

*f*REEDA<sup>TM</sup>

Programmer's Guide

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# Chapter 1

## fREEDA<sup>TM</sup> Architecture

### 1.1 Introduction

The architecture of fREEDA<sup>TM</sup> is based on faithful application of object oriented (OO) design practice [60, 61, 62, 63, 64, 65]<sup>1</sup>. The OO abstraction is well suited to modeling engineering systems, for example, in circuit simulation circuit elements are already viewed as discrete objects and at the same time as an integral part of a (circuit) continuum. The OO view is a unifying concept that maps extremely well onto the way humans perceive the world around them. Non-OO circuit simulators always become complicated with many layers of special cases. Referring to circuit elements again, traditional simulation implementations have many “if-then” like statements and individually identify every element in many places for special handling.

There are a few key premises that drove the architecture of fREEDA<sup>TM</sup>. One of these is the adoption of a very strong OO paradigm throughout to obtain a modular design. A second key premise is the separation of the core components embodying numerical methods from the modeling and solver formulation process with the result that numerical techniques developed by computer scientists and mathematicians can be formulated using formal correctness procedures. Thus, what is adopted here, is that the circuit abstraction is adapted so that highly reliable and efficient pre-developed libraries can be used.

C++ is the core implementation language and was once considered slow for scientific applications. Advances in compilers and programming techniques, however, have made this language attractive and in some benchmarks C++ outperforms Fortran [66, 67]. Several OO numerical libraries have been developed [68]. Of great importance to the work described here is the incorporation of the standard template library (STL) [69]. The STL is a C++ library of container classes, algorithms, and iterators; it provides many of the basic algorithms and data structures of computer science. The STL is a generic library, meaning that its components are heavily parameterized: almost every component in the STL is a template. The current ISO/ANSI C++ standard [70] has not been fully implemented and C++ compilers support a variable subset of the standard. The biggest areas of noncompliance being the templates and the standard library.

In this chapter the focus is on the design of the object-oriented structure of fREEDA<sup>TM</sup>. The goal in the design of fREEDA<sup>TM</sup> was to obtain speed in development, to use ‘off-the-shelf’ advanced numerical techniques, and to allow easy expansion and implementation of new models and numerical methods.

The design intent was to combine the advantages of previous OO circuit simulators with these new developments as well as expanding capability. fREEDA<sup>TM</sup> uses C++ libraries [71, 72] and several written in C or Fortran [51, 73, 58].

OO-design embodies a body of concepts that should be understood before a full appreciation of what follows can be made. This fits in beautifully with the idea of code reuse as here we utilize concept reuse. The first concept required is an understanding of the Unified Markup Language

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<sup>1</sup>These references are available on line at <http://www.objectmentor.com/>

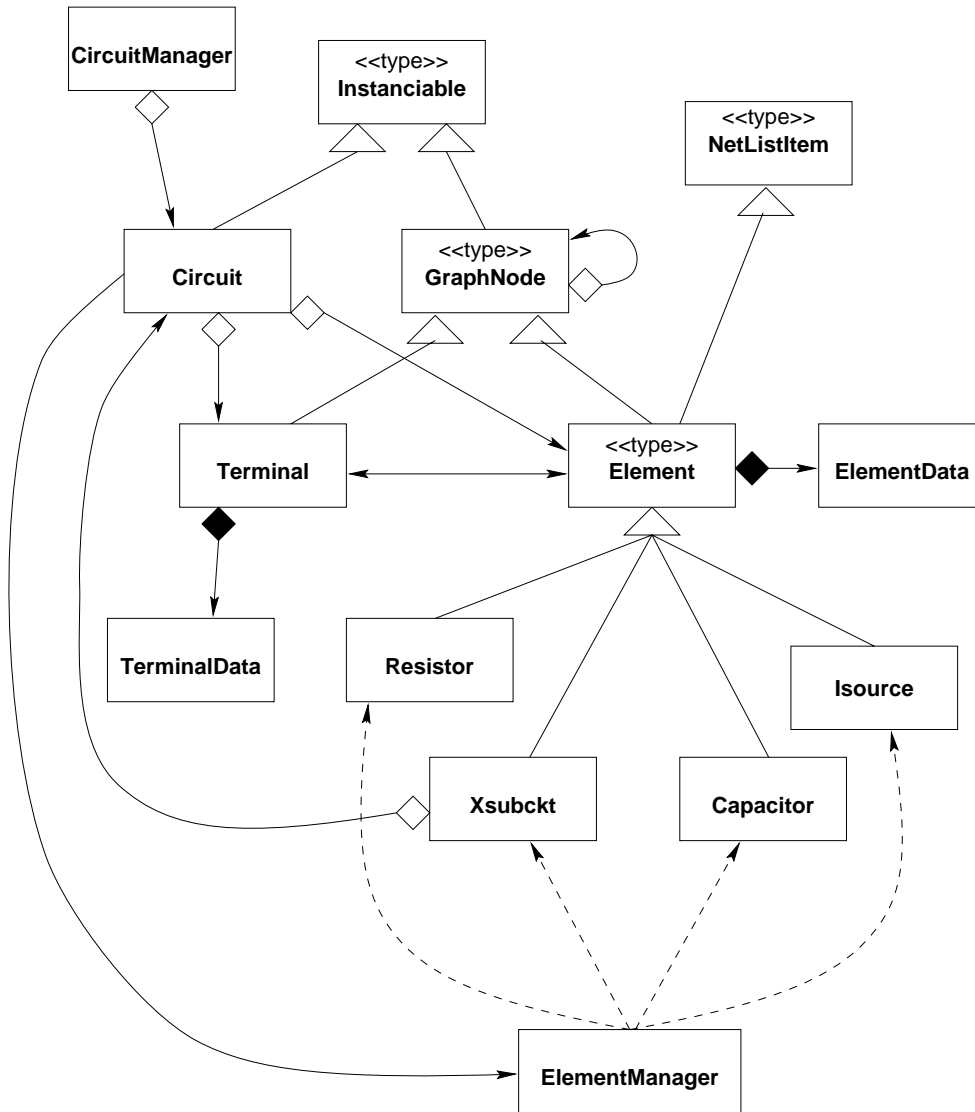


Figure 1.1: Class diagram: The network package is the core of the simulator.

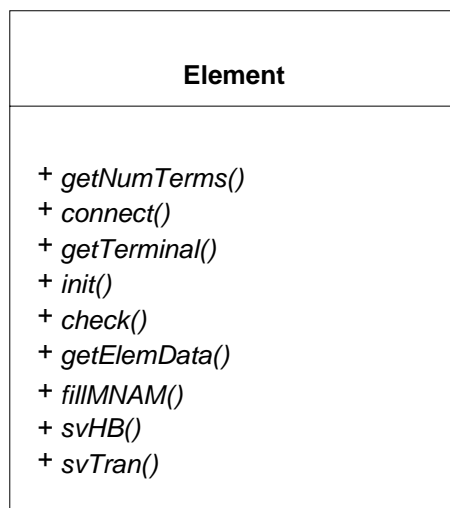
(UML). This is a universal way of presenting relationships among objects and is not specific to computer coding. UML can be viewed as a substitute for flowcharts. A flowchart permits a very limited set of relationships to be described and results in what is called spaghetti code: code that has many linear streams and complex connections between the threads. Common code is multiplied and perhaps slightly changed in each instance as the concept of using existing code and overriding only the differences is not supported. UML is different and fosters code reuse and much simpler coding. So if you are not familiar with UML study Appendix A before proceeding.

## 1.2 The Network Package

The network package is the core of the simulator. All the elements and the analysis classes are built upon it as shown in the class diagram of Figure 1.1. R(resistor), D(diode) and MesfetM shown here are examples of particular elements. Xsubckt is the SPICE X subcircuit element. The terminals of an element are specifically referred to as terminals and not as nodes as used in SPICE as the concept of a node is more general than that of a terminal. For example elements and terminals are both graph

Figure 1.2: The **NetListItem** class.

nodes. (See Appendix A for a description of the class diagram syntax.) Following the suggestion made in [8], there is a **NetListItem** class that is the base for all classes of objects that appear in the input netlist. This is the base class that handles parameters. Figure 1.2 shows some of the methods provided by this class. All the netlist items share a common syntax so that the element model developer does not need to worry about the details of element parsing and there is no need to modify the parser to add new elements. For compatibility reasons, Spice-type syntax (which does not have a consistent grammar) is supported by the parser outside the network package. Support for the elements using SPICE syntax is implemented for each element in the parser where each SPICE element is renamed as a *f*REEDA<sup>TM</sup> element.

Figure 1.3: The **Element** class.

