m3c/m2)/metal2 ss/ss  1.260 ~space/w,~space/a

# REPLACED perimc (m2,m2c/m2,m3c/m2)/metal2 ~(m2,m2c/m2,m3c/m2)/metal2  5.100
sideoverlap (m2,m2c/m2,m3c/m2)/metal2 ~(m2,m2c/m2,m3c/m2)/metal2 ss/ss  5.100 ~space/w,~space/a

# REPLACED areacap (m3,pad)/metal3  0.810
overlap (m3,pad)/metal3 ss/ss  0.810 ~space/w,~space/a

# REPLACED perimc (m3,m3c/m3,pad)/metal3 ~(m3,m3c/m3,pad)/metal3  3.300
sideoverlap (m3,m3c/m3,pad)/metal3 ~(m3,m3c/m3,pad)/metal3 ss/ss  3.300 ~space/w,~space/a
```

Finally, for the correction of the error of type 3:

```plaintext
# REPLACED sideoverlap (poly,pres,wcap,pc/a,wcap,pc/a)/active ~(poly,pres,wcap,pc/a,wcap,pc/a)/active ~space/w  15
sideoverlap (poly,pres,wcap,pc/a,wcap,pc/a)/active ~(poly,pres,wcap,pc/a,wcap,pc/a,cwnsd)/active ~space/w  15
# ADDED
sideoverlap (wcap)/active (cwnsd)/active ~space/w  40
```
Appendix D - Ext2spice Manual (by Lloyd Clonts)

NAME
ext2spice - convert from .ext format to .spice format

SYNOPSIS
ext2spice [-c cmin][-cs cdiv][-cdghmnrvBCLMNSV] extfile

DESCRIPTION
ext2spice reads a file in .ext format and creates a new file in .spice format. The output consists of a list of
subcircuits, containing diodes, capacitors, resistors, and transistors. Each .ext file is transformed into a
spice subcircuit.

Within an ext.defs file, the values of the different elements can be defined to automatically generate
PSPICE or HSPICE .model statements in the SPICE file. In addition, an option is available to create a .cir
file (if it does not exist) which calls the .spice file. Filescan be re-extracted without modifying the stimu-
loses which are in the .cir file. In addition, statements can be defined in the the ext.defs files which are
"echoed" into the .cir file. The echo statements are very useful to predefine .include statements or setup
parameters.

OPTIONS
-c cmin Do not output capacitors that are less than cmin femptoFarads. Default cmin is 1.0 FEMPTO-
Farads.
-cd Do not output any parastic capacitors.
-cs cdiv Scale all parasitic by value cdiv. This option is for processe with small lambda values where the
tech file has its output values scaled. For these technology file modifications, area type parasitics
should be scaled by a factor of the squareroot of cdiv and fringing parasitics should be scaled by a
factor of cdiv. This modification may be necessary because the magic extractor has a lower limit
on parasitic value of 1 FF/lambda.
-d Generate distributed models for devices and create WELL diodes and the GLOBAL node Vsub for
parasitics.
-D Generate WELL diodes and create a GLOBAL node Vsub for parasitics.
-g Create a separate .cir file to put the stimuliuses, to define analysis parameters, and other state-
ments (see echo in ext.defs).
-h Output devices in HSPICE format instead of PSPICE format.
-m Merge parallel/serial devices into an equivalent device. Applies for resistors, capacitors, and tran-
sistors.
-mm Transistors are defined with a multiplicity (M=). Averages are done on the widths, perimeters, and
areas to create an equivalent device.
Transistors are defined in terms of the geo= parameter. The MODEL statement for the device will
define the area/perimeter of the drain and sources for the transistor.

Output the NRG, NRD, and NRS spice parameter for each transistor.

Print all of the node names with their node numbers at the end of each subckt definition.

Output file in CaZM format.

Output transistor length as part of the MODEL name for MOS devices. For example, a p-transistor
of length = 5 will be given the MODEL name CMOSP_5.

Print all of the node names with their node numbers into a separate .merge file.

Print the root node names with their node numbers into a separate .names file.

Output all node and device names as strings instead of NODES.

Make well errors global nodes and insert voltage sources to drive these wells. This feature cor-
rects the floating well errors that occur when wells are not defined for certain devices. With this
option, simulation is possible, but CIF may not correctly generate wells to prevent them from
floating.

NODE CLEANUP OPERATIONS
Ext2spice checks to see if node is being used in the circuit. If the nodes is not used by device or not a
part of a subcircuit terminal list, then all parasitics related to the nodes are deleted. Shorted parasitic ca-
pacitors are also deleted.

NODE NUMBERS AND NAMES
An additional feature is in the identification of node numbers. Most older SPICE programs prefer node
numbers rather than node names. To aid in circuit verification and identification, ext2spice will number
nodes based on a scheme similar to that used for numbering transistors. Namely, a node named 'Nx',
where x is an integer, will be identified as node x. For example, a node labeled as 'N13' will be identified
as node number 13. No special ending character is needed. If more than one identifier is used on a node,
the lowest numbered one is used. All identifiers appear in the node listings.

For better clarity, ext2spice attempts to find the best node name to be matched with a node number for
the circuit's node listing. The search criteria is the following going from lowest to highest priority:

1) Ext2spice names or names with last character as a "#".
2) Hierarchical names or names containing a "/".
3) Names in number format Nx.

To force nodes as a name instead of a number, a $NAME will cause a node to be called NAME in the
circuit regardless of the -S option.

NODE MERGING
In older versions, any node name ending in the character '!" is assumed to be universal, so ext2spice
electrically connected these nodes. For example, node 'Vdd!' in one cell is merged with the node 'Vdd!' in
all other cells, whether or not they are physically connected. This option was removed to prevent simulation inaccuracy. Nodes should not be merged if they are not physically connected. Names ending in the character '!' have no special interpretation in terms of merging except for well errors (NWerror! and PWerror!) and the global substrate node. The substrate node (Vsub!) is the only true global node which should exist. The well error nodes should be corrected.

GLOBAL WELL ERRORS

The NWerror! and PWerror! signals are generated as globals to indicate that a well material is missing from the layout. With the -V option, voltage sources are added at the end of the spice file to prevent the wells from floating and to allow simulation. However, an ERROR could be present in the layout. CIF generation will add wells for transistor not going into the native substrate. However, no guarantees can be made that a well is connected to the correct bias if the well is not present for the device. The layout should be corrected by inserting a well under the devices with the errors. Ext2spice generates an error when NWerror! or PWerror! signals are found in the layout.

GENERAL DEVICE LABELLING

The user can control what identification number ext2spice gives to any particular device. The program looks to see if the gate argument is of the form "device character"name". If that is the case, then in the .spice file, the fet will be identified by "device character"name". For example, a transistor label'M12' will be called M12 in the .spice file. A resistor labeled 'RC1' will be called RC1. The same applies for bipolar transistors 'Qname', diodes 'Dname', and capacitors 'Cname'. A transistor whose identification number is not specified will be issued one by ext2spice. Ext2spice will create new names if conflicts occur.

TRANSISTOR LABELLING

The terminals of the MOS transistor are the source and drain. At least one terminal attribute must be 'S' or 'D' to be correctly identified. Both terminals may be labeled, but one is sufficient. In order for magic to place terminal attributes in the .ext file, the label 'S$' or 'D$' must be placed on the junction between the channel and the diffusion. When merging parallel transistors, it is sufficient to label only one of the transistors in order to specify the identification name and source-drain orientation of the merged transistor. Both Mx and S or D labels should be placed on the same transistor. S or D labels on transistors other than the one with Mx labels may be lost during the merging process.

For HSPICE users, the geo parameter can be extracted using the terminal attributes. The designations becomes 'Dx$' and 'Sx$'. If x=1,2, or 3, then geo=1 for shared drains, x=2 indicates shared sources, and x=3 or x=B means both shared drains and sources. Also, drains labelled 'DD$' indicates shared drains, sources labelled 'SS$' indicates shared sources, and labeling both the drain with 'DD$' and the source with 'SS$' will give in a geo=3 condition. For regular extraction, the geo designications are ignored.

RESISTOR DISTRIBUTED MODELLING

For improved accuracy of the resistor modelling, an option is available. A resistor becomes a subcircuit call which creates a number of individual resistive elements which equally divide the main resistor's value and include parasitics devices. The individual elements are defined using an equation define by distrib in the ext.defs file. For example, a resistor is nwell with a width of 10.0U and a length of 20.0U. The normal model is

R1 1 2 RNWELL_W10r0U 2.0

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For a distributed model of 5 elements, the devices becomes

XR1 1 2 3 D_RNWELL_W10r0U SQ=2.0 A=200.0P P=60.0U

.SUBCKT D_RNWELL_W10r0U 1 2 3
XR1 4 5 3 E_RNWELL_W10r0U SQ='SQ/5' A='A/5' P='P/5'
XR2 5 6 3 E_RNWELL_W10r0U SQ='SQ/5' A='A/5' P='P/5'
XR3 6 7 3 E_RNWELL_W10r0U SQ='SQ/5' A='A/5' P='P/5'
XR4 7 2 3 E_RNWELL_W10r0U SQ='SQ/5' A='A/5' P='P/5'
.ENDS

.SUBCKT D_RNWELL_W10r0U 1 2 3
R1 1 2 RNWELL_W10r0U SCALE=equation
D1 1 3 DNWELL AREA='A/2' PJ='P/2'
D2 2 3 DNWELL AREA='A/2' PJ='P/2'
.ENDS

SEARCH PATHS
Ext2spice looks for three different default files like magic. The first file is -/cad/lib/ext.defs, and it is the system default file that must exist. This file defines the different devices available and setups parameters for model values. Users can add or redefine all settings and include additional information using one or both of two custom files (-/ext.defs and ./ext.defs). The local ext.defs has the highest priority in terms of value defining.

The .ext search path uses the same search procedure as magic. The -/cad/lib/magic/sys.magic, -/.magic, and ./magic files are read in sequences and the path is determined via the path and addpath statements.

EXTRACTION DEFAULT FILE COMMANDS
The following are commands recognized in the ext.defs files.

SET
The set command (set nodename number) causes the node with the label nodename to have the node-number "number" in the spice file. For example:

set GND! 0
set input 7

would set the node with the label 'GND!' to 0, and the node with the label 'input' to 7.

DEF
The def command

def extname spicename spicechar flavor

specifies:

a extraction name (extname),
a spice model name (spicename),
a spice device designator (spicechar), and
a flavor for a fet (flavor).
For example:

def pfet PMOS M p
def nfet NMOS M n

would cause the transistors identified as 'pfet' in the .ext file to have model name 'PMOS' in the .spice file and a device definition of M"name". Likewise, 'nfet' transistors get the 'NMOS' model name. The flavor is an internal switch to indicate how a device is to be output and what parameters are needed.

**DISTRIB**
The distrib command is modelling statements in the form

distrib extname scale deviation method parfet equation

specifies:

- the extraction fet type (extname),
- a device value (scale),
- a deviation parameter (deviation),
- the method of calculating a device's value (method),
- the associated parasitic device (parfet), and
- an equation is for distributed modelling.

The parfet and equation are optional for devices without parasitics. For each type of device, the parameters are interpreted differently.

**CAPACITOR**
The scale becomes the number of FF/SQ.

**RESISTOR**
For method=0, the device's ohms/SQ is scale.
For method=1, the device's ohms/SQ is scale * exp(deviation/W) where W is the resistor's width. The model name will include the width. At present, the distrib statements are used to define just capacitor and resistor model statements. The equation defines how the unit element responds as a function of its voltages, currents, or passed parameters.

**PARAM**
The param command (param name value) specifies defaults for generating voltage sources to fix erroneous nodes in subcircuits. For example:

