

# Proposed Verilog-A Language Extensions for Compact Modeling

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**Version 7**

Describes the changes that should be made to the Verilog-A and Verilog-AMS language definition [1] to better support compact device modeling [2]. This document is a product of an Accellera subcommittee which started work in May 2003 and included participants from the compact modeling community, semiconductor companies, and EDA vendor companies.

## 1.0 Parameters and Variables

### 1.1 Descriptions and units

Parameter declarations should provide a method for specifying the units of the parameter and a text description. This information would be used to generate documentation and help messages. For example, “spectre -help bsim4” or “hpeesof\_sim -help BSIM4” instructs the respective simulator to print a description of Berkeley’s BSIM4 compact model, including a list of parameters with units and descriptions.

*Examples:*

```
parameter real freq=1G "Hz" from (0:inf) (*desc="Center frequency");
parameter real kvco=1G "Hz/V" from (0:inf) (*desc="Gain of VCO");
parameter integer resolution=8 "bits" from [1:30] (*desc="Quantization level of converter");
```

Variables declared at module scope (outside the analog block) should also be allowed to have units and descriptions. Those that do should be considered “output or operating point parameters,” meaning that simulators should print the name, value, units, and description of these variables (and no others) when printing operating-point information for the circuit. Operating point values, such as *vth*, *vdsat*, or *cgs* for the BSIM4 compact model, are frequently used in the design of circuits. (See also §3.2 for operating point values that are computed using derivatives.)

*Examples:*

```
real vdsat "V" (*desc="drain-source saturation voltage");
real cgs "F" (*desc="gate-source capacitance");
```

Since essentially all compact model parameters should be described in such a way, it is important to have an efficient and standard method of attaching this information. However, since the text description does not affect the numerical results of the simulation, it is reasonable to place this as an attribute; digital Verilog simulators will be able to ignore this attribute.

Note that there is no precedent for a standardized attribute in Verilog-AMS nor Verilog 2001. The SystemVerilog committee is apparently considering some standard attributes. We would like to standardize this attribute so that all SPICE-like simulators will work the same.

Units may affect the simulation, in particular through the ‘**timescale**’ directive in Verilog 1364-1995, and hence parameter units should not be implemented through an attribute. The ‘**timescale**’ directive is unfortunate, since it affects numbers throughout the module. We do not propose that any “units mathematics” be performed; however, it should be clear to the user what the units are. BSIM3 allowed one to specify the parameter U0 in non-MKS units; if the value was greater than 1.0, it was assumed to have been specified in cm<sup>2</sup>/Vs instead of m<sup>2</sup>/Vs.

**Proposal:** for parameters, change Syntax 3-2 and the corresponding lines of A.4 as follows.

```
declarator_init ::=
  parameter_identifier = constant_expression range_description
  | parameter_array_identifier range = constant_param_arrayinit range_description
range_description ::=
  [units_string] { opt_value_range } [ (*desc=description_string*)]
```

**Proposal:** for variables, create new declaration syntax for `output_integer_declaration` and `output_real_declaration`.

```
output_integer_declaration ::=
    integer list_of_output_identifiers ;

output_real_declaration ::=
    real list_of_output_identifiers ;

list_of_output_identifiers ::=
    output_var_name { , output_var_name }

output_var_name ::=
    variable_identifier
    | array_identifier array_range
    [units_string] [(*desc= description_string *)]

array_range ::=
    [ upper_limit_constant_expression : lower_limit_constant_expression ]
```

These new output declarations will be syntactically restricted to occur only at module scope; see §1.6. They replace `integer_declaration` and `real_declaration` at that scope, since the attribute is optional; `integer_declaration` and `real_declaration` will remain defined for `block_item_declarations`.

## 1.2 Detecting whether a parameter was specified

There are currently many instances in existing compact models where the code checks whether the user actually specified a value for a parameter. This detection is often used for overrides, where an unspecified parameter is determined from others; Verilog-A already supports computation of default values from (previously-declared) parameters. However, in some cases, the override equation is very complicated, and sometimes it is circular. One example of circularity involves `kp` and `uo` in MOS level-3 models; these parameters take their defaults from each other. If `uo` is specified, `kp` takes its default from  $kp=uo*tox$ . If `uo` is not specified, but `kp` is, then `uo` is calculated from `kp`. This circularity cannot be achieved presently in Verilog-A within the parameter declaration syntax. The BSIM3 threshold voltage is another example: one may specify it through the flatband voltage `vfb` or through `vth0`; the calculation of one from the other is much more complicated than in the MOS level-3 model. In the Verilog-A models that Silvaco has posted on the web [3], `vfb`, `vth0`, and `vtho` (an alias for `vth0`) all have default values of -99.0, which acts as a flag for making the correct computation. This method is not elegant nor generally applicable, since -99.0 (or any other value) might be a legitimate value for a particular parameter. Even when there are illegal values of the parameters that could be used as flags, this will likely force the parameter range to include other illegal values. Using a negative value for BSIM3's `cgdo` parameter means that the range for `cgdo` cannot forbid negative values. Further, any simulator that reports model card values for the simulation as part of an operating-point report will print these strange values.

A much better solution would be to provide a method of directly checking whether a parameter was specified. A reasonable default value may then be provided. ADMS [4] has a `$given` function and

Cadence reportedly provides the `$param_given` function (in internal versions of their simulator) for the purpose of determining whether a parameter value has been specified.

**Proposal:** add the `$param_given` system task, which can appear in a `genvar_expression` as well as an expression. The returned value would be one (1) if the parameter was specified either by its name or one of its aliases, or zero (0) if its value was computed from the default expression.

```
genvar_primary ::=
  constant_primary
  | genvar_identifier
  | genvar_identifier [ genvar_expression ]
  | analog_system_task

analog_system_task ::=
  analysis( arg_list )
  | $param_given(module_parameter_identifier)
  | $port_connected(port_scalar_expression)
```

See §4.1 for `$port_connected`.

The argument to `$param_given` must be a parameter name, not an alias (see the next section).

One might want to ask about more than parameter in a single call, but it is not clear whether the result should be true if all parameters are given or true if any of the parameters is given. The former would likely be more useful (if all the required parameters are specified, then use them to make a calculation). However, this would be inconsistent with the `analysis()` function, which returns true if any of its arguments matches the current analysis type.