The **Element** class contains basic methods common to all elements as well as the interface methods for the evaluation routines. Some of the methods of this class (Figure 1.3) need to be overridden by the derived classes. For example, in class **D**, *svTran()* is intended to contain the code to evaluate the time domain response of a diode. This function is used by DC and transient analyses. The same happens with *svHB()* and *fillMNAM()*. The overhead imposed by these virtual functions is

small compared to the time spent evaluating the functions themselves and so this approach is a good compromise between flexibility and efficiency. This idea has been used in [7, 8, 9]. *f*REEDA<sup>TM</sup> also offers a more elaborate mechanism for nonlinear element evaluation functions which will be described in Section 1.4.

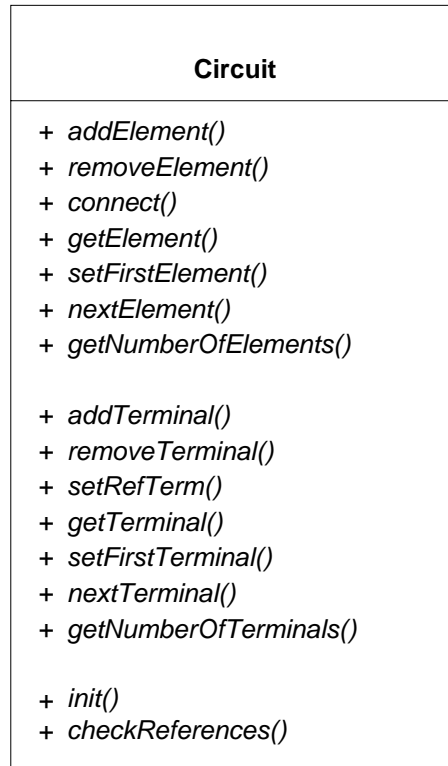


Figure 1.4: The **Circuit** class.

The **ElementManager** class is mainly responsible for keeping a catalog of all the existing elements. Note that this class is the only one that ‘knows’ about each and every type of element but this dependency is weak. The element list is included from an automatically generated file. **ElementManager** is also used to automatically generate documentation for the element in html format. The **Circuit** class represents either a main circuit or a subcircuit as a collection of elements and terminals. It provides methods to add, remove and find elements and terminals using different criteria and it also provides methods related to circuit topology. More details about this class are given in Figure 1.4. All the **Element** and **Terminal** instances must be stored in data structure inside the **Circuit** instance and the *map* container of the STL [69]. The map is a *Sorted Associative Container* that associates objects of type *Key* with objects of type *Data*. Here *Data* is either **Element** or **Terminal** and *Key* is *int* (the ID number). This is an example of where the features of C++ are used to reduce development time. This is achieved at no overhead as an optimum implementation of these concepts is embedded in the compiler. Subcircuit instances are represented by the **Xsubckt** class, see Figure 1.5. The method **attachDefinition()** is used to associate a **Circuit** instance where the actual subcircuit is stored to the particular **Xsubckt** instance and **expandToCircuit()** takes a **Circuit** pointer as argument and expands the subcircuit. Note that before expansion the complete hierarchy of a circuit is available in memory, so this engine could eventually be used to perform hierarchical simulation. The reason for this is that a subcircuit can be invoked (by an **X** element) before the subcircuit is defined (in a .SUBCKT block).



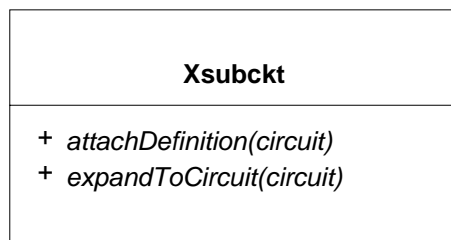


Figure 1.5: The **Xsubckt** class: This class handles the SPICE **X** element.

### 1.3 The Analysis Classes

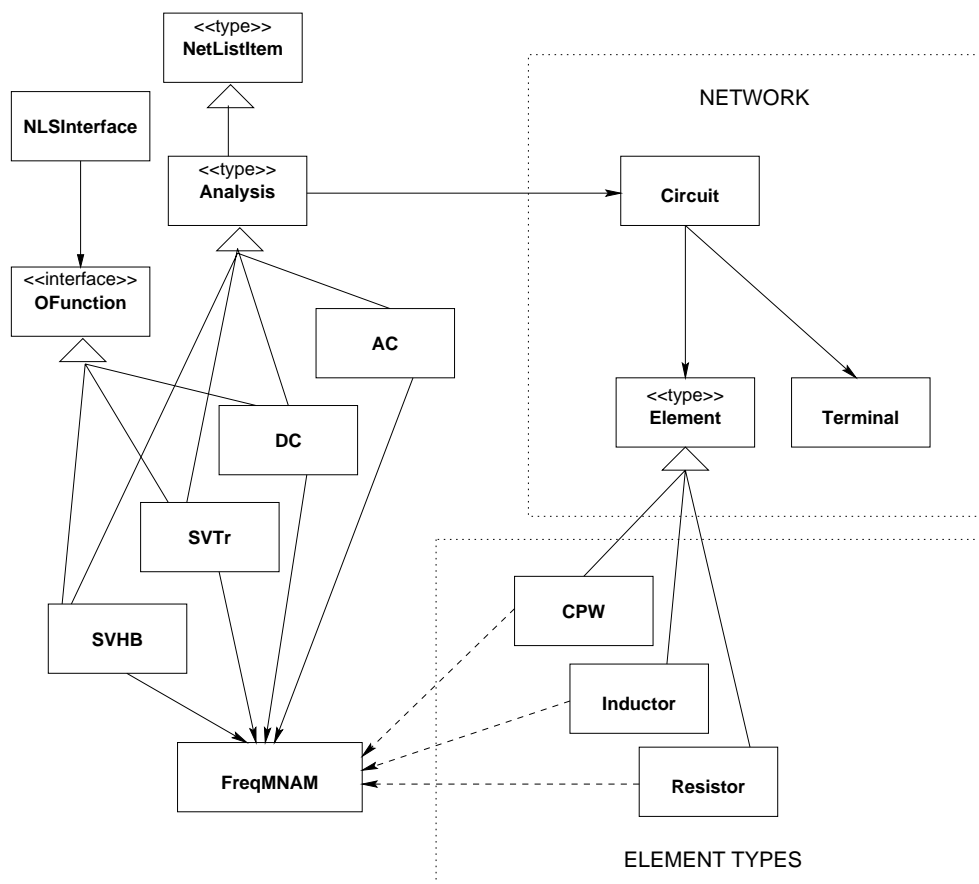


Figure 1.6: The analysis classes: **AC**, **DC**, **SVTr** and **SVHB** are different analysis types.

Figure 1.6 shows the relation between the network package, the elements and the analysis classes. Each of these classes stores analysis-specific data that would traditionally be global. This leads to the key desired attribute of flexibility in incorporating a new type of analysis, or even different implementations of the same analysis type. Examples are the **SVTr** and **SVHB** classes which contain the state-variable convolution transient and state-variable harmonic balance analyses described in [105].

There are some components common to two or more analysis types. The natural way of handling these components is by creating a class which is shared by the different analysis types. For example, the **FreqMNAM** class handles a Modified Nodal Admittance Matrix (MNAM) in the frequency domain. In a microwave simulator, the frequency domain admittance matrix is a key element for

most analysis types. Since it is used so often, special care was taken to optimize efficiency. The elements fill the matrix directly without the need for intermediate storage of the element stamp. They do that by means of in-line functions to reduce function call overhead. Elements can fill the source vector in a similar way. The elements depend on the **FreqMNAM** methods, but this is not a problem since the interface is very unlikely to change. The current implementation of MNAM uses the Sparse library, described in the next Section, and is completely encapsulated inside the **FreqMNAM** class. In the same way, the **NLSInterface** class encapsulates the nonlinear solver routines. Therefore it is possible to replace the underlying libraries, if that is desired, without the need for any code modification outside the wrapper classes. A final observation is that it is possible to add any kind of analysis type provided that the appropriate interface is defined and the member functions are written for each element type.