```
param VN WELL 5V
```

causes the a voltage source to be generated for the NWerror! node in a subcircuit.

**ECHO**
The echo command allows statements to be directly written to the .cir file.

```
echo statement
```

All statements should be in the form *message or SPICE will try to interpret them.

**RESISTANCE CLASS COMMANDS**
nrclass, prclass, plrclass, nwclass, and pwclass tell ext2spice which resistance classes (as specified by the magic technology file) correspond to layers which contain information for device. They are called re-

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istance classes because magic thinks it is extracting areas and perimeters to calculate resistance. In-
stead, ext2spice uses it as part of the spice transistor and capacitor description.

plrclass is for the bottom plate for the poly-poly capacitors, nrclass is nmos drain and source ar-
eas/perimeters, and prclass is pmos device. nwclass is for nwell diodes areas/perimeters and pwclass is
for pwell diodes. These commands have the form:

nrclass k
prclass k
nwclass k
pwclass k
plrclass k

where k is an integer specifying the first or second or third resistance class. The defaults are prclass 2;
nrclass 3;
pwclass 7; nwclass 8; plrclass 9.

SPECIAL NAMES
In cmos technology file, there are special nodes, defined in the magic technology file, which represent the
bulk (substrate) or well connections for a device. PWerror!, NWerror!, and Vsub! are commonly used
names. When this nodes are found, ext2spice will generate a global voltage source at the end of the
.spice file to correct the problem. The best scenario is to always define the wells for transistors.

RESISTOR EXTRACTION
Because resistors are extracted like FETs, the size of the resistor's terminals affect the value extracted.
The material for a resistor's terminals must be the same width as the resistor material or the FETs value
will not be extracted correctly. The magic file called resistors contains the correct layout of the various
types of resistors. However, a resistor's width is always printed on the following line as a comment.

WARNINGS
Since spice will be modeling the gate capacitance and the source and drain area and perimeter capaci-
tances, it would be redundant to include them as lumped capacitances . Therefore, it is best to set the
ndc, ndiff, ntrans, pdc, pdiff, and ptrans area and perimeter capacitances to 0 in the magic technology file.

SEE ALSO
magic(1) spice(1) ext(5)

AUTHOR
Core ext2spice originally done by Andy Burstein. Additions and modifications by Danny F. Newport for
MOSIS npn transistors, poly and well resistors, and poly-poly capacitors extractions and other capabilities.
Other modifications by Lloyd Clonts for enhancing performance, removing unused nodes, generating
automatic .model statements for resistors and capacitors.

BUGS
Diffusion areas that overlap between cells might not be measured correctly.
Appendix E - Ext.defs support file for Ext2spice (by Lloyd Clonts)

Ext2spice defaults file -cad/lib/ext.defs

******************************************************************************

Modified 12/03/96 by Lloyd Clonts

Grouped the different devices together and started to document the changes. It was out of control.

******************************************************************************

First letter of FET name must be in the following format:

p = PMOS device
n = NMOS device
r = Resistor
c = Capacitor
l = inductor

With this format, the ext2sim can be modified to have only one extraction format for all devices. The following actions will be taken for the given devices:

p,n = No changes
r,l = Short or a resistor
c = Change to a capacitor in the output

LGC - 12/16/96

******************************************************************************

Class definitions

prclass 1
nrclass 2
nwclass 3
pwclass 4
plrclass 5
Base structures - Present most CMOS processes

```
def nfet CMOSN M n
def nwell DNWELL D n
def ndiff DNDIFF D n
def pnet CMOSP M p
def pwell DPWELL D p
def pdiff DNDIFF D p
```

Bipolar Types

```
def qnpn NPN Q q
```

Float capacitor

```
def enet EFETN M n
def epet EFETP M p
```

Triquent GaAs Structures

```
def ngaas M1SSac B b
def dgaas DGAAS B b
def egaas EGAAS B b
def gaasdiode GAASDIODE D d
def resdiff RDEPL_M R r
def resdiffd RDEPL_D R r
def resdiffc RDEPL_E R r
def ngaasres NGAASRES R r
def nicrom NICROM R r
def lgaas LGAAS L 1
distrib resdiff 10.0 0.0 0
distrib resdiffd 625.0 0.0 0
distrib resdiffc 1000.0 0.0 0
distrib ngaasres 125.0 0.0 0
distrib nicrom 50.0 0.0 0
```

Generic Definitions - Older extraction format