### 1.3 Parameter aliases

To support backward compatibility to older models, and to provide convenient alternatives for parameter values, parameter aliases should be supported. For example, in older BJT models the `vaf` and `var` parameters are named `va` and `vb`. The values specified for aliased parameters are treated as though the user had specified using the target name. Thus, if the user specified `va=20`, the simulator treats it as if the user had specified `vaf=20`, and if the simulator reports the values of parameters for the simulation, it would list the value under `vaf` and not under `va`. Also, when the user asks for help, the `vaf` parameter is listed, but the `va` parameter is not.

Although some of this functionality can be handled by the language as it exists today by chaining defaults, there are two drawbacks. Consider `vaf` and `va` as an example. If the parameters are declared with

```
parameter real va = inf;
parameter real vaf = va;
```

and the model equations use `vaf`, then the user may specify either and get the right behavior. However, it is not possible to prevent both from being specified (with different values), and both parameters would appear in a list of parameter values generated by the simulator. Consider also the parameter used to indicate a difference in device temperature relative to the circuit, called variously `trise` (Cadence's Spectre), `dtemp` (Synopsys' HSpice), or `dta` (Philips' Mextram 504 model). Even if the aliased parameters all had the same value, there is potential confusion; if all three were specified as 10, or if the simulator printed a value of 10 for each alias, the user might think or expect that the device was 30 degrees above ambient.

**Proposal:** add the keyword **aliasparam** and use the syntax

```
parameter_alias_declaration ::=
  aliasparam alias_identifier = parameter_identifier;
```

Model equations must be written in terms of the original parameter name (*vaf*). Multiple aliases may be assigned to a single parameter. An *alias\_identifier* may not conflict with a *parameter\_identifier*. The simulator should report an error if a parameter is specified by more than one name (an alias and the original parameter name or two aliases), either on the instance line or through a paramset (see §2.0).

Note that **aliasparam** handles only exact aliases. SPICE allows one to specify the device temperature directly with the parameter *temp*, rather than as an offset from **\$temperature** as is done for *trise*. One would need to use **\$param\_given** to allow a model to use either.

#### 1.4 Multiplicity factor

This is an extremely important capability. It is heavily used and valued by designers. The multiplicity factor is a parameter  $m$  that can be applied to an instance of a model that acts to scale the instance in such a way that there appear to be  $m$  instances in parallel. This parameter must be handled hierarchically, so that if one instance is contained in another, then the effective multiplicity factor of the first instance is the product of multiplicity factors explicitly specified for both.

The omission of the multiplicity factor from the Verilog-AMS LRM led to some unfortunate inconsistencies. Within a Verilog-A netlist or module, one may reference a subcircuit defined in SPICE or in native Verilog-AMS, but only the SPICE subcircuit will accept the multiplicity factor.

The actual behavior of the multiplicity factor must be handled automatically by the simulator rather than forcing the user to explicitly code the proper behavior in to the model. The latter would result in multiplicity factors being inconsistently or incorrectly implemented and interpreted, which would eventually make the multiplicity factor useless. As such, the multiplicity factor must be an implicit parameter to each instance that multiplies the (deterministic) current contributed to any node or branch by the instance and divides the (deterministic) current probed by the instance. Note that, while deterministic current contributions should be scaled by  $m$ , noise currents should be scaled by  $m^{1/2}$  and noise voltage by  $m^{-1/2}$ .

In principle, mismatch parameters should also be scaled by  $m^{-1/2}$ . For example, if there are  $m$  resistors in parallel, each with a given standard deviation  $\sigma$  of their resistance, the parallel combination will have a standard deviation of resistance equal to  $\sigma/m^{1/2}$ . However, there is presently no method in Verilog-A for requesting a random variable with a particular standard deviation. Hence, the module can only treat the standard deviation as an offset, and any Monte-Carlo or Monte-Carlo-like simulations must be handled at a different level.

An unfortunate consequence of this decision to handle  $m$  automatically is that the simulator will likely have to implement this as a post-processing of all branch quantities and their derivatives for each iteration of each timepoint. Some compact models use the multiplicity factor to pre-scale parameters once for an analysis. We recognize this inefficiency, but feel that it is justified in the quest for ensuring correct implementation of the multiplicity factor for all models. We note that, for compact models of moderate complexity, the multiplications will probably not be a significant fraction of the floating-point operations involved in calculating the model. Further, the simulator may be able to find a way to optimize the scaling.

In addition to the automatic scaling of currents, there are times when the multiplicity factor would reasonably be expected to affect the values of output and operating point parameters (§1.4). For example, consider a resistor whose effective resistance is computed by  $r = (w*rsh)/(l*m)$ ; one might also want to

compute the effective current or power of the resistor. Not providing access to the multiplicity factor provides a certain consistency; all output and operating point parameters would necessarily be computed from the perspective of the individual elements of the parallel array. However, that is not always what the users want, so there is value in providing composite quantities. However, the model writer must not be allowed to interfere with the proper interpretation of the multiplicity factor, so that if the value of the multiplicity factor is made available in the module, it must not be allowed to affect the behavior of the model (the behavior of the model has already been appropriately modified by the simulator). For this reason, it should be an error for the output signal values to depend on the value of the multiplicity factor that is explicitly made available within the module.

The effect of OOMRs, or “out of module references,” must be considered. OOMRs are the observation of one module’s signals or variables from another module. The variables in a module will not be scaled by  $m$ , because the simulator will not know what the proper scaling is (unless it is an output parameter that the model writer has scaled), so necessarily variables referenced out-of-module must refer to a value for a one of the  $m$  parallel devices. On the other hand, it seems that signals, specifically currents, referenced out-of-module should refer to the actual current in the branch, for all the devices in composite. If the module that asks for the OOMR is a current-dependent source, it should get the correct current. This inconsistency may be confusing for model writers.

One last issue to consider is *rmin*. Some simulators (e.g., Cadence’s Spectre) eliminate a parasitic resistor if its value falls below a certain threshold, called *rmin*. If we do not allow access to  $m$  from within the model, we cannot determine the actual value of a resistor, only the unscaled value. This could be addressed with a branch parameter (a concept that does not yet exist in the language). The idea would be to pass *rmin* to the branch, and let it determine whether the branch should be shorted or not. Alternately, we could supply a function to determine if the scaled value is below a threshold. That function would be something like **m\_scaled\_value\_above**(quan, thresh, expon), which would return true if

$$\text{quan} * \text{pow}(m\text{factor}, \text{expon}) > \text{thresh}$$

Resistances would use  $\text{expon}=-1$ ; a conductance formulation or a parasitic capacitance might use  $\text{expon}=+1$ .