## 1.4 Nonlinear Elements

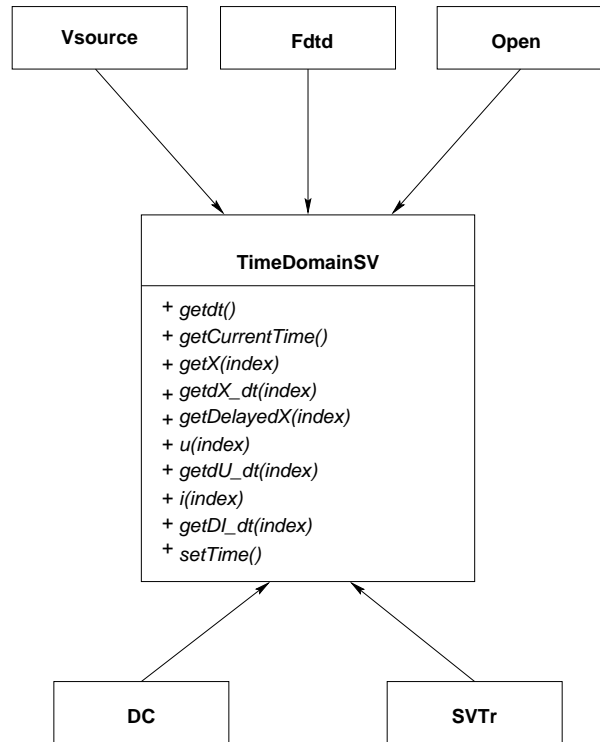


Figure 1.7: Dependency inversion was used to make the elements independent of the analysis classes.

Nonlinear elements often use service routines provided by the analysis classes. In order to maximize code reuse and to avoid the dependence of the element code on a particular analysis routines, in *fREEDA*™ elements depend on interface classes (Figure 1.7). The concept is similar to the *dependency inversion* [61]. This is a technique which relies on interface classes (normally implemented using abstract classes in C++) to make different parts of a program independent of each other. They only depend on the interfaces. To achieve greater efficiency, in *fREEDA*™ the dependency inversion is implemented using a concrete class with heavy in-lining and pass-by-reference.

In this way, the element routines and the analysis depend on an interface class, **TimeDomainSV** (not shown in Figure 1.1 for clarity). **TimeDomainSV** is a class that is used to exchange information between an element and a state variable based time domain analysis. It also provides some basic algorithms such as time differentiation methods. This approach enables the element routines to be reused by several analysis types without the need to modify the element code (as long as the

new analysis is state variable-based). For example, the **DC** analysis uses the same interface element as the **SVTr**.

*fREEDA*<sup>TM</sup> offers a more refined way to implement nonlinear elements, based on state variables (See chapter 3). By implementing those parametric equations using a special syntax in only one function, *fREEDA*<sup>TM</sup> can obtain the analysis functions **svHB()**, **svTran()** and derivatives automatically. This mechanism is termed *generic evaluation*.

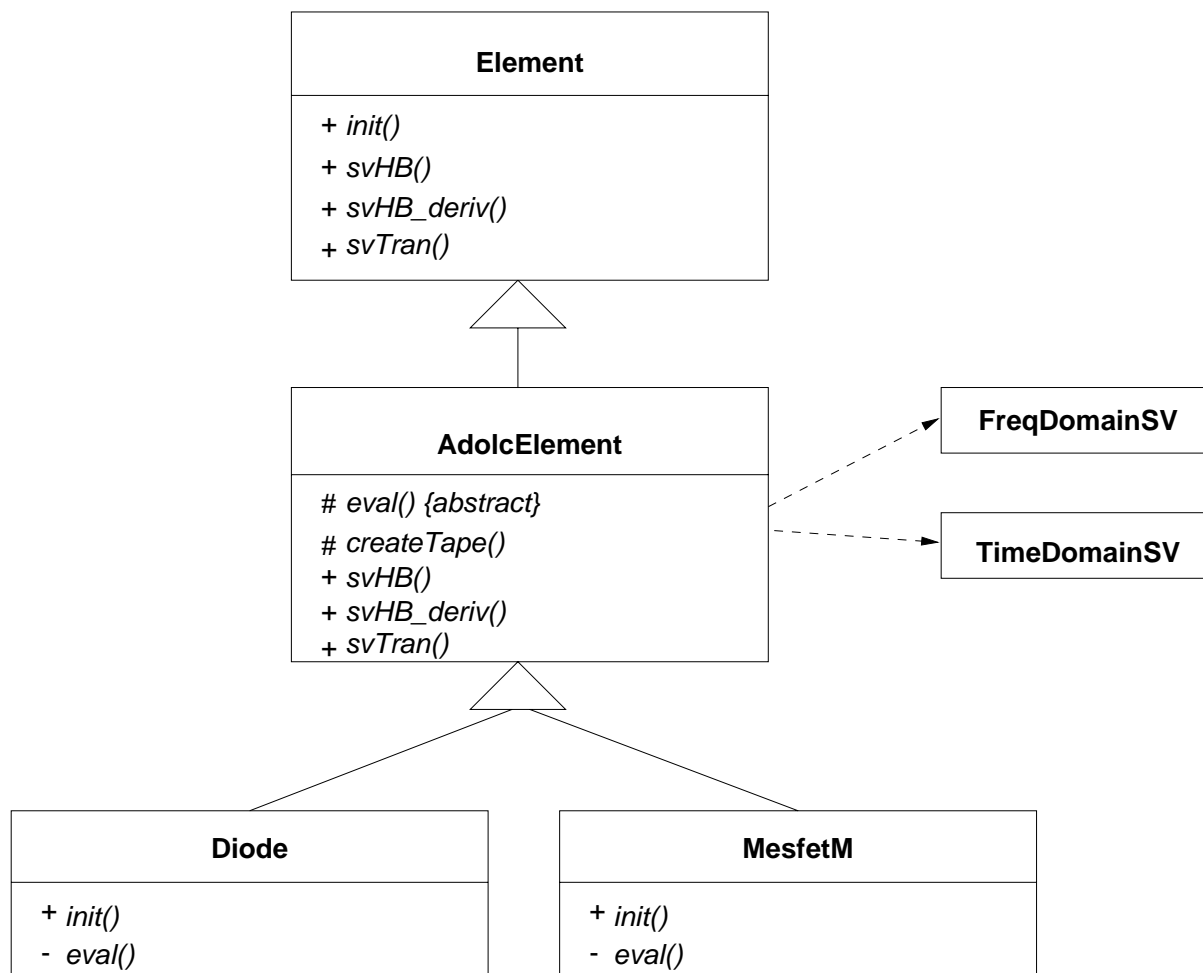


Figure 1.8: Class diagram for an element using generic evaluation. **D**(diode) and **MesfetM** are elements.

Figure 1.8 shows the class diagram for an element using this feature. Note that the **Diode** class is derived from a class (**AdolcElement**) which itself provides the analysis routines and deals with the analysis interfaces. The **Diode** class only needs to implement the `eval()` function with the parametric equations.

**AdolcElement** uses the Adol-C library (see Section B.5 for a detailed explanation) to evaluate the parametric function and derivatives, but the concept is independent of automatic differentiation. If automatic differentiation were not used, then the derived class (*e.g.* the **Diode** class) would have to provide the Jacobian for the parametric equations.

The idea is in a way similar to the one used in [6], *i.e.* the primitive equations are ‘wrapped’ in analysis-specific generic functions and so there is no need to write a separate routine for each analysis type. In the current work there are two additional features. The first is that the generic evaluation is combined with the state variable concept of Section ?? and automatic differentiation. This provides unprecedented simplicity in creating nonlinear element models. The second is that

a single mechanism is not mandatory. There are cases where it may not be practical to use this approach, or the overhead involved (which is completely acceptable even for simple models such as a diode) may not be properly amortized. In those cases, the element can implement the analysis functions directly, without using the **AdolcElement** class.

It is important to remark that generic evaluation is implemented efficiently so there are no superfluous calculations. The current implementation supports elements with any number of state variables. Each element selects the input variables as a subset of the following: the state variables, the first derivatives, the second derivatives and a time delayed version of the variables (the delay may be different for each). No derivation, time delaying nor transformation is performed on the unselected inputs.

For example, an element with only an algebraic nonlinearity (such as the VCT: voltage controlled transducer) only selects the state variables without any derivatives or delays. As another example, the **MesfetM** class implements the Materka-Kacprzac model for a MESFET. It requires two state variables, but only one of them needs to be delayed.

A consequence of having a library of elements using generic evaluation is that it is possible to add a new analysis type by just adding the appropriate evaluation routines to the **AdolcElement** class. Thus, the maintenance and expansion of the simulator is simplified. Put another way the code for a circuit element need only be implemented once and the identical code can be used by many different types of analyses including, hopefully any future analysis type implemented in the future.