```
def ccap CAP C c
def ccapp CAPP C c
```
def pcap PSCAP C c p
def pcap PSCAPP C c p
def ncap NSCAP C c n
def ncap NSCAP C c n
def hcap HPCAP C c
def hcap HPCAP C c
distrib pcap 0.475 0.0 0 pcap
distrib pcap 0.475 0.0 0 pcap
distrib pcap 0.930 0.0 0
distrib pcap 0.930 0.0 0
def ndres RNDIFF R r
def pdres RPDIFF R r
def polyres RPOLY R r
def nwres RNWELL R r
def pwres RPWELL R r
distrib polyres 90.0 0.000 0 polyres 90.0 0.000 0

def ndres200u RNDIFF R r
def pdres200u RPDIFF R r
def polyres200u RPOLY R r
def nwres200u RNWELL R r
def pwres200u RPWELL R r
distrib polyres200u 22.5 0.000 0 polyres200u 22.5 0.000 0

<table>
<thead>
<tr>
<th>12.0 ORBIT micron definitions</th>
</tr>
</thead>
</table>

def pcap200u PSCAP200u C c p
def pcap200u PSCAP200u C c p
def ncap200u NSCAP200u C c n
def ncap200u NSCAP200u C c n
def pcap200u PSCAP200u C c p
def ncap200u NSCAP200u C c n
def pcap200u PSCAP200u C c p
def ncap200u NSCAP200u C c n
distrib pcap200u 0.475 0.0 0 pcap200u
1.2 ORBIT micron

The NWELL measurements were done by Dr. Kennedy 1994 for a ORBIT nwell process.

1.2 HP micron
def polyreshp120u  RHPPOLY120  R r
distrib polyreshp120u  2.6  0.000 0  chppoly120u  SQ

def chppoly120u  CHPPOLY120  C c
distrib chppoly120u  0.57  0.000 0

def nwreshp120u  RHNWELL120  R r
distrib nwreshp120u  821.0  0.000 0  dnwell  SQ

def hcap120u  HPCAP1r2  C c
distrib hcap120u  1.260  0.0 0  hcapp120u

# The parasitic capacitance is between the cwell and substrate.
#
# def hcap120u  HPCAP1r2P  C c
distrib hcap120u  0.001  0.0 0

|***************************************************************************************************|
|***************************************************************************************************|
| Generic Definitions - New Format|
|***************************************************************************************************|
|***************************************************************************************************|

def cap  CAP  C c
def capp  CAPP  C c
def cpfet  PSCAP  C c p
def cpfetp  PSCAPP  C c p
def cnfet  NSCAP  C c n
def cnfetp  NSCAPP  C c n
def cwhp  HPCAP  C c
def cwhpp  HPCAPP  C c
distrib cap  0.475  0.0 0  capp
distrib capp  0.088  0.0 0
distrib cpfet  0.475  0.0 0  cpfetp
distrib cpfetp  0.930  0.0 0
distrib cnfet  0.475  0.0 0  cnfetp
distrib cnfetp  0.930  0.0 0
def rndiff  RNDIFF  R r
def rpdiff  RPDIFF  R r
def rpoly  RPOLY  R r
def rnwell  RNWELL  R r
def rpwell  RPWELL  R r
distrib rpoly  90  0.000 0  capp  SQ
distrib rndiff  25.0  0.000 0  dndiff  SQ
distrib rpdiff  77.5  0.000 0  dpdiff  SQ
distrib rnwell  2392.0  1.289 1  dnwell  SQ*(1+0.02*(v(1,3)+v(2,3))+0.02*ABS(v(1,2)))
distrib rpwell  1921.0  1.347 1  dpwell  SQ*(1+0.02*(v(1,3)+v(2,3))+0.02*ABS(v(1,2)))

|***************************************************************************************************|
|***************************************************************************************************|
| Generic Definitions - New format|
|***************************************************************************************************|
|***************************************************************************************************|

58
def cap   CAP   C c
def capp  CAPP  C c
def cpfet  PSCAP  C c p
def cpfetp PSCAPP  C c p
def cnfet  NSCAP  C c n
def cnfetp NSCAP  C c n
def hcap   HPCAP  C c
def hcapp  HPCAPP  C c
distrib cap  0.475  0.0  0 capp
distrib capp  0.057  0.0  0
distrib cpfet  0.475  0.0  0 cpfetp
distrib cpfetp  0.930  0.0  0
distrib cnfet  0.475  0.0  0 cnfetp
distrib cnfetp  0.930  0.0  0
def rndiff RNDIFF R r
def rpdiff RPDIFF R r
def rpoly RPOLY R r
def rnwell RNWELL R r
def rpwell RPWELL R r
distrib rpoly  22.5  0.000  0 capp  SQ
distrib rndiff  25.0  0.000  0 dndiff  SQ
distrib rpdiff  77.5  0.000  0 dpdiff  SQ
distrib rnwell  2392.0  1.289  1 dnwell  SQ*(1+0.02*(v(1,3)+v(2,3))+0.02*ABS(v(1,2)))
distrib rpwell  1921.0  1.347  1 dpwell  SQ*(1+0.02*(v(1,3)+v(2,3))+0.02*ABS(v(1,2)))

2.0 ORBIT micron definitions - New Format

def rndiff200u RNDIFF R r
def rpdiff200u RPDIFF R r
def rpoly200u RPOLY R r
def rnwell200u RNWELL R r
def rpwell200u RPWELL R r
distrib rpoly200u  22.5  0.000  0 capp200u  SQ
distrib rndiff200u  25.0  0.000  0 dndiff200u  SQ
distrib rpdiff200u  77.5  0.000  0 dpdiff200u  SQ
distrib rnwell200u  2392.0  1.289  1 dnwell200u  SQ*(1+0.02*(v(1,3)+v(2,3))+0.02*ABS(v(1,2)))
distrib rpwell200u  1921.0  1.347  1 dpwell200u  SQ*(1+0.02*(v(1,3)+v(2,3))+0.02*ABS(v(1,2)))
def cap200u CAP2r0  C c
def capp200u CAP2r0P  C c
def cpfet200u PSCAP2r0  C c p
def cpfetp200u PSCAP2r0P  C c p
def cnfet200u NSCAP2r0  C c n
def cnfetp200u NSCAP2r0P  C c n
def pfcap200u PFCAP2r0  C c p
def nfcap200u NFCAP2r0 C c n
distrib cap200u 0.475 0.0 0.0 capp200u
distrib capp200u 0.057 0.0 0
distrib cpfet200u 0.475 0.0 0.0 cpfetp200u
distrib cpfetp200u 0.930 0.0 0
distrib cnfet200u 0.475 0.0 0.0 cnfetp200u
distrib cnfetp200u 0.930 0.0 0
distrib nfcap200u 0.475 0.0 0.0

distrib pfcap200u 0.475 0.0 0.0

1.2 ORBIT micron - New Format

The NWELL measurements were done by Dr. Kennedy 1994 for a ORBIT nwell process.

1.2 HP micron - New Format

|****************************|
|****************************|
|****************************|
|****************************|

def cap120u CAP1r2 C c
def capp120u CAP1r2P C c
def cpfet120u CAP1r2 C c p
def cpfetp120u CAP1r2P C c p
def cnfet120u CAP1r2 C c n
def cnfetp120u CAP1r2P C c n
# def cpfet120u PSCAP1r2 C c
# def cpfetp120u PSCAP1r2P C c
# def cnfet120u NSCAP1r2 C c
# def cnfetp120u NSCAP1r2P C c

distrib cap120u 0.0 0.0 0.500 capp120u
distrib capp120u 0.0 0.0 0.0
distrib cpfet120u 0.0 0.0 0.0
distrib cpfetp120u 0.0 0.0 0.0
distrib cnfet120u 0.0 0.0 0.0
distrib cnfetp120u 0.0 0.0 0.0
def rndiff120u RNDIFF R r
def rpdiff120u RPDIFF R r
def rpoly120u RPOLY R r
def rnwell120u RNWELL R r
def rpwell120u RPWELL R r
distrib rpoly120u 22.4 0.000 0.000 capp120u
distrib rndiff120u 39.5 0.000 0.000 cdiff

distrib rpdiff120u 77.5 0.000 0.000 ddiff
distrib rnwell120u 953.0 7.160 1.000 dnwell

distrib rpwell120u 1080.6 6.968 1.000 dpwell

|****************************|
|****************************|
|****************************|
|****************************|
|****************************|

1.2 HP micron - New Format

|****************************|
|****************************|

60
```python
def cwellhp120u HPCAP120 Cc
def cwellhp120up HPCAP120P Cc
def cpolyhp120up CHPPOLY120 Cc