*Example:*

```
rmin = 0.001;
if (m_scaled_value_above(rseff, rmin, -1.0))
    I(s,si) <+ V(s,si)/rseff;
else
    V(s,si) <+ 0.0;
```

The main obstacle to the inclusion of a multiplicity factor in Verilog-AMS at this point is the concern that defining  $m$  as a reserved keyword would break a number of existing modules. One solution would be to use *mfactor* as the keyword for Verilog-AMS netlists, and recommend that a Verilog-AMS simulator reading a SPICE netlist should translate  $m$  to *mfactor* if it encounters  $m$  on an instance of a Verilog-AMS module in that SPICE netlist.

**Proposal:** implement the  $m$ -factor automatically; OOMRs, *rmin*, output parameters, and mismatch need to be addressed.

## 1.5 String parameters

For compact modeling, essentially all transistor models have one set of equations that is used for both p- and n-type devices, generally by multiplying voltages and currents by  $-1$ . In printing device information, simulators always use a string to tell the user whether the device is p-type or n-type (NMOS or PMOS, NPN or PNP, etc.). Users will expect Verilog-A device models to give the same information, without

requiring them to translate a numerical value of +1 or -1. (Most compact model equations are written for the n-type device, and values are multiplied by -1 for p-type; however, one could imagine a novice model writer or a designer to expect “negative one” to mean n-type, causing a great deal of confusion and/or wrong results.) Some simulators also use strings in the netlist to specify the device, eg, type=“npn”.

String variables would also be helpful for debugging; one might want to print debugging information only for certain instances of a model. Here, one would want to compare a string constant to the value of special function such as \$instanceName or \$modelName.

A sort of “string variable” is permitted in Verilog-AMS for the digital domain: an array of type **reg**, but these are excluded from Verilog-A. We do not need the ability to operate on the variables (concatenation or character replacement); we only need to be able (1) to pass the value in from a netlist and (2) to compare a value from a netlist or a special function to a constant string. We do not need our strings to interact at all with **reg** arrays from the digital domain.

Strings, that is, constant strings, are also permitted in Verilog-AMS expressions. Strings are used as arguments to some functions (e.g., **analysis()**), so Annex C.2 should be updated to remove the exclusion of strings from Verilog-A.

SystemVerilog has a variable type **enum** which addresses some of the needs here: one could have values such as npn and pnp. However, SV does not allow parameters of type enum, and it is not clear how the netlist would support this: enum identifiers are reserved in the module, but should not be in the netlist.

It seems cleaner to simply allow string parameters.

**Proposal:** allow string parameter declarations and allow comparison with the “==” and “!=” operators.

**string\_parameter\_declaration ::=**

**parameter string parameter\_identifier = string from {string {,string}};**

The set of allowed strings for a parameter must be specified explicitly in the declaration.

## 1.6 Module item declarations

The following is the new definition of module\_item\_declaration, including the items from §§1.1, 1.3, and 1.5.

```
module_item_declaration ::=
  parameter_declaration
  string_parameter_declaration
  parameter_alias_declaration
  digital_input_declaration
  digital_output_declaration
  digital_inout_declaration
  ground_declaration
  output_integer_declaration
  output_real_declaration
  net_discipline_declaration
  genvar_declaration
  branch_declaration
  analog_function_declaration
```

*digital\_function\_declaration*  
*digital\_net\_declaration*  
*digital\_reg\_declaration*  
*digital\_time\_declaration*  
*digital\_realtime\_declaration*  
*digital\_event\_declaration*  
*digital\_task\_declaration*

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## 2.0 Paramsets

The following items are covered in a separate proposal entitled “paramset: a Verilog-A/MS Implementation of SPICE. model Statements.” I need to merge these two proposals and formalize the syntax.

### 2.1 Different model levels

Used to support multiple versions of the models optimized for different applications. For example, one might use a simple version of the model to support digital transistors, a more sophisticated model for analog transistors, and an even more sophisticated model for RF transistors; additionally, the model file might contain rules for Monte-Carlo simulations of process variation. The model compiler would then create multiple versions of the model that all have the same name. Then, through an as yet undefined mechanism, the user would specify which flavor of the model to use for each instance.

This might consist simply of a parameter or parameters passed in that indicate the level or flavor of the model desired. Simulator should treat this parameter as special in that the user will want to be able to hierarchically set its value (perhaps with wildcard characters). In this aspect, it may be similar to the multiplicity factor.

### 2.2 Model and instance parameters

Some parameters for compact models are generally associated with the manufacturing process, whereas others are specific to instances. For example, the oxide thickness of a MOSFET model is a “model parameter,” whereas the gate length is an “instance parameter.” It is inefficient to require the simulator to keep an array of values for every parameter of a compact model for each instance; simulators generally use model cards to accumulate shared model parameters. Verilog-A does not distinguish between the two types of parameters, and thus some simulators do, in fact, have a complete array of parameters for each instance.

One company (Tiburion-DA) has implemented a method that allows use of model cards with Verilog-A modules; although one can specify “instance” parameters in a model card (which the model writer might like to prevent), it does not require the simulator to keep track of a full array of parameters for each instance.

The subcommittee will investigate whether this approach is viable for other simulators. Many SPICE-like simulators would suffer from memory constraints if they cannot implement a similar scheme.

### 2.3 Binning

Binning is the ability to chose a model based on instance parameters.

## 3.0 Functions

### 3.1 More flexible functions

Analog functions in Verilog-A are restricted to having only one return value. This is a problem for the BSIM4 model, which contains a geometry function that returns four values: the drain and source areas and perimeters; intermediate calculations are made that would have to be repeated if the four values had to be computed in four separate functions. We should extend Verilog-A to support multiple return values.

One suggestion is to allow function definitions to use the **output** and **inout** keywords in addition to **input** to describe their arguments.

```

analog function integer BSIM4PAeffGeo;
  input geo, minSD, Weffcj, DMCG, DMCI, DMDG;
  output Ps, Pd, As, Ad;
  integer geo, minSD;
  real Weffcj, DMCG, DMCI, DMDG, Ps, Pd, As, Ad;
  ...
endfunction

```

The SystemVerilog LRM says that **inout** arguments are copied in at the start of the function and copied-out when the function completes; SystemVerilog has a new type, **ref**, for variables that are passed by reference. However, the analog block in Verilog-AMS does not allow for blocking or waiting, so from a simulation standpoint, analog function evaluation is instantaneous, and the copy-in, copy-out mechanism is unnecessary. Simulators may implement **inout** arguments for analog functions as pass-by-reference; it is not necessary to designate “ref” as a keyword. Since **inout** and **output** are allowed for module ports, there is some symmetry to allowing these for functions as well.