## 1.5 Example: Use of Polymorphism

The concept of polymorphism is briefly explained in the introduction to object oriented programming in Appendix A. The scheme presented in Section 1.4 constitutes a good example of the use of polymorphism to solve a complex problem. Consider the segment of code of Figure 1.9. This code evaluates the nonlinear element functions in state variable convolution transient analysis. **elem\_vec** is a vector of **Element** pointers implemented using the vector container of the C++ STL. There

```
// Number of elements
int n_elem = elem_vec.size();
int i = 0;
// Go through all the nonlinear elements
for (int k = 0; k < n_elem; k++) {
    // Set base index in interface object
    tdsv->setIBase(i);
    // nonlinear element evaluation
    elem_vec[k]->svTran(tdsv);
    i += elem_vec[k]->getNumberOfStates();
}
```

Figure 1.9: Nonlinear element function evaluation in convolution transient.

is no need to keep the size of the vector in a separate variable as **elem\_vec.size()** returns the size of the vector. Also, the memory management of the vector is dynamic and automatic; and **elem\_vec[k]** returns the **Element** pointer at position *k* in the vector.

Each pointer inside **elem\_vec** points to different kinds of elements. For the transient routine the actual type of each element does not matter. The line containing **elem\_vec[k]->svTran(tdsv)** will call the appropriate evaluation routine depending on the actual type of **Element** pointer. The element may implement the routine directly or through generic evaluation, but it makes no difference for the analysis routine.

The resulting code is therefore simple and there is no need for lists of “if-then” statements. Those lists would be very difficult to maintain, because each time an element is added or removed, all the

lists would have to be updated.

## 1.6 Documentation

*f*REEDA<sup>TM</sup> uses LaTeX as the main documentation language. Documentation is distributed in the directory *freeda-version/doc*. Element (in *.../doc/fr\_elements.html*) and Analysis (in *.../doc/fr\_analysis.html*) as well as the entire contents of the *.../doc/elements* directory are automatically generated. Elements are handled as special way, as various implementations of *f*REEDA<sup>TM</sup> can have different model sets (other than the standard distribution to include proprietary models for example) *f*REEDA<sup>TM</sup> which generate html documentation of individual elements. This documentation is generated from the internal data structures so it is known to be right. The convention is to also to provide PDF documentation of individual elements which are in the directory structure for the element. Environment Variables are discussed in the User's Guide in the subsection "Setting up the Cygwin Environment." To create a set of analysis and element documentation ready for web use you should set the environment variables as follows

```
$ENV_FREEDA_DOCUMENTATION    .../freeda-version/doc
$ENV_FREEDA_WEB_DOCUMENTATION .../freeda-version/doc
```

where *freeda-version* is either just */myhome/freeda/freeda* (if soft linking has been used) or something like */myhome/freeda/freeda/freeda-1.3* and *myhome* is the user name. Then 'run *freeda -c*' and 'run *freeda -a*'. This will put *fr\_eleemnst.html* and *fr\_analysis.html* in *.../freeda-version/doc* and all of the element pdf files from the tree *\$ENV\_FREEDA\_SIMULATOR/elements* tree in *.../freeda-version/doc/elements*.

### 1.6.1 Automatically Generated Documentation

*f*REEDA<sup>TM</sup> supports automatic generation of documentation for elements and analyses. These are derived from the internal data structures and is likely to be more up-to-date than separately generated documentation. The command lines to generate documentation are

- *freeda -c*
- *freeda -c elementName*
- *freeda -a*

or, if using the cygwin environment:

- *freeda.exe -c*
- *freeda.exe -c elementName*
- *freeda.exe -a*

#### Analysis Catalog, *freeda -a*

This generates the analysis catalog and puts it in the file *fr\_analysis.html* in the directory *\$ENV\_FREEDA\_DOCUMENTATION*.

#### Element Catalog, *freeda -c*

This generates a catalog of all elements and puts it in the file *fr\_elements.html* in the directory *\$ENV\_FREEDA\_DOCUMENTATION* (this is the environment variable). Individual html files of each element are also generated and put in the directory *\$ENV\_FREEDA\_DOCUMENTATION/elements*. The individual element html files will include a link to a more detailed pdf file for the element. This must be generated by the developer who created the element model. There are two ways of handling the links to the pdf files.

If the environment variable `$ENV_FREEDA_WEB_DOCUMENTATION` points to a local filesystem then the element pdf's are copied from the `.../simulator/elements` tree and put in the directory `$ENV_FREEDA_WEB_DOCUMENTATION/elements` and the individual element html's point to the local pdfs.

If the environment variable `$ENV_FREEDA_WEB_DOCUMENTATION` does not point to a local filesystem (e.g. `http://www.freedda.org/doc` ) then the individual element html file `xxxx.html` internally points to `$ENV_FREEDA_WEB_DOCUMENTATION/elements/xxxx.pdf`. Typically this would be `http://www.freedda.org/doc/elements/xxxx.pdf` .

Environment Variables are discussed in the User's Guide.

### Individual Element Documentation, `freeda -c ElementName`

This generates an individual html file for the element with the name *ElementName*. This is put in the directory `$ENV_FREEDA_DOCUMENTATION/elements`. The html file will include a link to a more detailed pdf file for the element. This must be generated by the developer who created the element model. There are two ways of handling the links to the pdf files.

If the environment variable `$ENV_FREEDA_WEB_DOCUMENTATION` points to a local filesystem then the element's pdf's points to the pdf directly in the same directory, that is in `$ENV_FREEDA_DOCUMENTATION/elements`.

If the environment variable `$ENV_FREEDA_WEB_DOCUMENTATION` does not point to a local filesystem then the individual element html file `xxxx.html` internally points to `$ENV_FREEDA_WEB_DOCUMENTATION/elements/xxxx.pdf`.

## 1.6.2 Enhanced Element Documentation

The static information for an element is contained in the following data structure in the file `NetListItem.h` :

```
// Structure to hold static information about the item.
struct ItemInfo
{
    char* name; // Name of netlist item
    char* description; // Expanded name of netlist item
    char* author; // Acknowledge the author
    char* documentation; // documentation string
    char* version; // include version number yyyy_mm_dd
};
```

Enhanced element documentation is contained in the `documentation` variable. This is not currently used (as of *fREEDA*<sup>TM</sup> version 1.3) but will be included at some time. New elements should set this string. The syntax is a comma and semicolon delimited string:

`type1:arg1,arg2;type2:arg1`

The string is terminated in either a semicolon or a null.

An example for a microstrip line is

`category:microstrip,transmission line`

The intent is for `category:` to be used to sort documentation into categories. The developer can use any argument in category. Other fields (e.g. `tyupe2` ) are for future expansion but particularly for enhancing documentation.

## 1.7 Contributors

The following contributed to this chapter

Carlos Christoffersen

Michael Steer.

## Chapter 2

# Adding Linear Elements to **fREEDA**<sup>TM</sup>

### 2.1 The **fREEDA**<sup>TM</sup> Circuit Simulator

**fREEDA**<sup>TM</sup> is a netlist-based circuit simulator. The input format of the netlist file is similar to the SPICE format with extensions for new device models, variables, sweeps, and repetitive simulation. The program provides a variety of output data and plots. The addition of new circuit element models and analysis types in **fREEDA**<sup>TM</sup> is much simpler than in other circuit simulators such as Spice. For example, new element models are coded and incorporated into the program without modification to the high-level simulator. The circuit analysis types currently available in **fREEDA**<sup>TM</sup> are DC, AC, harmonic balance (HB) [2], convolution transient [102], wavelet transient [103], and time-marching transient [105]. Some insight into the program architecture is given in [104].

This tutorial describes the addition of linear device models to **fREEDA**<sup>TM</sup><sup>1</sup>. We assume the reader is familiar with C++ syntax and basic concepts of object-oriented programming. Some issues such as the creation of an element with multiple reference nodes are not yet described.

In the majority of cases the code for a new element can be written by following the code for an existing similar element. For example, to write a new MOS transistor mode the code for an existing MOS model can be followed.

### 2.2 Example: Adding a Linear Resistor

We illustrate the addition of linear device models using a step-by-step example with a simple model: a linear resistor.

#### 2.2.1 Netlist syntax

A brief description of the netlist will help in understanding the rest of this section.

The standard **fREEDA**<sup>TM</sup> netlist syntax is common to all elements. This differs from the SPICE syntax but this is also supported but is less general in that the grammar of the standard **fREEDA**<sup>TM</sup> netlist syntax is common to all elements while the grammar of the SPICE syntax is not consistent and each SPICE element must be handled separately in the parser.

The standard **fREEDA**<sup>TM</sup> netlist syntax is

`<netlist Name>:(<element ID> <terminal1>...<terminaln> [<parameter> = <value>]...]`

The **fREEDA**<sup>TM</sup> netlist syntax for a resistor (the **r** element) is, for example, **r:r1 n1 n2 r=100**.

---

<sup>1</sup>Further details may be found at <http://www.freeda.org>

**r:1** is the **elementID**, **n1** and **n2** are the names of two terminals and **r** is a parameter syntax name.

SPICE syntax is also supported for the original SPICE elements. In SPICE syntax the equivalent input is **r1 n1 n2 100**.