def rndiffhp120u RHPNDIFF120 R r
def rpdifffhp120u RHPPDIFF120 R r
def rpolyhp120u RHPPOLY120 R r

def rndiffhp120u RHPNWELL120 R r

distrib cwellhp120u 1.260 0.0 0

distrib cwellhp120up 0.57 0.0 0

distrib cpolyhp120up 2.4 0.000 0 dndiff SQ

distrib cpolyhp120up 2.1 0.000 0 cpolyhp120up SQ

distrib cpolyhp120up 2.6 0.000 0 dpdiff SQ

distrib cpolyhp120up 821.0 0.000 0 dnwell SQ

0.8 HP micron - New format
```

```python

def cwellhp80u HPCAP80 Cc

def cwellhp80up HPCAP80P Cc

def cpolyhp80up CHPPOLY80 Cc

def rndiffhp80u RHPNDIFF80 R r

def rpdifffhp80u RHPPDIFF80 R r

def rpolyhp80u RHPPOLY80 R r

def rpolysbhp80u RHPPOLYSBSO R r

def rnwellhp80u RHPNWELL80 R r

distrib cwellhp80u 1.260 0.0 0

distrib cwellhp80up 0.57 0.0 0

distrib cpolyhp80up 2.4 0.000 0 dndiff SQ

distrib cpolyhp80up 2.1 0.000 0 cpolyhp80up SQ

distrib cpolyhp80up 2.6 0.000 0 dpdiff SQ

distrib cpolyhp80up 821.0 0.000 0 dnwell SQ

1.0 HP micron - format
```

```python

def cwellhp50u HPCAP50 Cc

def cwellhp50up HPCAP50P Cc

def cpolyhp50up CHPPOLY50 Cc

def rndiffhp50u RHPNDIFF50 R r

def rpdifffhp50u RHPPDIFF50 R r

def rpolyhp50u RHPPOLY50 R r

def rpolysbhp50u RHPPOLYSBSO R r

def rnwellhp50u RHPNWELL50 R r

```
distrib cwellhp50u 3.56 0.0 0
distrib cpolyhp50up 0.088 0.0 0
distrib rndiffhp50u 2.4 0.000 0 dndiff SQ
distrib rpolyhp50u 2.1 0.000 0 cpolyhp50up SQ
distrib rpolysbhp50u 90.0 0.000 0 cpolyhp50up SQ
distrib rpdiffhp50u 2.6 0.000 0 dpdiff SQ
distrib rnwellhp50u 821.0 0.000 0 dnwell SQ

0.35 HP micron - format

---

def cwellhp35u HPCAP35 C c
def cwellhp35up HPCAP35P C c
def cpolyhp35up CHPPOLY35 C c
def rndiffhp35u RHPNDIFF35 R r
def rpdiffhp35u RHPPDIFF35 R r
def rpolyhp35u RHPPOLY35 R r
def rpolysbhp35u RHPPOLYSB35 R r
def rnwellhp35u RHPNWELL35 R r
distrib cwellhp35u 3.56 0.0 0
distrib cpolyhp35up 0.57 0.0 0
distrib rndiffhp35u 2.4 0.000 0 dndiff SQ
distrib rpolyhp35u 2.1 0.000 0 cpolyhp35up SQ
distrib rpolysbhp35u 90.0 0.000 0 cpolyhp35up SQ
distrib rpdiffhp35u 2.6 0.000 0 dpdiff SQ
distrib rnwellhp35u 821.0 0.000 0 dnwell SQ

---

0.35 IMEMS - format

---

def cbaseimems CBASE C c
def cemitimems CEMIT C c
def rndiffimems RNDIFFIMEMS R r
def rpdiffimems RPDIFFIMEMS R r
def rtflimems RTFLIMEMS R r
def rnwellimems RNWELLIMEMS R r
def lpnp LPNP Q q
def spnp SPNP Q q
def vnnp VNPN Q q
distrib rndiffimems 2.4 0 0
distrib rpdiffimems 2.4 0 0
distrib rtflimems 2.4 0 0
distrib rnwellimems 500 0 0
param VNWELL 5V
param NWELL DNWELL
param PWELL DPWELL
param VPWELL 0V
param VCWELL 0V

I Info into .cir files

include /usr/local/cad/lib/spice/level3/hp50_n56s.l3
include /usr/local/cad/lib/spice/level3/hp50_n61y.l3
include /usr/local/cad/lib/spice/level3/hp50_n62e.l3
include /usr/local/cad/lib/spice/level3/hp50_n64o.l3
include /usr/local/cad/lib/spice/level3/hp50_n65a.l3
include /usr/local/cad/lib/spice/level3/hp50_n6ay.l3
include /usr/local/cad/lib/spice/level3/hp50_n6bg.l3
include /usr/local/cad/lib/spice/level3/hp50_n71s.l3
include /usr/local/cad/lib/spice/level3/hp50_n72a.l3
include /usr/local/cad/lib/spice/level3/hp50_n73d.l3
include /usr/local/cad/lib/spice/level13/hp50_n56s.l13
include /usr/local/cad/lib/spice/level13/hp50_n61y.l13
include /usr/local/cad/lib/spice/level13/hp50_n62e.l13
include /usr/local/cad/lib/spice/level13/hp50_n64o.l13
include /usr/local/cad/lib/spice/level13/hp50_n65a.l13
include /usr/local/cad/lib/spice/level13/hp50_n6ay.l13
include /usr/local/cad/lib/spice/level13/hp50_n6bg.l13
include /usr/local/cad/lib/spice/level13/hp50_n71s.l13
include /usr/local/cad/lib/spice/level13/hp50_n72a.l13
include /usr/local/cad/lib/spice/level13/hp50_n73d.l13
echo *.include /usr/local/cad/lib/spice/level13/hp50_n6bg.l13
echo *.include /usr/local/cad/lib/spice/level13/hp50_n71s.l13
echo *.include /usr/local/cad/lib/spice/level13/hp50_n72a.l13
echo *.include /usr/local/cad/lib/spice/level13/hp50_n73d.l13
echo *

echo *********************************************

echo * ORBIT 1.2U SPICE models
echo *********************************************

echo *

echo *.include /usr/local/cad/lib/spice/level2/n1nom1u2.12

echo *

echo *.include /usr/local/cad/lib/spice/level3/n1fst1u2.13

echo *

echo *.include /usr/local/cad/lib/spice/level3/n1lom1u2.13

echo *

echo *.include /usr/local/cad/lib/spice/level3/n1sl01u2.13

echo *

echo *.include /usr/local/cad/lib/spice/level13/xxxxx1u2.113

echo *

echo *********************************************

echo * MOSIS/ORBIT 2.0U DIODE model parameters
echo *********************************************

echo *

echo *.include /usr/local/cad/lib/spice/level1/diodc2u0.11

echo *

echo *********************************************

echo * ORBIT 2.0U SPICE models
echo *********************************************

echo *

echo *.include /usr/local/cad/lib/spice/level13/m961n2u0.113

echo *

echo *.include /usr/local/cad/lib/spice/level13/n07sn2u0.113

echo *

echo *.include /usr/local/cad/lib/spice/level13/n13in2u0.113