A completely different alternative would be to allow return values to be arrays; however, this idea was rejected by the Verilog-AMS committee in another context.

**Proposal:** allow **output** and **inout** arguments for functions

```

function_item_declaration ::=
  input_declaration
  | inout_declaration
  | output_declaration
  | block_item_declaration

```

Note that the compiler will have to check that any call of the analog function has an lvalue for any argument that the function declares to be **inout** or **output**.

### 3.2 Access to derivatives

Many operating point parameters are derivatives with respect to the signals on terminals or nodes. Most if not all of these derivatives are internally calculated by the simulator, along with other derivatives, for the purpose of loading the Jacobian matrix. Rather than forcing the user to manually compute these values, which would be both tedious and error-prone, it is desirable to give direct access to the derivatives. For example, consider computing the gm of a transistor using

$$gm = \text{ddx}(I_{ds}, V(g))$$

where **ddx** is a function that takes a variable and a node potential identifier, and returns the symbolic partial derivative of the variable with respect to the node potential, holding all other node potentials fixed, and evaluated at the current operating point.

In the case

```
if (V(g) > 0)
  Ig = V(g) * ggcond;
else
  Ig = -V(g) * ggcond;
```

the derivative of  $I_g$  is not continuous (mathematically, it is said not to exist). The **ddx** function will evaluate the symbolic derivative based on the branch of the **if** statement that was actually taken, so that when  $V(g)=0$ ,  $\text{ddx}(I_g, V(g)) = -\text{ggcond}$ . The derivative is symbolic, so no tolerance is necessary (unlike the **ddt** function).

The name **ddx** was chosen to be similar to **ddt**, which is the time derivative. It might be preferable to use **\$ddx** so as not to reserve another keyword; however, we feel it is better to be consistent with the notation **ddt**. If **ddt** becomes **\$ddt**, then **ddx** should also change to **\$ddx**. Another thought was that **ddv** might be more intuitive for electrical models; however, **ddx** preserves the generality of the language.

The first argument to the function can be any `analog_expression`, that is, anything that can be used as the right-hand side of an `analog_branch_contribution`, because the simulator needs to compute the derivative of anything contributed to a branch.

The second argument to the function must be a single node potential or a branch flow; it may not be a branch potential. Node potentials and branch flows are the independent variables in modified nodal analysis used by SPICE-like simulators. By restricting the second argument to be one of the independent variables, it is clear that all the other variables are held constant when taking the partial derivative. While the BSIM3 model generally uses  $V_{gs}$ ,  $V_{ds}$ , and  $V_{bs}$  as its independent variables, it sometimes uses  $V_{gd}$ ,  $V_{sd}$ , and  $V_{bd}$  (in “reverse mode”), and so it would be difficult for the simulator to automatically determine what is meant by an expression like  $\text{ddx}(I_{ds}, V(g,s))$  if  $I_{ds}$  is not calculated directly from  $V_{gs}$ .

**Proposal:** implement the **ddx()** function

```
ddx_call ::=
```

```
ddx(expression, potential_access_identifier(net_or_port_scalar_expression));
| ddx(expression, flow_access_identifier(branch_identifier));
```

---

## 4.0 Ports and Nodes

### 4.1 Optional ports

Optional ports (optional terminals in the language of device modelers) would be helpful for supporting the SPICE BJT model, as well as models that use a variable number of ports, such as switches, controlled-sources, diffusion or polynomial resistors, etc. In the case of the Mextram 504 or VBIC BJT models, which each have both an optional substrate terminal and an optional thermal terminal, a model writer might be forced to write four separate Verilog-A modules to cover all the cases.

Verilog already has the concept of optional ports; if a connection is not specified, then the port is left floating. An unconnected port of a digital module is in the high-impedance state, which acts similar to unknown for an input and also allows the module to drive it as an output.

Floating the port works in many cases, such as the thermal terminal of a BJT model with self-heating. Sometimes the designer may want to connect this port to model heat flow, or simply to monitor the device temperature. If the port is not connected, the self-heating effects should still be calculated.

However, floating the port would not work for optional ports that have only a capacitive connection to the rest of the device. An example might be a 3-terminal polysilicon resistor model, where the substrate terminal would be capacitively connected to the device, if connected, but if left unconnected, the model might be expected to work as a simple 2-terminal resistor for faster simulation. Most SPICE-like simulators would object to the “no dc path to ground” condition arising from floating this port.

A solution to this problem would be the implementation of a `$port_connected` function that would return 1 if the port is connected in the netlist and 0 if not; this would be similar to the `$param_given` proposed in §1.2. Although it is somewhat inelegant, and perhaps dangerous, to allow such direct access to the circuit topology, this proposal covers all the situations we can imagine.

Another possibility would be to allow specification of defaults for ports, as is done for parameters. For example,

```
module mosfet(d,g,s,b=s)
```

in which the body node defaults to the source. The use of defaults could considerably simplify the writing of multi-port behavioral models, such as controlled sources; using the `$port_connected` function would require testing each optional port explicitly. However, most compact models do not have many optional terminals (the most we are aware of is BSIMSOI with three). It is also relevant to note that BSIMSOI uses parameters to determine whether the extra terminals are allowed (only for “level=9”) and to determine the meaning of the optional terminals, if fewer than three are specified. It does not seem possible to handle this functionality with port defaults.

It may be reasonable to add port defaults to the language to handle multi-port behavioral models, but this request should not come from the compact modeling subcommittee. In fact, the specification of a default in the example above might cause headaches for a designer: if the body terminal of a MOSFET is not connected in a netlist, the circuit would still simulate as intended, but the actual silicon would not perform correctly.

The determination by the simulator of which ports are connected for an instance will follow standard Verilog rules for connecting ports by name or by order. Note that these rules do not make special considerations for optional ports: connection by ordered list will connect following the order in the module definition, even if that means that a port that was intended to be optional (because it is tested with `$port_connected`) is connected but a non-optional port is left floating. SPICE-like simulators suffer a similar limitation: if you instantiate

```
q1 c b s vnpn
```

the simulator will not know that you intended to connect the node “s” to the optional substrate terminal instead of the emitter.