### 2.2.2 Class Name, Required Files, etc.

A good class name for the element is **Resistor**. By convention (in *fREEDA*<sup>TM</sup>), class names should begin with capital letters and contain no underscores. We need a netlist name for our element. Assume we call it “r”.

The files containing the model description are located in the **elements/** directory. There is a header file (**Resistor.h**) containing the declaration of the class that defines the element and a file containing the definition of the class with the actual model (**Resistor.cc**). Those are the only files needed to define this element.

### 2.2.3 The Header File

The header file starts with comment lines describing the element type and sometimes a simple ASCII drawing of the element schematic. The figure can be used to describe terminal numbering.

```
// This may look like C code, but it is really -*- C++ -*-
//
// This is an resistor model
//
//          res
//      o-----/\ /\ \-----o
//
// by Carlos E. Christoffersen
```

Header files may be included more than once in C++ programs. To avoid multiple declarations of the classes defined in the body of the header file, the definitions in the header file are enclosed by following preprocessor directives:

```
#ifndef Resistor_h
#define Resistor_h
class Resistor : public
Element {
    :
}
#endif
```

The **Resistor** class is derived from the base class **Element**. This is so for all elements in *fREEDA*<sup>TM</sup>. For simple elements, the only public functions that must be declared are the following:

```
class Resistor : public Element
{
public:

    Resistor(const string& iname);

    ~Resistor() {}

    static const char* getNetlistName()
    {
        return einfo.name;
    }
}
```



```
// fill MNAM
virtual void fillMNAM(FreqMNAM* mnam);
```

```
virtual void fillMNAM(TimeMNAM* mnam);
```

The constructor always takes the same argument, `iname`. This is the name of the particular instance of the resistor being created (e.g. “r1”).

The destructor in this case is trivial. The next function, `getNetlistName()` must be defined in every `fREEDATM` element. It returns the netlist name of the element (“r” in our example) and it is used during the netlist parsing. It is declared static because this function is called before the actual element instance is created.

The last two functions are the declarations of the functions to fill the entries corresponding to this element in the *modified nodal admittance matrix* (MNAM) of the circuit being simulated.

The private members of this class are:

```
private:
```

```
// Parameter variables
double r;

// Element information
static ItemInfo einfo;

// Number of parameters of this element
static const unsigned n_par;

// Parameter information
static ParmInfo pinfo[];

};
```

The netlist parameters of this element are declared as normal member variables. Here we have only one parameter called “r”. There is no conflict with the netlist name of this element. The last three variables can be left unchanged for any other element in `fREEDATM`. We will describe their use later.

### 2.2.4 The Class Source File

The include preprocessor directives go first. The `ElementManager.h` file must be always included first. For linear elements, we also need to include the declarations for the MNAM classes. There are currently two implementations of the MNAM in `fREEDATM`. One in the frequency domain (`FreqMNAM`) and one in the time domain (`TimeMNAM`). The last included file is the declaration of the class for our element, `Resistor.h`.

```
#include "../network/ElementManager.h"
#include "../analysis/FreqMNAM.h"
#include "../analysis/TimeMNAM.h"
#include "Resistor.h"
```

Now we must define the static member variables:

```
// Static members
const unsigned Resistor::n_par = 1;

// Element information
ItemInfo Resistor::einfo = {
    "r",
    "Resistor",
    "Carlos E. Christoffersen",
    ""
};
```

```
// Parameter information
ParmInfo Resistor::pinfo[] = {
    {"r", "Resistance value (Ohms)", TR_DOUBLE, true}
};
```

"r" under Element information is the NetlistName. "r" under Parameter information is where the parameter name is defined.

The `n_par` variable must be equal to the number of parameters defined for the element. In our example, only one parameter is defined. Note the use of the scope (::) operator. It is used to indicate that the `n_par` variable is a member of the **Resistor** class. The `einfo` structure contains strings describing the element. The first string is the netlist name of the element ("r"). This name must always be given in lowercase letters. The second string is a more verbose description of the element. The third string should list the authors of the element code and the last string is used to store a web/ftp/e-mail address where more information about the element can be found. The idea here is give full exposure to the creator of an element and that an element is 'owned' by the person or group that creates the element.

The `pinfo` vector contains the description of each parameter defined in the element. The first field (always in lowercase) is the netlist name of the parameter followed by a more verbose description. The units of the parameter (if any) should be included in this description. The third field is a flag denoting the type of the parameter. The possible flags are defined in the `NetListItem.h` file. The corresponding types for each flag are given in Table 2.1. The last field in the parameter description is a flag that is *true* if the parameter

Flag	Corresponding type
TR_INT	int
TR_LONG	long int
TR_FLOAT	float
TR_DOUBLE	double
TR_CHAR	char
TR_STRING	string
TR_COMPLEX	double_complex
TR_BOOLEAN	bool
TR_INT_VECTOR	IntVector
TR_DOUBLE_VECTOR	DoubleVector
TR_INT_MATRIX	IntMatrix
TR_DOUBLE_MATRIX	DoubleMatrix

Table 2.1: Possible parameter types.

is required in the netlist (*i.e.* an error occurs if the parameter is not specified) and *false* otherwise.

The constructor definition follows:

```
Resistor::Resistor(const string& iname) : Element(&einfo, pinfo, n_par,
iname)
{
    // Value of r is required
    paramvalue[0] = &res;

    // Set the number of terminals
    setNumTerms(2);

    // Set flags
    setFlags(LINEAR | ONE_REF | TR_FREQ_DOMAIN);
}
```

The constructor for the **Element** class takes the static member variables we defined before and the instance name (`iname`) as arguments. In the body of the **Resistor** constructor we must point each element

of the **paramvalue** vector to the address of the member variables corresponding to the netlist parameters. The order of the elements in the **paramvalue** vector must be the same order that we used in the description in **pinfo**.

The member function **setnumTerms()**, which is derived from the **Element** class, is used to set how many terminals our element has.

The last function used in this constructor sets some flags that are useful to classify the different elements in a circuit. Table 2.2 summarizes the meaning of each possible flag.

Flag	Meaning
LINEAR	Linear in HB and tran2 analyses.
NONLINEAR	Nonlinear in HB and tran2 analyses.
ONE_REF	One internal reference node ( <i>e.g.</i> a normal element).
MULTI_REF	Two or more internal reference nodes.
TR_TIME_DOMAIN	The element is treated in time domain in convolution transient analysis. This includes nonlinear elements.
TR_FREQ_DOMAIN	The element is treated in frequency domain in convolution transient analysis. This includes linear elements.
SOURCE	The element is a source in tran2 analysis.

Table 2.2: Currently defined flags.

### 2.2.5 Filling the Modified Nodal Admittance Matrix

The last two methods in the source file are the routines used to fill the MNAM of the circuit. The first function takes a pointer to an object of the **FreqMNAM** class.

```
void Resistor::fillMNAM(FreqMNAM* mnam)
{
    // Ask my terminals the row numbers
    mnam->setAdmittance(getTerminal(0)->getRC(), getTerminal(1)->getRC(),
        one/res);
}
```

Here **setAdmittance** sets the four element stamp of an admittance **g**. Thus the stamp is

$$\begin{bmatrix} g & -g \\ -g & g \end{bmatrix} \quad (2.1)$$

The first terminal of the element is obtained using **getTerminal(0)** and the second terminal by **getTerminal(1)**. **getRC()** gets the row/column index and  $g = 1/r$  (i.e. **one/res**) is the conductance of the element.

This is a class to represent a set of modified nodal admittance matrices in the frequency domain with their respective source vectors. The following is the list of functions available to fill the matrices and the source vectors:

```
//-----
// Methods to fill the matrices used by the elements
//-----

inline void setElement(const unsigned& row, const unsigned& col,
    const double_complex& val);

inline void setAdmittance(const unsigned& term1, const unsigned& term2,
    const double_complex& val);
```

```

inline void setQuad(const unsigned& row1, const unsigned& row2,
                   const unsigned& col1, const unsigned& col2,
                   const double_complex& val);

```

```

inline void setOnes(const unsigned& pos, const unsigned& neg,
                   const unsigned& eqn);

```

```

//-----
// Methods to fill the source vectors
//-----

```

```

// Set one element of the current source vector

```

```

void setSource(const unsigned& row, const double_complex& val);

```

```

// Add value to a pair of elements of the current source vector

```

```

void addToSource(const unsigned& pos, const unsigned& neg,
                 const double_complex& val);

```

The **FreqMNAM** class also provides a set of methods to retrieve information about the matrices. One of the most used in the element routines is:

```

// Get the current frequency
inline const double& getFreq();

```

This method returns the frequency at which the the MNAM is expected to be calculated.