echo *

echo *.include /usr/local/cad/lib/spice/level13/n21nh2u0.113

echo *

echo *.include /usr/local/cad/lib/spice/level13/n2cqn2u0.113

echo *

echo *.include /usr/local/cad/lib/spice/level13/n02sn2u0.113

echo *

echo *.include /usr/local/cad/lib/spice/level13/n09enn2u0.113

echo *

echo *.include /usr/local/cad/lib/spice/level13/n15n2u0.113

echo *

echo *.include /usr/local/cad/lib/spice/level13/n23q2u0.113

echo *

echo *.include /usr/local/cad/lib/spice/level13/n32an2u0.113

echo *

echo *.include /usr/local/cad/lib/spice/level13/n04zn2u0.113

echo *

echo *.include /usr/local/cad/lib/spice/level13/n0bpm2u0.113

echo *

echo *.include /usr/local/cad/lib/spice/level13/n18mm2u0.113

echo *

echo *.include /usr/local/cad/lib/spice/level13/n29tm2u0.113

echo *

echo *.include /usr/local/cad/lib/spice/level13/n05in2u0.113

echo *

echo *.include /usr/local/cad/lib/spice/level13/n11bn2u0.113

echo *

echo *.include /usr/local/cad/lib/spice/level13/n1avn2u0.113

echo *

echo *.include /usr/local/cad/lib/spice/level13/n2ben2u0.113

echo *

echo *********************************************

echo * HSPICE options
echo *********************************************

echo *

echo *.options POST

echo *.options METHOD=GEAR

echo *

echo *********************************************

echo *
echo * Used for distributed modelling or when well
diodes are generated.

Used for distributed modelling or when well

---

echo *Vsub Vsub 0 DC 0V $NWell Process
echo *Vsub Vsub 0 DC 5V $PWell Process
echo *
Appendix F - Ext2spice.corr code

# Replaces all "=" with " = " in order to get values.
sed '/=// s/=/ = /g' $1!

# Replaces all ",_L" with " ," in order to get the value of NL.
sed '/,L// s/,/= /' !

# Eliminates capacitors with value 0.0F.
sed '/0.0F/d' !

nawk 'BEGIN {LAMBDA=0.3e-6}
{RSQ=90}
{RSQPOLY=3.5}
{RDW=0.066}
{RACAP=8.5e-5}
{RPCAP=1.22e-10}
{WMIN=1.5e-6}
{RSQM1=0.07}
{RSQM2=0.07}
{RSQM3=0.05}
{VC1=0.001}
{VC2=0.0005}
{TC1=0}
{TC2=0}
{NRSQ=2.5}
{PRSQ=2}
{NDC=4}
{PDC=4}
{PC=7}
{CLT=5}
{CMOSN="CMOSN"}
{CMOSP="CMOSP"}

# Sets NL as parameter and calculates R, W, Weff, Cpar
if ($5=="D_RPOLY") print $1,$2,$3,$4,$5,$6,$7,$8,$9,$10,$11,$12,"(NL="0""); print
"*",$1="2*PC+2*RSQPOLY*2*LAMBDA/sqrt($12*le-12/$9)+(RSQ*(9-0.45)/(1-
RDW/sqrt($12* 1 e- 12/$9))),"(W="sqrt($12* 1 e- 12/$9)", Weff="(sqrt($12* 1e-12/$9)-RDW)",
Cpar="(RACAP*$12* 1 e-l2+RPCAP*2*sqrt($12* le-12*$9))")"
else if ($6=="D_RPOLY") print $1,$3,$4,$5,$6,$7,$8,$9,$10,$11,$12,$13,"(NL="0""); print
"*",$1="2*PC+2*RSQPOLY*2*LAMBDA/sqrt($13*1e-12/$10)+(RSQ*(9-0.45)/(1-
RDW/sqrt($13*1e-12/$10))),"(W="sqrt($13*1e-12/$10)", Weff="(sqrt($13*1e-12/$10)-RDW)",
Cpar="(RACAP*$13*1e-12+RPCAP*2*sqrt($13*1e-12*$10))")"
else if ($2=="D_RPOLY") print $1,$2,$3,$4,$5,$6,$7,$8,$9,$10,$11,$12,"(NL="0""); print
"*",$1="2*PC+2*RSQPOLY*2*LAMBDA/sqrt($13*1e-12/$10)+(RSQ*(9-0.45)/(1-
RDW/sqrt($13*1e-12/$10))),"(W="sqrt($13*1e-12/$10)", Weff="(sqrt($13*1e-12/$10)-RDW)",
Cpar="(RACAP*$13*1e-12+RPCAP*2*sqrt($13*1e-12*$10))")"
else if ($2=="E_RPOLY") print $1,$2,$3,$4,$5,$6,$7,$8,$9,$10,$11,$12,"(NL="0""); print
"*",$1="2*PC+2*RSQPOLY*2*LAMBDA/sqrt($13*1e-12/$10)+(RSQ*(9-0.45)/(1-
RDW/sqrt($13*1e-12/$10))),"(W="sqrt($13*1e-12/$10)", Weff="(sqrt($13*1e-12/$10)-RDW)",
Cpar="(RACAP*$13*1e-12+RPCAP*2*sqrt($13*1e-12*$10))")"
else if ($5=="E_Rpoly") print $1,$2,$3,$4,$5,$6,$7,$8,$9,$10,$11,$12,"(NL="0""); print
"*",$1="2*PC+2*RSQPOLY*2*LAMBDA/sqrt($13*1e-12/$10)+(RSQ*(9-0.45)/(1-
RDW/sqrt($13*1e-12/$10))),"(W="sqrt($13*1e-12/$10)", Weff="(sqrt($13*1e-12/$10)-RDW)",
Cpar="(RACAP*$13*1e-12+RPCAP*2*sqrt($13*1e-12*$10))")"
else if ($4=="RPOLY" && $5=="[SQ]") print
$1,$2,$3,("2*PC+2*RSQPOLY*2*LAMBDA/VSQRT(A*1p/VSQ)+RSQ*(SQ-NL*0.45)/(1-
RDW/SQRT(A*1p/VSQ))")

66
else if ($4=="CAPP" && $5=="{A/2}\) print $1,$2,$3,("{RACAP*A*1pV2+RPCAP*2*SQRT(A*1p*SQ)}")
else
if ($1==".MODEL" && $2=="RPOLY") print ".PARAM RSQ="RSQ","RDW="RDW","SQ=1 A=1p NL=0"
else
if ($1==".MODEL" && $2=="CAPP") print ".PARAM RACAP="RACAP","RPCAP="RPCAP"
else

# Modifies NRS, NRD and NRG and adds .PARAM line for GRSQ
if ($1="/M/ && $6==CMOSN && $7=="W")
{W=$9*1e-6
L=$12*1e-6
print $0
getline
(if ($1="/+" && $2=="NRG") print
$1,"RG","3,(RSQPOLY*(1/$4+4*M*4*LAMBDAL)L)/4+PC/2/(L/CLT/LAMBDAL),"RD","6,(NRSQ*(
$7-3*LAMBDAL/W)+NDC/(W/CLT/LAMBDAL),"RS","9,(NRSQ*(10-
3*LAMBDAL/W)+PDC/(W/CLT/LAMBDAL))
else print $0})
else
if ($1="/M/ && $6==CMOSN && $7=="M")
{M=$9
W=$12*1e-6
L=$15*1e-6
print $0
getline
(if ($1="/+" && $2=="NRG") print
$1,"RG","3,(RSQPOLY*(1/$4+4*M*4*LAMBDAL)L)/4+PC/2/(L/CLT/LAMBDAL),"RD","6,(NRSQ*(
$7-3*LAMBDAL/W)+NDC/(W/CLT/LAMBDAL),"RS","9,(NRSQ*(10-
3*LAMBDAL/W)+PDC/(W/CLT/LAMBDAL))
else print $0})
else
if ($1="/M/ && $6==CMOSN && $7=="W")
{W=$9*1e-6
L=$12*1e-6
print $0
getline
(if ($1="/+" && $2=="NRG") print
$1,"RG","3,(RSQPOLY*(1/$4+4*M*4*LAMBDAL)L)/4+PC/2/(L/CLT/LAMBDAL),"RD","6,(PRSQ*(
$7-3*LAMBDAL/W)+NDC/(W/CLT/LAMBDAL),"RS","9,(PRSQ*(10-