**Proposal:** implement the `$port_connected` function. See §1.2 for the proposed syntax.

## 4.2 Descriptions

Similar to (§1.1), module ports should have descriptions, so that one may name the collector terminal of a bipolar transistor as “c” to match SPICE behavior and save typing, yet have the full description appear in any help information generated by the simulator (“spectre -help bjt”).

```
module BJT(c,b,e);
  inout c, b, e;
  electrical (*desc="collector"*) c,
```

```
(*desc="base") b,
(*desc="emitter") e;
```

**Proposal:** allow descriptions of terminals as follows:

```
net_discipline_declaration ::=
  discipline_identifier [range] list_of_nets;
list_of_nets ::=
  [(*desc= string*)] net_identifier [range]
  | [(*desc= string*)] net_identifier [range], list_of_nets
```

The description applies to all *net\_identifier* in the *list\_of\_nets* until the next description is encountered. For example,

```
module AND(in1,in2,in3,out);
  electrical (*desc="input pin") in1, in2, in3, (*desc="output pin") out;
```

would attach the description “input pin” to all three input pins.

It has been proposed elsewhere that **inout** and **electrical** be allowed in the same line, e.g.,

```
inout electrical c, b, e;
```

for a cleaner-looking syntax; the description attribute could precede the list of nets in this form, as well.

## 5.0 Simulation

### 5.1 Non-repetitive warnings/notices

It is often useful to monitor device behavior and print messages when they change state. For example, imagine a BJT model that included code that monitored the region of the transistor and printed warnings when the transistor entered saturation. The warning should be printed only once, when the transistor enters saturation, and not on every timestep or step of a dc analysis.

This could be handled by printing the warning from within an event block, such as one defined by the **cross** function. Unfortunately, the **cross** event does not have behavior defined for dc analysis, and thus simulators may have implemented different behavior, or model writers may have assumed that it does nothing in a dc analysis.

Cadence’s Spectre has added a new event, **above**, that performs just like **cross** during a transient analysis, but also triggers during a dc analysis. In particular, if the inequality is true for the dc (time-0) solution preceding a transient analysis, the **above** event is triggered, unlike the **cross** event.

The **above** event as implemented in Cadence’s Spectre remembers its state between points of a dc sweep, just as it and **cross** do for timepoints of a transient analysis. (The state is cleared for each new analysis.) According to Cadence, this was not an intentional design goal; however, as implemented, the **above** event works perfectly for avoiding repetition of a message during a dc sweep.

**Proposal:** standardize Cadence’s **@above** event for detecting crossings in dc analysis

This might be a good point to mention that the current Verilog-AMS LRM does not specify what happens to variables and internal state during a dc sweep or between analyses. We would like to standardize on the following:

1. Variables and any internal state of models should be remembered from point to point of a dc sweep.

2. All variables and internal state should be reset or re-initialized at the start of each new analysis, that is, the simulator should not remember the values of variables from one analysis to the next.

## 5.2 Simulator parameter access

There are a variety of simulator parameters that models need access to. Examples include *gmin* and *scale*, which are present in almost all SPICE-like simulators. In addition, there may be many simulator-specific parameters, such as *cmin*, *shrink*, and perhaps a “homotopy” or “continuation” parameter (used to improve convergence). Some mechanism must be made to provide access to these values. This mechanism could also be used to pass a diagnosis flag into the models.

We recommend providing a function that takes a string and returns a value, similar in operation to the **analysis** function. And like with the **analysis** function, recommended names should be given to help ensure consistency between the various implementations. Some consideration should be made on how to handle unknown parameters. For example, a model might be written to work with a specific homotopy that is not available in all simulators; rather than having the model keep track of the names of all the simulators that support the homotopy, the model could simply ask whether the homotopy variable is known to the simulator. The access function takes one additional optional argument. If the second argument is specified, it is the value returned by the access function if the parameter name is unknown to the simulator. If the optional argument is not specified, then an error should be generated by the simulator if the parameter name is not known. Thus,

```
gmin = $simparam("gmin");
```

would obtain the value of *gmin*, if it is known, and generate an error otherwise;

```
sourcescale = $simparam("source_scale_factor", 1);
```

would obtain the value of the scale factor for source-stepping, if that continuation method is supported; otherwise the value 1 would be returned but no error would be generated, so the model could simply multiply any independent sources by the returned value, without needing to know if source-stepping is supported and active, supported but not active, or not supported.

The value returned by **\$simparam** would be the value used by the simulator responsible for evaluating the module that makes the call; this distinction is necessary for multi-simulator mixed-signal simulations.

Verilog 2001 uses **\$value\$plusargs** to obtain a value passed to the simulator. According to the Verilog 2005 draft LRM (2001 is assumed to be similar), if the simulator is invoked with the argument +FINISH=10000, then **\$value\$plusargs**("FINISH", endtime) will return the integer one and set the variable endtime to 10000. If the simulator is invoked without the argument, then the function call returns zero and does not alter the value of endtime. Following this pattern, one could use **\$value\$simargs** for simulator values. The main drawback is that, if a module wanted to require a value for *gmin*, it could only generate a user-level message (eg, with **\$strobe**), rather than generating a simulator error.

**Proposal:** implement the **\$simparam()** function with three arguments; create a table with “standard” parameter names

```
simparam_call ::=
  $simparam(string)
  | $simparam(string, constant_expression)
```

The following is a tentative list of standard parameter names, which should go in a table in Annex E, SPICE compatibility. This table should include descriptions such that simulator vendors can identify the intended parameter in case the name is not the same.

```

gmin
gdev
scale
shrink
simulatorVersion
simulatorSubversion
tnom
imelt
imax
iteration
source_scale_factor

```

### 5.3 Simulator specificity

Individual simulators have individual quirks that must be supported. For example, Cadence’s Spectre wants to install *gmin* slightly differently than most simulators. These differences need to be accommodated. Further, some simulators may support non-standard syntax such as further proprietary extensions to the language. Thus, some mechanism must be provided to allow models to determine which simulator is running them.

In order to handle the non-standard syntax, this mechanism must be in the form of a **‘define** that is specific to the simulator. This is similar to how C code is made portable; the source contains statements like

```
#ifdef HPUX11
```

or

```
#ifdef SUNOS4
```

and a compiler running on a SunOS4 system will automatically define SUNOS4 to be true.

We recommend pre-defining tokens using the format “company\_simulator” to ensure that the tokens are unique (recall that “spectre” was originally the name of a harmonic-balance simulator at Berkeley). For example, a module could contain the following code:

```

#ifdef CADENCE_SPECTRE
    install gmin their way
#else
    install gmin the usual way
#endif

```

When running in Cadence’s Spectre simulator, CADENCE\_SPECTRE would be pre-defined true by the simulator, and thus *gmin* would be installed appropriate to that simulator.