The last method is used to fill the elements in the time-domain MNAM.

```

void Resistor::fillMNAM(TimeMNAM* mnam)
{
    // Ask my terminals the row numbers
    mnam->setMAdmittance(getTerminal(0)->getRC(), getTerminal(1)->getRC(),
                       one/res);
}

```

For a simple element such as our resistor example, this function looks very similar to the frequency-domain case. Nevertheless, the interfaces of the **FreqMNAM** and **TimeMNAM** classes are not exactly the same. The following are the methods available to fill the MNAM in the time domain:

```

//-----
// Methods to fill the matrices used by the elements
//-----

```

```

// Methods for filling M

```

```

inline void setMElement(const unsigned& row, const unsigned& col,
                       const double& val);

```

```

inline void setMAdmittance(const unsigned& term1, const unsigned& term2,
                          const double& val);

```

```

inline void setMQuad(const unsigned& row1, const unsigned& row2,
                    const unsigned& col1, const unsigned& col2,
                    const double& val);

```

```

inline void setMOnes(const unsigned& pos, const unsigned& neg,
                    const unsigned& eqn);

```

```

// Methods for filling Mp

```

```

inline void setMpElement(const unsigned& row, const unsigned& col,
                       const double& val);

```

```

inline void setMpAdmittance(const unsigned& term1, const unsigned& term2,
                          const double& val);

```

```

inline void setMpQuad(const unsigned& row1, const unsigned& row2,
                    const unsigned& col1, const unsigned& col2,
                    const double& val);

```

```

inline void setMpOnes(const unsigned& pos, const unsigned& neg,
                    const unsigned& eqn);

//-----
// Methods to fill the source vector
//-----
// Reminder: reference source element is not stored

// Set one element of the source vector
inline void setSource(const unsigned& row, const double& val);

// Add value to a pair of elements of the source vector
inline void TimeMNAME::addToSource(const unsigned& pos, const unsigned&
neg, const double& val);

// Get the current time
inline const double& getTime();

```

Note that we must build two matrices denoted in the source code as **M** and **Mp**. These correspond to the **G** and **C** matrices in Chapter 3 of Reference [105]. Another difference with the frequency domain case is that the source vector is handled separately, as it will demonstrated in a future example.

### 2.2.6 Modifications to the rest of the **fREEDA**<sup>TM</sup> Source Files

Refer to the file **readme.txt** in the **freeda-0.1** subdirectory for instructions on compiling a new element into **fREEDA**<sup>TM</sup>.

## 2.3 Template Files for New Linear Elements

The files in this section are provided to simplify the creation of new elements. All that is required is to replace **Element1** by the name of the actual class (or name of the element), copy the files with appropriate names in the **elements/** directory and write the code to fill the MNAME.

### 2.3.1 Header File

```

// This may look like C code, but it is really -*- C++ -*-
//
// This is a generic element template
//
//      +-----+
//      o-----+ +-----o
//      +-----+
//
#ifndef Element1_h
#define Element1_h 1

class Element1 : public Element
{
public:

    Element1(const string& iname);

    ~Element1() {}

    static const char* getNetlistName()

```

```

{
    return einfo.name;
}

// fill MNAM
virtual void fillMNAM(FreqMNAM* mnam);

virtual void fillMNAM(TimeMNAM* mnam);

private:

    // Parameter variables (replace by your parameter variables here)
    double par1, par2;

    // Element information
    static ItemInfo einfo;

    // Number of parameters of this element
    static const unsigned n_par;

    // Parameter information
    static ParmInfo pinfo[];

};

#endif

```

### 2.3.2 Class Source File

```

#include "../network/ElementManager.h"
#include "../analysis/FreqMNAM.h"
#include "../analysis/TimeMNAM.h"
#include "Element1.h"

// Static members (set to the number of parameters of your element)
const unsigned Element1::n_par = ;

// Element information
ItemInfo Element1::einfo = {
    "res",
    "Element1",
    "Your Name",
    "Reserved for documentation use"
};

// Parameter information
ParmInfo Element1::pinfo[] = {
    {"par1", "Parameter 1 (Ohms)", TR_DOUBLE, true},
    {"par2", "Parameter 2 (F)", TR_DOUBLE, false}
};

```

```

Element1::Element1(const string& iname) : Element(&einfo, pinfo, n_par,
iname)
{
    // Value of r is required
    paramvalue[0] = &par1;
    paramvalue[1] = &(par2 = 1e-12); // Example of how to set a default value.

    // Set the number of terminals (Set to the number of terminals of
    // your element)
    setNumTerms(2);

    // Set flags (do not change these for normal linear elements)
    setFlags(LINEAR | ONE_REF | TR_FREQ_DOMAIN);
}

void Element1::fillMNAME(FreqMNAME* mnam)
{
    // Put your model here
}

void Element1::fillMNAME(TimeMNAME* mnam)
{
    // Put your model here
}

```

## 2.4 Contributors

The following contributed to this chapter

Nikhil Kriplani

Carlos Christoffersen

Michael Steer.





## Chapter 3

# Adding Nonlinear Elements to fREEDA<sup>TM</sup>

### 3.1 The fREEDA<sup>TM</sup> Circuit Simulator

This tutorial describes the addition of nonlinear device models to fREEDA<sup>TM</sup><sup>1</sup>. We assume the reader is familiar with C++ syntax and basic concepts about object-oriented programming.

Presently, there are two ways to write nonlinear element models, using two different base classes. One class, AdolcElement, has been extensively used and is the base class of the case study to follow. Unfortunately, a limitation recently discovered in the Adol-C library is reflected directly into a limitation on the elements that can be defined by the AdolcElement class. This class has three function prototypes for the createTape() function. One of these prototype functions is designed to allow the declaration of a second derivative state variable intended to track one of the declared state variables. **This second derivative capability has been found to be non-functional.** Usage of the function prototype is still permitted, as it is the only way to obtain delayed instances of state variables, but the argument for the second derivative IntVector to createTape() in this case should be 'novar' and this is noted herein.

The fREEDA<sup>TM</sup> team is engaged in the on-going development of a new nonlinear base class called NAdolcElement which is intended to alleviate the limitation in Adol-C and allow evaluation of second and third derivatives in models. While NAdolcElement is subject to change, model writers requiring these higher order derivatives are invited to inspect the header files and source codes for NMOSx and PMOSx elements (where x is 1,2,3, or 14) and the CompactDiode element and use them as templates for writing a model based on the NAdolcElement class. Model writers are encouraged to use NAdolcElement for writing all new elements, since over a period of time, AdolcElement will take on full legacy status and its continued use will be discouraged.

A future revision of this document will also contain a case study of an element based on the NAdolcElement base class.

### 3.2 Example: Adding a Nonlinear Electro-Thermal Resistor

We illustrate the addition of nonlinear device models using a step-by-step example with a simple model: a nonlinear electro-thermal resistor<sup>2</sup>.

---

<sup>1</sup>Further details may be found at <http://www.freedea.org>

<sup>2</sup>Note that the electro-thermal resistor behaves as a linear resistor when the parameter "pdr" is set to *false*

### 3.2.1 Netlist syntax

A brief description of the *freeda*<sup>TM</sup> netlist syntax will assist in understanding of the details of model construction. The standard *freeda*<sup>TM</sup> netlist syntax is common to all elements. This differs from the *SPICE* syntax. *SPICE* syntax is also supported but its grammar is less general than the standard *freeda*<sup>TM</sup> netlist syntax. This is because the *freeda*<sup>TM</sup> grammar is common to all elements while the grammar of the *SPICE* syntax is not consistent and each *SPICE* element must be handled separately in the netlist parser.

The standard *freeda*<sup>TM</sup> netlist syntax is

```
<netlist Name>:<element ID> <terminal1>...<terminaln> [<parameter> = <value>...]
```

For example, the *freeda*<sup>TM</sup> netlist syntax for the thermal resistor element (the **tr** element), which will be shown here for tutorial purposes, is:

```
tr:tr1 n1 n2 n3 n4 r0=100 l=1 w=1 t=35 rsh=100 pdr=true
```

**tr1** is the **elementID** and **n1** through **n4** are the names of four terminals. **r0**, **l**, **w**, **t**, etc. are parameter syntax names, and the values on the right side of the = symbol are parameter values specific to this instance of **tr:1**. Some optional parameters have been omitted in this example.