3*LAMBDAL/W)+PDC/(W/CLT/LAMBDAL))
else print $0})
else
if ($1="/M/ && $6==CMOSN && $7=="M")
{M=$9
W=$12*1e-6
L=$15*1e-6
print $0
getline
(if ($1="/+" && $2=="NRG") print
$1,"RG","3,(RSQPOLY*(1/$4+4*M*4*LAMBDAL)L)/4+PC/2/(L/CLT/LAMBDAL),"RD","6,(PRSQ*(
$7-3*LAMBDAL/W)+NDC/(W/CLT/LAMBDAL),"RS","9,(PRSQ*(10-
3*LAMBDAL/W)+PDC/(W/CLT/LAMBDAL))
else print $0})
else print $0 }
else
  if ($1=="*" && $2=="Model")
    print ".PARAM RSQPOLY="RSQPOLY,"PC="PC
  else

# Sets model for linear capacitors.
  if ($1=="MODEL" && $2=="LCAP") print $1,$2,"CAP (C=l
  VC1="VC1,"VC2="VC2,"TC1="TC1,"TC2="TC2,"
else

# Sets expression for R in the non distributed case.
  if ($5=="RPOLY") print $1,$3,$4,2*PC+2*RSQPOLY*2*LAMBDA/WMIN+(RSQ*($6-$2*0.45)/(1-
  RDW/WMIN))
else

# Evaluates the resistance of Metal1 fets
  if ($6=="RM1")
    print ".R$1,$2,$4,(RSQM1*(($15+$21)/$9/$9+1))
    getline
    if ($1=="+")
      print *
    else
      print $0
  else

# Evaluates the resistance of Metal2 fets
  if ($6=="RM2")
    print ".R$1,$2,$4,(RSQM2*(($15+$21)/$9/$9+1))
    getline
    if ($1=="+")
      print *
    else
      print $0
  else

# Evaluates the resistance of Metal3 fets
  if ($6=="RM3")
    print ".R$1,$2,$4,(RSQM3*(($15+$21)/$9/$9+1))
    getline
    if ($1=="+")
      print *
    else
      print $0
else

# Separates linear capacitors LCAP
  nawk '{ BEGIN { K=0 }
    { k=0 }
    { flag=0 }
    { if ($4=="LCAP")
      { K=K+1 }
      N1[K]=$2
      N2[K]=$3
      getline }
    else
      { while (flag<=0)
        { if (($2==N1[k] && $3==N2[k] && $1-"/C") || ($2==N2[k] && $3==N1[k] && $1-"/C")}
          print $1,$2,$3,"LCAP",$4
          flag=1 }
        else
          { if (k==K)
            print $0
            flag=1 }
          } }'}
else
  {k=k+1}}}

# Replaces all " = " with "=".
sed '/' = /s/=/g' l

# Delete extra * R line generated by extspice.
sed 'A* R.*$d'
Appendix G - Technology File: Summary of Replacements and Additions

Planes section:

# ADDED
substrate, ss

Types section:

# ADDED
metal 1 m1res, m1r
metal 2 m2res, m2r
metal 3 m3res, m3r

Styles section:

# ADDED
mlres 20
mlres 12
mlres 13
m2res 21
m2res 12
m2res 13
m3res 22
m3res 12
m3res 13

Connect section:

# MODIFIED
m1, m1res, ndc/m1, nsc/m1, cwc/m1, pdc/m1, psc/m1, cwc/m1, pc/m1, m2c/m1
ndc/m1, nsc/m1, cwc/m1, pdc/m1, psc/m1, cwc/m1, pc/m1, m2c/m1
m2, m2res, m2c/m2, m3c/m2, m3c/m2
m2c/m2, m3c/m2, m3c/m2
m3, m3res, m3c/m3, m3c/m3
poly, wcap, nfet, wcap, pfet, pc/a poly, wcap, nfet, wcap, pfet, pc/a

Cifoutput section:

# MODIFIED
layer CMF m1, m1res, ndc/m1, nsc/m1, cwc/m1, pdc/m1, psc/m1, cwc/m1, pc/m1, m2c/m1
labels m1, m1, ndc/m1, nsc/m1, cwc/m1, pdc/m1, psc/m1, cwc/m1, pc/m1, m2c/m1
layer CMS m2, m2res, m2c/m2, m3c/m2, m3c/m2
labels m2, m2, m2c/m2, m3c/m2, m3c/m2
layer CMT m3, m3res, m3c/m3
labels m3, m3, m3c/m3

All Extract section:

extract
style HP 0.5um(hpcmos14b)from:N81D
cscale 1
lambda 30
step 100
sidehalo 8
# MODIFIED ORDER
planeorder ss 0
planeorder well 1
planeorder implant 2
planeorder select 3
planeorder active 4
planeorder metal1 5
planeorder metal2 6
planeorder metal3 7
planeorder oxide 8
planeorder xp 9
planeorder comment 10
planeorder via1 12
planeorder via2 13

# REPLACED resist (ndiff,anres,ndc/a,nsd,nsc/a)/active 1900
resist ((ndiff,ndc/a)/active,(pdiff,pdc/a)/active) 0
# REPLACED resist (pdiff,apres,pdc/a,psd,psc/a)/active 2000
resist (ml/metal1,m2/metal2,m3/metal3) 0
resist (poly,pres,wcap,pc/a,wcap,pc/a,nfet,pfet)/active 1800
# DELETED resist (m1,ndc/ml,nsc/ml,cwc/ml,pdc/m1,psc/ml,cwc/ml,pc/ml,m2c/ml,m2c/ml)/metal1 70
# DELETED resist (m2,m2c/m2,m3c/m2,m3c/m2)/metal2 70
# DELETED resist (m3,m3c/m3,pad)/metal3 50

contact ndc 4 2000
contact pdc 4 2000
contact pc 4 1700
contact m2c 4 700
contact m3c 4 360

#wellcap
overlap wcap capwell 207.900

#nwell,cwell,pwell
# REPLACED areacap (nwell,cwell)/well 8.460
# REPLACED areacap (cwell)/well 8.460
overlap (nwell,cwell)/well ss/ss 8.460

#ndiff
areacap (anres)/active 48.330
overlap (anres)/active -space/w 48.330
perimc (anres)/active -(anres)/active 67.800
sideoverlap (anres)/active -(anres)/active -space/w 67.800
# MODEL HANDLES THIS: areacap (ndiff,anres,ndc/a)/active 48.330
# MODEL HANDLES THIS: overlap (ndiff,anres,ndc/a)/active -space/w 48.330
# MODEL HANDLES THIS: perimc (ndiff,anres,ndc/a)/active -(ndiff,anres,ndc/a,nfet,pfet)/active 67.800
# MODEL HANDLES THIS: sideoverlap (ndiff,anres,ndc/a)/active -(ndiff,anres,ndc/a,nfet,pfet)/active -space/w 67.800

#pdiff
areacap (apres)/active 84.600
overlap (apres)/active -space/w 84.600
perimc (apres)/active -(apres)/active 67.500
sideoverlap (apres)/active -(apres)/active -space/w 67.500
# MODEL HANDLES THIS: areacap (pdiff,apres,pdc/a)/active 84.600
MODEL HANDLES THIS:
overlap (pdiff,apres,pdc/a)/active −space/w 84.600
MODEL HANDLES THIS: perimc (pdiff,apres,pdc/a)/active −(pdiff,apres,pdc/a,nfet,pfet)/active 67.500