One might want to check what version of the simulator is running; however, this would require a preprocessor directive **‘if** and some inequality operator such as  $\geq$ . Rather than checking the version for each simulator, the module should be able to check whether the compact modeling extensions in this proposal have been implemented by the simulator. Simulators that support Verilog-AMS will **‘define** `__VAMS_ENABLE__`; we propose **‘define** `__VAMS_COMPACT_MODELING__`. The Verilog-AMS committee should consider other tokens when further blocks of revisions are incorporated into the LRM.

**Proposal:** require simulators to provide (and document) a token that will be pre-defined by the simulator; define the token `VAMS_COMPACT_MODELING` if the extensions in this proposal have been fully implemented.

## 5.4 Convergence aids

Compact model equations are frequently strongly nonlinear and Newton-Raphson iteration is often unable to find solutions, particularly for the dc operating point or time=0 solution. Most SPICE-like simulators employ some sort of modified or damped Newton-Raphson algorithm; the damping algorithms are known as “limiting.” Presently, Verilog-A only supports limiting through the **limexp** function. Other limiting functions are necessary to provide the level of performance users expect for small circuits. The **limexp** function is specific to the exponential, but there are many other nonlinear functions that cause trouble for Newton-Raphson. Also, the **limexp** function is an analog operator, which restricts its use. For example, bipolar transistor models make frequent use of the diode current equation, though with different branch voltages and saturation currents. It would be convenient to write the current equation once in an analog function, but **limexp** may not be used in this context. The SPICE BJT model also uses the same branch voltage in two diode current computations (“ideal” and “nonideal”); it would be more efficient to limit the branch voltage once, rather than making two calls to **limexp**.

Large circuits frequently fail to converge even with limiting, and homotopy methods are needed. We believe that homotopy algorithms can be implemented using the simulator parameters from §5.2. For example, a *gmin*-stepping homotopy will be supported automatically if the compact model module references the simulator’s *gmin* rather than using a model parameter or constant.

Ideally, the Verilog-A simulator would recognize the nonlinear equations and automatically determine appropriate algorithms to improve convergence. For example, the simulator could automatically use **limexp** in place of **exp**. However, current implementations of Verilog-A are not sufficiently sophisticated and simulator vendors tend to treat their limiting algorithms as trade secrets.

The art of writing limiting functions is subtle and difficult, and the appropriate function may depend on the simulator. Model writers should not have to worry about details of the limiting function, however, they are aware of the two main traditional limiting functions, known as “pnjlim” (for PN junctions, including those in bipolar transistors) and “fetlim” (for MOSFETs and JFETs). Some simulators convert calls to **limexp** into calls to pnjlim. Versions of these two algorithms are implemented in SPICE and many SPICE-like simulators. We propose the following syntax to allow model writers to access these two common limiting functions, as implemented in the current simulator.

```
vdio = $limit(V(a,c), "pnjlim", vcrit);
vgs = $limit(V(g,s), "fetlim", vto);
```

The simulator need not actually implement the limiting function requested; Spectre does not use fetlim because Cadence feels that circuit performance is better without it (multiple homotopies aid in dc convergence, and limiting requires storage of a double-precision number). If the simulator does not implement the limiting algorithm, or if limiting is not appropriate for the analysis, the **\$limit** function simply returns the value of the first argument.

The syntax could also be used simply to alert the simulator to a strong nonlinearity, in the case that the model writer does not know the appropriate limiting function:

```
vds = $limit(V(d,s));
```

Some simulators put a minimum conductance, similar to *gmin*, in parallel with strongly nonlinear branches; this “gdev” is decreased all the way to zero for a converged solution (*gmin* homotopies typically reduce *gmin* to some small but non-zero value). This **\$limit** call could be a hint to the simulator to add “gdev” in this branch.

If the simulator does implement the requested limiting algorithm, it is responsible for storing the previous value between iterations and restoring the correct value in the case of a rejected timepoint. The simulator is also responsible for generating a value to use for the first iteration. In SPICE, pnjlim typically returns the argument *vcrit* for the first iteration, unless a better value is known from a nodeset file; simi-

larly, `fetlim` returns the argument `vto`. These values are chosen to initialize the device in such a way that the conductance of the nonlinear branch is reasonable. If limiting occurs, the simulator may need to apply a so-called “limiting correction” to correct for the fact that currents were not computed at the voltages requested by the simulator. (SPICE itself does not use the currents directly because it solves for the new voltage vector instead of the voltage increment; the so-called SPICE right-hand side does not need a limiting correction.) When correcting a current for limiting, one adds to the current a term composed of the derivative of the current with respect to the limited voltage multiplied by the difference between the voltage requested by the simulator and that used in the equations.

The proposed syntax was tested in the internal circuit simulator at Analog Devices, using test cases from the “CircuitSim93” set of circuits [2]. The results were rather convincing for MOSFET circuits: 13 of the 14 MOS level-3 circuits converged in dc Newton using a built-in model with limiting, whereas only 3 converged using a Verilog-A model without any limiting. Adding limiting to the Verilog-A model resulted in 13 circuits converging (though the non-converging circuit was different). For the BJT circuits, 9 of 12 converged with a built-in model with limiting, but only 4 with the Verilog-A model without limiting. Adding limiting to the Verilog-A resulted in convergence in the same 9 circuits as the built-in model. The actual number of iterations varied somewhat, because the proposed syntax does not provide for specifying an initial guess; the base-collector voltage is traditionally started at 0V whereas the base-emitter starts at `vcrít`. For this set of examples, the ability to specify the initial guess was not necessary to obtain similar performance to the built-in models.

**Proposal:** implement the `$limit` function

`limit_call ::=`

`$limit(access_function_reference [, string, expression])`

The strings “`pnjlim`” and “`fetlim`” should connect with the appropriate algorithm. Simulators are free to accept other strings, if they have other limiting algorithms. The expression should not depend on the voltages; however, it may depend on `$temperature`, so a `genvar_expression` is too restrictive. Perhaps the restriction on the expression is semantic.

## 5.5 Diagnosis modes

Frequently, when developing a model, the author needs to understand the achieved behavior of the model and why it differs from the expected behavior. Generally, this requires printing extra values during simulation. In particular, it is frequently useful to print the value of a variable on every iteration rather than only after convergence.

Cadence’s `$debug` function prints the value on every iteration and accepts the same arguments as `$strobe`. There is a proliferation of functions that print values: `$strobe`, `$write`, `$display`, `$monitor`; unfortunately, each has a particular meaning and it does not appear possible to adapt one of them for the purpose of printing on each iteration.