### 3.2.2 Class Name, Required Files, etc.

A good class name for the element is **Tresistor**. By convention (in *freeda*<sup>TM</sup>), class names should begin with capital letters and contain no underscores. A netlist name is needed for the element. Assume we call it "tr". *The netlist name must always be given in lowercase letters because the freeda<sup>TM</sup> netlist parser converts a netlist to lower case before attempting to identify elements or parameters.*

The files containing the model description are located in the **elements/** directory. There is a header file (**Tresistor.h**) containing the declaration of the class that defines the element and a file containing the definition of the class with the actual model (**Tresistor.cc**). Those are the only files needed to define this element. They are described next.

### 3.2.3 The Header File

The header file starts with comment lines describing the element type and sometimes a simple ASCII drawing of the element schematic as shown in Listing 3.1. The figure can be used to describe terminal numbering.

Listing 3.1: Header Comments of Tresistor.h

---

```

1
2  // This may look like C code, but it is really -*- C++ -*-
3  //
4  // This is an electro-thermal resistor model
5  //
6  //
7  //      ++++++
8  //      +   tr   +
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```

---

Header files may be included more than once in C++ programs. To avoid multiple declarations of the classes defined in the body of the header file, the definitions in the header file are enclosed by following preprocessor directives as in Listing 3.2.

Listing 3.2: Preprocessor Directives of Tresistor.h

---

```

17
18 #ifndef Tresistor_h
19 #define Tresistor_h 1
20
21 #include "../network/AdolcElement.h"
22
23
24 class Tresistor : public AdolcElement

```

---

The **Tresistor** class is derived from the base class **AdolcElement**. This is so for all nonlinear elements in *fREEDA*<sup>TM</sup> that rely upon ADOL-C for evaluating the first and higher derivatives of vector functions. For simple elements, the only public functions that must be declared are shown in Listing 3.3.

Listing 3.3: Class declaration and public functions of Tresistor.h

---

```

24
25 class Tresistor : public AdolcElement
26 { public:
27
28   Tresistor(const string& iname);
29
30   ~Tresistor() {}
31
32   static const char* getNetlistName()
33   {
34     return einfo.name;
35   }
36
37   // Do some local initialization
38   virtual void init() throw(string&);
39
40   // Get a vector with the indexes of the local reference nodes.

```

---

---

```

41 // Get terminal pointers in term_list
42 // ordered by local reference node:
43 //
44 // t0 t1 t2 t3
45 //
46 //      ^      ^
47 //   LRN1 LRN2
48 //
49 // local_ref_vec contains: {1, 3}
50 //
51 virtual void getLocalRefIdx(UnsignedVector& local_ref_vec, //
52                             TerminalVector& term_list); //
53
54 // fill MNAM
55 virtual void fillMNAM(FreqMNAM* mnam);
56
57 virtual void fillMNAM(TimeMNAM* mnam);

```

---

The constructor always takes the same argument, **iname**. This is the name of the particular instance of the thermal resistor being created (*e.g.* “tr1”). It is passed from the input netlist at the time *f*REEDA<sup>TM</sup> is run.

The destructor in this case is trivial. The next function, **getNetlistName()** must be defined in every *f*REEDA<sup>TM</sup> element. It returns the netlist name of the element (“tres” in our example) and it is used during the netlist parsing. It is declared static because this function is called before the actual element instance is created.

The function **init()** must be defined for every nonlinear element based on Adol-C, since it is here where we create the Adol-C tape. It is also here where we perform some initializations that cannot be done by the constructor, *i.e.* when some parameters are not known at construction time.

The function **getLocalRefIdx()** must be defined whenever we want to implement an element with multiple reference nodes *e.g.* spatially distributed elements or electro-thermal ones. It is important to know that this function has nothing to do with linearity. So we may have a linear element with one reference node, or with multiple reference nodes. We may also have a nonlinear element with one reference node, or with multiple reference nodes. Local reference nodes are covered in further detail in Section 3.2.4.

The last two functions are the declarations of the functions to fill the entries corresponding to this element in the *modified nodal admittance matrix* (MNAM) of the circuit being simulated. Please refer to [106] for a detailed explanation on how to implement a linear element.

The private members of this class are shown in Listing 3.4.

Listing 3.4: Private functions of Tresistor.h

---

```

58
59 private:
60
61
62 virtual void eval(adoublev& x, adoublev& vp, adoublev& ip);
63
64
65
66 // Element information
67 static ItemInfo einfo;
68
69 // Number of parameters of this element
70 static const unsigned n-par;
71
72 // Parameter variables
73 double r0, rsh, l, w, narrow, defw, t, tnom, tc1, tc2;
74 bool pdr;
75
76 // Parameter information

```

---

```

77  static ParmInfo pinfo [];
78
79  // The calculated resistor value
80  double r;
81  };
82
83 #endif

```

---

The netlist parameters of this element are declared as normal member variables. Here we have 11 parameters, and again we emphasize that *parameter names must always be given in lowercase letters because the fREEDA<sup>TM</sup> netlist parser converts a netlist to lower case before attempting to identify elements or parameters*. Please refer to reference [108] for a description of the parameters and the device equations. The variable “r” is a temporary variable used to compute the resistance value when the resistor is linear i.e. when `pdr == false`. The last three variables can be left unchanged for any other element in fREEDA<sup>TM</sup>. We will describe their use later.

The only private function that must be defined is the `eval` function. It takes 3 arguments of type `adoublev&`, the first one is an input argument, and the second and the third are output arguments. Things will be more clear when we discuss the `createTape` function.

### 3.2.4 The Class Source File

#### Preprocessor Directives

The include preprocessor directives go first; see Listing 3.5. The `ElementManager.h` file must be always included first. The next file to be included is `AdolcElement.h`. This file must be included in all nonlinear elements that are based on Adol-C. Since our element is also a linear one, we also need to include the declarations for the MNAM classes. There are currently two implementations of the MNAM in fREEDA<sup>TM</sup>. One is the frequency domain (`FreqMNAM`) and the other is the time domain (`TimeMNAM`). The last included file is the declaration of the class for our element, `Tresistor.h`.

---

Listing 3.5: Preprocessor directives of `Tresistor.cc`

---

```

1
2 #include "../network/ElementManager.h"
3 #include "../analysis/FreqMNAM.h"
4 #include "../analysis/TimeMNAM.h"
5 #include "Tresistor.h"

```

---

#### Static Variables and Model Parameters

Now we must define the static member variables as in Listing 3.6.

---

Listing 3.6: Static members and Parameter info of `Tresistor.cc`

---

```

7
8 // Static members
9 const unsigned Tresistor::n_par = 11;
10
11 // Element information
12 ItemInfo Tresistor::einfo = {
13     "tr",
14     "Tresistor",
15     "Houssam S. Kanj",
16     DEFAULT_ADDRESS"elements/Tresistor.h.html"
17 };
18
19 // Parameter information
20 ParmInfo Tresistor::pinfo [] = {
21     {"r0", "Resistance value (Ohms)", TR_DOUBLE, false},

```

```

22 {"l", "length (meters)", TR_DOUBLE, false},
23 {"w", "width (meters)", TR_DOUBLE, false},
24 {"t", "system Temperature (Celsius)", TR_DOUBLE, false},
25 {"rsh", "sheat resistance (Ohms/sq)", TR_DOUBLE, false},
26 {"defw", "default device width (meters)", TR_DOUBLE, false},
27 {"narrow", "narrowing due to side etching (meters)", TR_DOUBLE, false},
28 {"tnom", "initial Temperature (Celsius)", TR_DOUBLE, false},
29 {"tc1", "Temperature Coefficient (1/Celsius)", TR_DOUBLE, false},
30 {"tc2", "Temperature Coefficient (1/Celsius)", TR_DOUBLE, false},
31 {"pdr", "Power Deperent Resistor", TR_BOOLEAN, false}
32
33 };

```

The **n\_par** variable must be equal to the number of parameters defined in our element. In our example, 11 parameters are defined. Note the use of the scope (::) operator. It is used to indicate that the **n\_par** variable is a member of the **Tresistor** class. The **einfo** structure contains strings describing the element. The first string is the netlist name of the element ("tr"). *This name must always be given in lowercase letters because the FREEDA<sup>TM</sup> netlist parser converts a netlist to lower case before attempting to identify elements or parameters.* The second string is a more verbose description of the element. The third string should list the authors of the element code and the last string is used to store documentation information, see Section ??

The **pinfo** vector contains the description of each parameter defined in the element. The first field (*again, always in lowercase*) is the netlist name of the parameter followed by a more verbose description. We recommend to include the units of the parameter (if any) in this description. The third field is a flag denoting the type of the parameter. The possible flags are defined in the **NetListItem.h** file. The corresponding types for each flag are given in Table 3.1. The last field in the parameter description is a flag

Flag	Corresponding type
TR_INT	int
TR_LONG	long int
TR_FLOAT	float
TR_DOUBLE	double
TR_CHAR	char
TR_STRING	string
TR_COMPLEX	double_complex
TR_BOOLEAN	bool
TR_INT_VECTOR	IntVector
TR_DOUBLE_VECTOR	DoubleVector
TR_INT_MATRIX	IntMatrix
TR_DOUBLE_MATRIX	DoubleMatrix

Table 3.1: Possible parameter types.

that is *true* if the parameter is required in the netlist (*i.e.* an error occurs if the parameter is not specified) and *false* otherwise.

### Constructor for a Nonlinear Instance of Tresistor

The constructor definition for a nonlinear instance of **Tresistor** (where **pdr == true**) follows in Listing 3.7.

Listing 3.7: Constructor definition of Tresistor.cc