MODEL HANDLES THIS: sideoverlap (pdiff,apres,pdc/a)/active −(pdiff,apres,pdc/a,nfet,pfet)/active −space/w 67.500

# polycap

# poly sidewall (poly,pres,wcap,pc/a,wcap,pc/a)/active −(poly,pres,wcap,pc/a,wcap,pc/a)/active −(poly,pres,wcap,pc/a,wcap,pc/a)/active 2.136
# REPLACED areacap (poly,pres,wcap,pc/a,wcap,pc/a)/active −space/w 7.920
overlap (poly,pres,wcap,pc/a,wcap,pc/a)/active ss/ss 7.920 −space/w
overlap (poly,pres,p/a,pc/a)/active −space/w 7.920
# REPLACED perimc (poly,pres,wcap,pc/a,wcap,pc/a)/active −(poly,pres,wcap,pc/a,wcap,pc/a)/active 15
sideoverlap (poly,pres,wcap,pc/a,wcap,pc/a)/active −(poly,pres,wcap,pc/a,wcap,pc/a)/active ss/ss 15 −space/w
# REPLACED sideoverlap (poly,pres,wcap,pc/a,wcap,pc/a)/active −(poly,pres,wcap,pc/a,wcap,pc/a)/active −space/w 15
sideoverlap (poly,pres,wcap,pc/a,wcap,pc/a)/active −(poly,pres,wcap,pc/a,wcap,pc/a)/active −space/w 15
# ADDED
sideoverlap (wcap)/active (cwnsd)/active −space/w 40
MODEL HANDLES THIS: overlap (nfet)/active (ndiff,anres,ndc/a)/active 324.360
MODEL HANDLES THIS: sideoverlap (nfet)/active −(nfet)/active (ndiff,anres,ndc/a)/active 15
MODEL HANDLES THIS: overlap (pfet)/active (pdiff,apres,pdc/a)/active 308.160
MODEL HANDLES THIS: sideoverlap (pfet)/active −(pfet)/active (pdiff,apres,pdc/a)/active 15

# poly2

# metal1 sidewall (ml,ndc/ml,nsc/ml,cwc/ml,pdc/m1,psc/ml,cwc/ml,pc/ml,m2c/ml)/metal1
−(ml,ndc/ml,nsc/ml,cwc/ml,pdc/m1,psc/ml,cwc/ml,pc/ml,m2c/ml)/metal1
−(ml,ndc/ml,nsc/ml,cwc/ml,pdc/m1,psc/ml,cwc/ml,pc/ml,m2c/ml)/metal1
−(ml,ndc/ml,nsc/ml,cwc/ml,pdc/m1,psc/ml,cwc/ml,pc/ml,m2c/ml)/metal1
−(ml,ndc/ml,nsc/ml,cwc/ml,pdc/m1,psc/ml,cwc/ml,pc/ml,m2c/ml)/metal1 6.690
# REPLACED areacap (ml,ndc/ml,nsc/ml,cwc/ml,pdc/m1,psc/ml,cwc/ml,pc/ml,m2c/ml)/metal1 2.700
overlap (ml,ndc/ml,nsc/ml,cwc/ml,pdc/m1,psc/ml,cwc/ml,pc/ml,m2c/ml)/metal1 ss/ss 2.700 −space/w,−space/a
# metal-sub blocked by −space/a,−space/c
# REPLACED overlap (ml,ndc/ml,nsc/ml,cwc/ml,pdc/m1,psc/ml,cwc/ml,pc/ml,m2c/ml)/metal1 −space/w 2.700 −space/a
overlap (ml,ndc/ml,nsc/ml,cwc/ml,pdc/m1,psc/ml,cwc/ml,pc/ml,m2c/ml)/metal1 −space/w 2.700 −space/a
# REPLACED perimc (ml,ndc/ml,nsc/ml,cwc/ml,pdc/m1,psc/ml,cwc/ml,pc/ml,m2c/ml)/metal1
−(ml,ndc/ml,nsc/ml,cwc/ml,pdc/m1,psc/ml,cwc/ml,pc/ml,m2c/ml)/metal1 9.300
sideoverlap (ml,ndc/ml,nsc/ml,cwc/ml,pdc/m1,psc/ml,cwc/ml,pc/ml,m2c/ml)/metal1
−(ml,ndc/ml,nsc/ml,cwc/ml,pdc/m1,psc/ml,cwc/ml,pc/ml,m2c/ml)/metal1 ss/ss 9.300 −space/w,−space/a
# REPLACED sideoverlap (ml,ndc/ml,nsc/ml,cwc/ml,pdc/m1,psc/ml,cwc/ml,pc/ml,m2c/ml)/metal1
−(ml,ndc/ml,nsc/ml,cwc/ml,pdc/m1,psc/ml,cwc/ml,pc/ml,m2c/ml)/metal1 −space/w 9.300 −space/a
sideoverlap (ml,ndc/ml,nsc/ml,cwc/ml,pdc/m1,psc/ml,cwc/ml,pc/ml,m2c/ml)/metal1
−(ml,ndc/ml,nsc/ml,cwc/ml,pdc/m1,psc/ml,cwc/ml,pc/ml,m2c/ml)/metal1 −space/w 9.300 −space/a
# metal-diff blocked by −space/c
# REPLACED overlap (ml,ndc/ml,nsc/ml,cwc/ml,pdc/m1,psc/ml,cwc/ml,pc/ml,m2c/ml)/metal1
(ndiff,anres,ndc/a)/active 2.790
overlap (ml,ndc/ml,nsc/ml,cwc/ml,pdc/m1,psc/ml,cwc/ml,pc/ml,m2c/ml)/metal1 (ndiff,anres,ndc/a)/active 2.790
# REPLACED sideoverlap (m1,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1
- (m1,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1
  (ndiff,anres,ndc/a)/active 9.300
sideoverlap (m1,nsc/m1,cwc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1
- (m1,nsc/m1,cwc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1 (ndiff,anres,ndc/a)/active 9.300
# REPLACED overlap (m1,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1
- (m1,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1
  (pdiff,apres,pdc/a)/active 2.790
overlap (m1,nsc/m1,cwc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1 (pdiff,apres,pdc/a)/active 2.790
# REPLACED sideoverlap (m1,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1
- (m1,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1
  (pdiff,apres,pdc/a)/active 9.300
sideoverlap (m1,nsc/m1,cwc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1
- (m1,nsc/m1,cwc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1 (ndiff,anres,ndc/a)/active 9.300

#metal-polycap blocked by poly
#metal-polycap blocked by poly2
# REPLACED overlap (m1,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1
(poly,pres,wcap,pc/a,wcap,pc/a,nfet,pfet)/active 5.400
overlap (m1,nsc/m1,cwc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1
(poly,pres,wcap,pc/a,wcap,pc/a,nfet,pfet)/active 5.400
sideoverlap (m1,nsc/m1,cwc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1
(poly,pres,wcap,pc/a,wcap,pc/a,nfet,pfet)/active 15
sideoverlap (poly,pres,wcap,pc/a,wcap,pc/a,nfet,pfet)/active ~ (poly,pres,wcap,pc/a,wcap,pc/a,nfet,pfet)/active
(m1,ndc/m1,nsc/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1 15

#metal2 not blocked
#metal2
sidewall (m2,m2c/m2,m3c/m2)/metal2 ~ (m2,m2c/m2,m3c/m2)/metal2 ~ (m2,m2c/m2,m3c/m2)/metal2
(m2,m2c/m2,m3c/m2)/metal2 9.090
# REPLACED areacap (m2,m3c/m2)/metal2 1.260
overlap (m2,m3c/m2)/metal2 ss/ss 1.260 ~space/w,~space/a