Model authors may also want to enable extra diagnostic information for users of the model, conditional on a diagnosis flag being set in the simulator. This flag (assuming the simulator has one) may be accessed `$simparam` from §5.2 to turn on trace modes or print special warnings (notices) only for diagnosis mode.

The `$simparam` function should have an argument that requests the iteration number.

**Proposal:** standardize Cadence’s `$debug` function; specify that `$simparam`(“iteration”) should return the iteration number.

---

## 6.0 Convenience Items

### 6.1 New format specifier for engineering notation

When printing warning, error, or diagnostic messages, designers find it easier to read numbers in engineering notation rather than exponential. For example, a capacitance might be reported as “15fF” rather than “1.5e-14F.” We propose the addition of a new “percent” code for use in `display_tasks` such as `$strobe`.

**Proposal:** add a new format specification for real numbers, `%r`, for displaying numbers in engineering notation using the scale factors listed in the LRM (T,G,M,K,k,m,u,n,p,f,a).

### 6.2 Variable initialization

To bring Verilog-A in line with other modern programming languages, it should be allowed to initialize variables where they are declared. In fact, this capability is present in Verilog 2001. We should be clear that the inability to do so is not an impediment to creating device models, though it is an inconvenience and be reminiscent of older languages like FORTRAN.

Determining a domain for variables declared and initialized at top level should not be a problem. Top-level variables have their domain determined by whether they are assigned a value in a digital or analog block. This method can still be used, even if the variable also has an initialization value, because it still can only have a value assigned in at most one domain. If it has no assignment in either domain, then the compiler could either treat it as a digital variable, or perhaps even as a numerical constant for further optimization. Another possibility mentioned in the Verilog-AMS committee meeting would be to prefix analog variable declarations with the keyword **analog**, which would also allow a user to force the domain of a variable.

**Proposal:** allow variables to be initialized where they are declared.

Since this is not a critical issue for device modeling, it can be postponed until Verilog-AMS is updated to merge with IEEE 1364-2001 or 2003 Verilog, or with SystemVerilog.

### 6.3 Declare variables where used

The issue of variable declaration is another convenience issue not specific to compact modeling.

In order to make models modular, it would be nice to declare the variables close to the code where they are used. Primarily, this is a coding style issue, but if the variables are declared inside of blocks and thus quickly go out of scope, it is possible to dramatically reduce the amount of stack space required for the model. It also allows effects to be easily ‘ifdef-ed out, copied and pasted, etc.

Named blocks already satisfy much of this requirement. For example, one could write

```
begin : avalanche
  real a, b;
  ...
end
```

and the variables  $a$  and  $b$  would be local to the avalanche block. We also understand that SystemVerilog is moving towards the C++ style of allowing variable declarations to be placed anywhere in the code.

**Proposal:** copy the SystemVerilog syntax that allows variables to be declared anywhere.

Since this is not a critical issue for device modeling, it can be postponed until Verilog-AMS is updated to merge with SystemVerilog.

---

## 7.0 Concepts Not Requiring Extensions

The following items were discussed by the subcommittee, but the consensus was that no extensions were appropriate. Either the simulator should automatically do the right thing, or the required changes to the language were too dramatic or cumbersome.

### 7.1 Required parameters

There was some interest in removing the requirement that a parameter declaration include a default value. If no value is specified for the module, then the variable would be initialized to *unknown* and the simulator should complain about a missing required parameter if that value is ever used.

However, it would be unfortunate if a simulation ran for a long time before a device entered a new region of operation where the *unknown* parameter was used and caused the simulation to abort. One could check all parameters at the start of a simulation, but a parameter might only be required contingent on other parameters (e.g., an avalanche coefficient needed only when the flag requesting avalanche modeling is on).

Since we are proposing to allow the detection of whether a parameter is specified (§1.2), the model writer can determine whether specifying the parameter is required (possibly based on values of other parameters) and then explicitly check if the parameter is specified and print an error message if it is not. Although this violates the Verilog-A design approach of providing powerful defaults and range limits as a way of avoiding the burden writing code to ‘manage’ the parameters, this work-around is sufficient to handle the examples the subcommittee is aware of.

### 7.2 Infinity as a valid parameter value

In OVI-1.0 *inf* was allowed valid value for real numbers, however in OVI-2.0 it is only allowed in the ranges, (e.g.,  $bv = 1e6$  from  $(0:inf)$ ). This is problematic because many parameters in compact models default to infinity. The Early voltage parameter of the BJT (*vaf*) is an example.

Since we are proposing the ability to detect whether a parameter was specified (§1.2), the model could be written to behave as though an unspecified parameter had the value of *inf*. In most cases, compact models will not actually use the value of *inf* in a calculation, but rather use the value as a flag to skip a certain block of equations. However, when listing the parameter values, the simulator will show the default value instead of infinity; if the user then specifies that default value, the results will be different.

In some cases, it might be possible to use 0 to mean infinity. This could be confusing for the user, but many models/simulators already use this work-around. Many simulators do not allow *inf* for parameter values and it would be impossible to specify that value in a netlist. Even if we allowed *inf* in Verilog-A, it might not be possible to use it in all simulators.

It might also be possible to use MAXFLOAT or MAXDOUBLE constants for IEEE 754 floating-point values. MAXFLOAT=3.40282346638528860e+38 and MAXDOUBLE=1.79769313486231570e+308.

Unfortunately, MAXFLOAT is not large enough to stand in for *inf* in all circumstances, and any operations on MAXDOUBLE could generate a floating-point overflow exception. One would also want the simulator to handle these values specially when printing parameter lists.

If this issue is ever revisited, it should be noted that the IEEE definition of *inf* uses a bit pattern that propagates. Thus, the simulator does not need to check for infinite values at each step of a calculation. The simulator should only check -- and complain -- if the value *inf* were ever contributed to a branch. Since what constitutes a “large” number depends on the situation, a general-purpose modeling language really ought to have the original capabilities of *inf*.

### 7.3 Partitioning of code based on phase of simulation

In compact models there is a considerable amount of code that depends only on the input parameters and perhaps temperature. As such, it only need run when the model is initialized. Exploiting this can dramatically speed up a simulation. There is sometimes code that depends on time but not on the signal values. This code should be run at the beginning of each timestep. Finally, there is code that depends on the signal values, but that do not affect the signal values produced by the module (e.g., operating point parameters). In this case, it is not necessary to execute the code on every iteration; once after convergence is sufficient.