#metal2-sub blocked by ~space/a,~space/m1,poly2/poly2/polycap
overlap (m2,m3c/m2)/metal2 ~space/w 1.260 ~space/a,~space/m1
# REPLACED perimc (m2,m2c/m2,m3c/m2)/metal2 ~ (m2,m2c/m2,m3c/m2)/metal2 5.100
sideoverlap (m2,m2c/m2,m3c/m2)/metal2 ~ (m2,m2c/m2,m3c/m2)/metal2 ss/ss 5.100 ~space/w,~space/a
sideoverlap (m2,m2c/m2,m3c/m2)/metal2 ~ (m2,m2c/m2,m3c/m2)/metal2 ~space/w 5.100 ~space/a,~space/m1
overlap (m2,m3c/m2)/metal2 (ndiff,anres,ndc/a)/active 1.260 ~space/m1
sideoverlap (m2,m2c/m2,m3c/m2)/metal2 ~ (m2,m2c/m2,m3c/m2)/metal2 (ndiff,anres,ndc/a)/active 5.100
~space/m1
overlap (m2,m2c/m2,m3c/m2)/metal2 (pdiff,apres,pdc/a)/active 1.260 ~space/m1
sideoverlap (m2,m2c/m2,m3c/m2)/metal2 ~ (m2,m2c/m2,m3c/m2)/metal2 (pdiff,apres,pdc/a)/active 5.100
~space/m1

#metal2-polycap by ~space/m1,poly
#metal2-poly blocked by ~space/m1,poly2
overlap (m2,m3c/m2)/metal2 (poly,pres,wcap,pc/a,wcap,pc/a,nfet,pfet)/active 1.620 ~space/m1
sideoverlap (m2,m3c/m2)/metal2 ~ (m2,m2c/m2,m3c/m2)/metal2
(poly,pres,wcap,pc/a,wcap,pc/a,nfet,pfet)/active 6.300 ~space/m1
sideoverlap (poly,pres,wcap,pc/a,wcap,pc/a,nfet,pfet)/active ~ (poly,pres,wcap,pc/a,wcap,pc/a,nfet,pfet)/active
(m2,m2c/m2,m3c/m2)/metal2  6.300 ~space/m1
#metal2-poly2 blocked by -space/m1

#M2->M1
overlap (m2,m3c/m2)/metal2 (m1,ndc/m1,ns/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1  4.050
sideoverlap (m2,m2c/m2,m3c/m2)/metal2 ~(m2,m2c/m2,m3c/m2)/metal2
(m1,ndc/m1,ns/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1  11.700
sideoverlap (m1,ndc/m1,ns/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1
~(m1,ndc/m1,ns/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1
(m2,m2c/m2,m3c/m2)/metal2  11.700

#metal3
 sidewall (m3,m3c/m3,pad)/metal3 ~(m3,m3c/m3,pad)/metal3 ~(m3,m3c/m3,pad)/metal3 (m3,m3c/m3,pad)/metal3  12.900
# REPLACED areacap (m3,pad)/metal3  0.810
overlap (m3,pad)/metal3 ss/ss  0.810 ~space/w,~space/a

#metal3-sub blocked by ~space/a,~space/m1,~space/m2,poly2/polycap
overlap (m3,pad)/metal3 ~(m3,m3c/m3,pad)/metal3 ~(m3,m3c/m3,pad)/metal3 (m3,m3c/m3,pad)/metal3  1.440
# REPLACED perimc (m3,m3c/m3,pad)/metal3 ~(m3,m3c/m3,pad)/metal3  3.300
sideoverlap (m3,m3c/m3,pad)/metal3 ~(m3,m3c/m3,pad)/metal3 ss/ss  3.300 ~space/w,~space/a
sideoverlap (m3,m3c/m3,pad)/metal3 ~(m3,m3c/m3,pad)/metal3 ~(m3,m3c/m3,pad)/metal3 ~space/w  3.300 ~space/a,~space/m1,~space/m2

#metal3-*diff blocked by ~space/m1,~space/m2,poly2/polycap
overlap (m3,pad)/metal3 (ndiff,anres,ndc/a)/active  0.810 ~space/m1,~space/m2
sideoverlap (m3,m3c/m3,pad)/metal3 ~(m3,m3c/m3,pad)/metal3 (ndiff,anres,ndc/a)/active  3.300
~space/m1,~space/m2
overlap (m3,pad)/metal3 (pdfiff,apres,pdc/a)/active  0.810 ~space/m1,~space/m2
sideoverlap (m3,m3c/m3,pad)/metal3 ~(m3,m3c/m3,pad)/metal3 (pdfiff,apres,pdc/a)/active  3.300
~space/m1,~space/m2

#metal3-polycap by ~space/m1,~space/m2,poly

#metal3-poly blocked by ~space/m1,~space/m2,poly2
overlap (m3,pad)/metal3 (poly, pres, wcap, pc/a, wcap, pc/a, nfet, pfet)/active  0.990 ~space/m1,~space/m2
sideoverlap (m3,m3c/m3,pad)/metal3 ~(m3,m3c/m3,pad)/metal3 (poly, pres, wcap, pc/a, wcap, pc/a, nfet, pfet)/active
3.600 ~space/m1,~space/m2
sideoverlap (poly, pres, wcap, pc/a, wcap, pc/a, nfet, pfet)/active ~(poly, pres, wcap, pc/a, wcap, pc/a, nfet, pfet)/active
(m3,m3c/m3,pad)/metal3  3.600 ~space/m1,~space/m2

#metal3-poly2 blocked by ~space/m1

#M3->M1

#metal3-metal1 blocked by ~space/m2
overlap (m3,pad)/metal3 (m1,ndc/m1,ns/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1  1.440
~space/m2
sideoverlap (m3,m3c/m3,pad)/metal3 ~(m3,m3c/m3,pad)/metal3
(m1,ndc/m1,ns/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1  5.400 ~space/m2
sideoverlap (m1,ndc/m1,ns/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1
~(m1,ndc/m1,ns/m1,cwc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1)/metal1
(m3,m3c/m3,pad)/metal3  5.400 ~space/m2

#M3->M2
overlap (m3,pad)/metal3 (m2,m2c/m2,m3c/m2)/metal2  4.230
sideoverlap (m3,m3c/m3,pad)/metal3 ~(m3,m3c/m3,pad)/metal3 (m2,m2c/m2,m3c/m2)/metal2  10.800
sideoverlap (m2,m2c/m2,m3c/m2)/metal2 ~(m2,m2c/m2,m3c/m2)/metal2 (m3,m3c/m3,pad)/metal3  10.800

#metal4

#fets

fet pfet pdiff,pdc 2 pfet Vdd! nwell 15 308
fet pfet pdiff,pdc 1 pfet Vdd! nwell 15 308

# REPLACED fet nfet ndiff,ndc 2 nfet Gnd! pwell 15 324
fet nfet ndiff,ndc 2 nfet $$G_{V_{sub}}!$$ pwell,ss 15 324
# REPLACED fet nfet ndiff,ndc 1 nfet Gnd! pwell 15 324
fet nfet ndiff,ndc 1 nfet $$G_{V_{sub}}!$$ pwell,ss 15 324

fetresis pfet linear 15658
fetresis pfet saturation 15658
fetresis nfet linear 4267
fetresis nfet saturation 4267

# ADDED
fet pres poly 2 polyres $$G_{V_{sub}}!$$ ss,nwell,pwell 0 0

# ADDED
fet wcap cwnsd 1 wcap $$G_{V_{sub}}!$$ 0 0

# ADDED
fet m1res m1,ndc/m1,nsc/m1,pdc/m1,psc/m1,cwc/m1,pc/m1,m2c/m1 2 m1res $$G_{V_{sub}}!$$ 0 0
# ADDED
fet m2res m2,m2c/m2,m3c/m2,m3c/m2 2 m2res $$G_{V_{sub}}!$$ 0 0
# ADDED
fet m3res m3,m3c/m3 2 m3res $$G_{V_{sub}}!$$ 0 0end

end