Although model writers would be able to optimize their modules in the short term by placing properties on begin/end blocks, this solution is not very sophisticated and places the burden on the model writer to hand-optimize the code. We believe it would be much better to expect the simulator to form a dependency tree and use that to partition the code automatically.

### 7.4 More flexible noise specifications

A posting at The Designer’s Guide Forum asked how to implement the induced gate noise of the channel from the Philips MOS11 model, whose power spectral density is [5]

$$S_{ig} = \frac{\frac{1}{3} \cdot N_T \cdot (2 \cdot \pi \cdot f \cdot C_{ox})^2 / g_m}{1 + 0.075 \cdot (2 \cdot \pi \cdot f \cdot C_{ox} / g_m)^2} \quad (1)$$

This noise function can be specified through Laplace transforms, specifically,

```
x = white_noise(1/3*NT*Cox*Cox/gm);
l(g,s) <+ laplace_nd(x, {0, 1}, {1, sqrt(0.075)*Cox/gm});
```

Simulators may implement this inefficiently if they do not realize the optimization possible because the expression is only used in a noise contribution, but this is an implementation detail just like the dependency tree of §7.3, not something that requires an extension to the language.

Correlation of non-white noise from two-ports should be possible using the same method. Any physically-realizable noise coloring must be obtained from an H(s) filter that colors the noise as |H(s)|<sup>2</sup>. Indeed, if a simulator tries to implement the noise in the time domain, then it must construct the filter.

### 7.5 Optional internal nodes

In compact models there are times when internal nodes become unnecessary, either because they are shorted to another node, or because they are left to float. In these cases, models would run faster if these nodes were eliminated.

This can be done by using optimizations to eliminate the nodes. First one would identify whether a node were shorted to another node or terminal, or if it were completely floating. When such nodes were found, they would not be created. In the case of shorted nodes, the name of the node being eliminated is used as an alias for the node to which it is shorted.

Simulators must be responsible for checking whether a node is shorted or not, and whether that determination depends on a swept parameter.

Although model writers are perhaps accustomed to explicitly coding the short, they should not object to being relieved of that burden. Tiburon-DA's simulator is able to detect this condition automatically. Thus, no extension is necessary.

### **7.6 Initial guess for Newton's method**

Some compact models recommend an initial value for branch voltages, such as those across the junctions of a BJT or diode. These values sometimes give a better starting point for Newton's method, which is provably convergent only for starting points "close" in some sense to the solution.

Note that these initial guesses differ from nodeset values, which are already supported by the Verilog-AMS language. Nodeset values are enforced (typically through small resistors) for some number of Newton iterations, but these initial guesses are only applied once for the first iteration.

Unfortunately, these initial values often depend on some knowledge of the external circuit. For example, diode branches are often initialized at a positive bias to give a reasonable conductance; this is tremendously helpful if the diode is driven by a current source and frequently helpful in other situations because the linearized equation presented to the simulator tells it not to change the node potentials drastically to change the current. However, if the diode is being held in reverse bias by voltage sources on its terminals, then initializing the diode in forward bias is not helpful at all. It would be tedious for a model author to attempt to write code to handle all the possible cases.

Further, the efficacy of these initial guesses is limited for large circuits. Hence, the subcommittee does not recommend an extension to support this concept.

---

## **8.0 Extensions Requiring Further Research**

The following items are more complicated, and the correct solution is not apparent. We expect a second iteration through the compact modeling extensions to address these issues and others that were not brought up in the subcommittee.

### **8.1 Inheritance**

One might be interested in defining a set of models starting from a basic model and adding features. The advanced models would inherit various things from the base model, such as structure, ports, parameters, and then add further items. The idea is very much like the class inheritance of C++. Bell Labs is reported to have used this sort of inheritance for a set of transistor models called "asim." The simulator was able to switch between model levels automatically when attempting to solve the circuit; it would start with the base model and then work up to the complicated models.

The subcommittee feels that this extension would entail a great deal of effort on the part of the simulator vendors, akin to moving their whole development system from C to C++. Further, the inheritance method is not commonly used by authors of compact models.

## 8.2 Table models

Agilent is reportedly very interested in table models for devices. However, they did not communicate their needs or suggestions to the subcommittee. Cadence's Verilog-A implementation includes some support for table models, and their approach has been forwarded to the Verilog-AMS committee for consideration separate from these compact modeling extensions. We hope that these will be sufficient.

## 8.3 Frequency-domain descriptions

Many people have proposed adding frequency-domain modeling extensions to Verilog-A, but it is an endeavor that is filled with risk. In addition, while appealing in concept, it is not very useful in practice because there are very few models that can be easily described using formulas given in the frequency domain that are also not easily given in the time domain, and the time domain formulas do not entail the same degree of risk. Consider a few examples. Lumped components such as resistors, capacitors, and inductors are easily described in the frequency domain, but they are also easily described in the time domain, and time domain models are inherently more general as they can include nonlinear effects. This leaves distributed components, which are nearly impossible to specify in the time-domain using only the *ddt* and *idt* operators available in Verilog-A. However, in almost all cases, these models are also very difficult to specify using formulas in the frequency domain. It is almost always easier to describe these models in a tabular fashion in the frequency domain. So Verilog-A should provide a primitive instance that is capable of reading industry standard *s*-parameter files, but that would not require supporting frequency-domain extensions to the behavioral modeling capabilities in Verilog-A. There are a few models that are both distributed, and so not easily described with time-domain models, and for which the frequency domain formula is both simple and well known. Skin effect is perhaps the best known instance. However, it is hard to think of many others. For these cases, we have two choices. One alternative is simply to identify these cases and provide functions to implement the desired behavior (e.g., a skin effect function). Simply implementing a function that provided a transfer function of  $Y(s) = s\alpha X(s)$  where  $\alpha$  could be any number between  $-1$  and  $1$  would go a long way in addressing the need for frequency domain models.

If there were a strong desire to provide frequency domain modeling in Verilog-A, one would need to address the causality issue. For frequency domain models to be causal, their real and imaginary parts must be related in a complex and subtle way. The language must be designed in such a way that these conditions are met, otherwise the models will not be compatible with time-domain simulators.

We believe that any extensions of this type should originate from the RF extensions subcommittee.

## 8.4 Asserts

Some have requested an assert capability that would generate a simulator error, not a user error message, and allow checking/reporting of multiple assertion failures.

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## 9.0 References

- [1] Verilog-AMS Language Reference Manual, version 2.1, available from Accellera, [www.accelera.com/](http://www.accelera.com/).
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