```

35
36 Tresistor::Tresistor(const string& iname) : AdolcElement(&einfo,
37 pinfo, n_par, iname) {
38

```

---

```

39 // Set parameters
40 paramvalue[0] = &(r0);
41 paramvalue[1] = &(l);
42 paramvalue[2] = &(w);
43 paramvalue[3] = &(t);
44 paramvalue[4] = &(rsh);
45 paramvalue[5] = &(defw = 1e-6);
46 paramvalue[6] = &(narrow = zero);
47 paramvalue[7] = &(tnom = 27);
48 paramvalue[8] = &(tc1= zero);
49 paramvalue[9] = &(tc2= zero);
50 paramvalue[10] = &(pdr=false);
51
52
53 }

```

---

The constructor for the **AdolcElement** class takes the static member variables we defined before and the instance name (**iname**) as arguments. In the body of the **Tresistor** constructor we must point each element of the **paramvalue** vector to the address of the member variables corresponding to the netlist parameters. The order of the elements in the **paramvalue** vector must be the same order that we used in the description in **pinfo**.

Then the **init()** function definition follows in Listing 3.8.

Listing 3.8: Initialization function of Tresistor.cc

---

```

54
55 void Tresistor::init() throw(string&)
56 {
57     if (!isSet(&w))
58         w = defw;
59     if (!isSet(&t))
60         t = tnom;
61
62     if (pdr){
63         // Set the number of terminals
64         setNumTerms(4);
65         // Set flags
66         setFlags(NONLINEAR | MULTLREF | TR_TIME_DOMAIN);
67         // Set number of states
68         setNumberOfStates(2);
69
70         if (!isSet(&r0)) {
71             if (!isSet(&l))
72                 throw("r0 or l must be specified for the resistance");
73             if (!isSet(&rsh))
74                 throw("rsh must be specified for the resistance");
75         }
76         // create tape
77         IntVector var(2,0);
78         var[1] = 1;
79         createTape(var);
80     }

```

---

The **init()** function starts by doing some conditional assignments. The condition test cannot be carried out in the constructor since the parameters value are not known at construction time. Then if **pdr == true**, i.e. if it is an electro-thermal resistor, we use the member function **setnumTerms()**, which is derived from the **Element** class, to set how many terminals our element has.

The next function is used to sets some flags that are useful to classify the different elements in a circuit. Table 3.2 summarizes the meaning of each possible flag. Function **setNumberOfStates()** is used to set the number of state variables used to implement the model. It is important to know that the the functions **setnumTerms()**, **setFlags()**, & **setNumberOfStates()** could be called from the constructor, but in our

Flag	Meaning
LINEAR	Linear in HB and tran2 analyses.
NONLINEAR	Nonlinear in HB and tran2 analyses.
ONE_REF	One internal reference node ( <i>e.g.</i> a normal element).
MULTI_REF	Two or more internal reference nodes.
TR_TIME_DOMAIN	The element is treated in time domain in convolution transient analysis. This includes nonlinear elements).
TR_FREQ_DOMAIN	The element is treated in frequency domain in convolution transient analysis. This includes linear elements).
SOURCE	The element is a source in tran2 analysis.

Table 3.2: Currently defined flags.

special case, we needed to put them in the `init()` function since we had to take some conditional decisions based on the parameters which are not known at construction time. Note, for example, that it is vitally important to set the `NONLINEAR` flag when an element requires the Adolc library facilities to implement a nonlinear function.

Next, the code makes some tests on the input parameters, and then throws an exception if some parameters are missing.

The final thing that you need to do is to create the Adolc Tape. This is done by calling the `createTape()` function with the proper arguments. All nonlinear elements that are derived from the `AdolcElement` class must do this.

First the statement `IntVector var(2,0);` create a vector called `var` of two elements and initialize them to 0. Then the statement `var[1] = 1;` initialize the second element to 1. Last, the statement `createTape(var)` creates the tape.

### Building Arguments for `createTape( )`

We will digress from the discussion of the `Tresistor` model for a moment in order to discuss the process of creating state variables in some detail. The `createTape()` function is a member function of the `AdolcElement` class. There are three versions of it corresponding to three different valid function prototypes. The choice of function prototype depends upon the function that the model is to implement.

#### 1: `createTape(IntVector& var)`

- Takes 1 argument, a vector of state variables.
- This is the form used in the `Tresistor` model.

#### 2: `createTape(IntVector& var, IntVector& dvar)`

- Takes 2 arguments – a vector of state variables and a vector corresponding to 1st derivatives of state variables.
- This is the most widely used form, since many physical models use physical variables and their first derivatives.

#### 3: `createTape(IntVector& var, IntVector& dvar, IntVector& d2var, IntVector& t_var, DoubleVector& delay)`

- Takes 5 arguments – a vector of state variables, first derivatives, second derivatives, delayed state variables, and a vector of state variable delays.
- This form must be used 2nd derivatives or time delays are required in models.

For example, suppose that we need to create a new `fREEDATM` element with the following description (`f( )` can be `v( )` or `i( )`):



$$f(x_0, x_1, x_2, \frac{dx_0}{dt}, \frac{dx_1}{dt}, \frac{dx_2}{dt}, \frac{d^2x_2}{dt^2}, x_0(t - \tau_0), x_2(t - \tau_2))$$

Then the arguments of the `createTape( )` function would be as follows:

`createTape(i_vec1, i_vec2, i_vec3, i_vec4, d_vec)`

In this case 4 vectors of type `IntVector&` and one vector of type `DoubleVector&` are needed. The first one, `i_vec1`, consists of the indexes of the state variables. The second one, `i_vec2`, consists of the indexes of the derivatives of the state variables. The third one, `i_vec3`, consists of the indexes of the second derivatives of the state variables. The fourth one, `i_vec4`, consists of the indexes of the state variables that are delayed. The last vector, `d_vec`, contains the value of the delays applied to the state variables created by passing `i_vec4` to `createTape( )`. For this example, let  $\tau_0$  be 1 picosecond and  $\tau_2$  be 0.1 picosecond. To prepare this set of vectors for passing to `createTape( )`, the portion of C++ code in Listing 3.9 would appear in the `init( )` function prior to the call to `createTape( )`.

Listing 3.9: Example creating input vectors passed to `createTape( )`

---

```

1 // Set number of states
2 setNumberOfStates(3); // Three main state vars x0, x1, and x2
3
4 // Argument vector for creating main state variables
5 IntVector i_vec1(3,0); // Initialize 3-element integer array to 0
6 i_vec1(1) = 1; // Set elements for additional state variables
7 i_vec1(2) = 2;
8
9 // Argument vector for 1st derivatives of state variables
10 IntVector i_vec2(3,0); // Initialize 3-element integer array to 0
11 i_vec2(1) = 1; // Set elements for additional 1st derivatives
12 i_vec2(2) = 2;
13
14 // Argument vector for 2nd derivatives of state variables
15 IntVector i_vec3(1,0); // Initialize 1-element integer array to 0
16 i_vec3(0) = 2; // Set element to get 2nd derivative of state var x2!
17
18 // Argument vector for delayed state variables
19 IntVector i_vec4(2,0); // Initialize 2-element integer array to 0
20 i_vec4(1) = 2; // Set element to get delayed copy of state var x2!
21
22 // Argument vector for time delays
23 DoubleVector d_vec(2,0); // Init. 2-element double array to 0
24 d_vec(0) = 1e-12; // Delay state variable x0 by 1 picosecond
25 d_vec(1) = 1e-13; // Delay state variable x2 by 0.1 picosecond

```

---

Notice that `i_vec1` and `i_vec2` are equivalent, so in this case it was not necessary to create `i_vec2`, and one could instead have simply passed `i_vec1` to `createTape( )` twice, in succession:

`createTape(i_vec1, i_vec1, i_vec3, i_vec4, d_vec)`

This emphasizes the point that the variables of type `IntVector&` and `DoubleVector&` are used only to create the collection of state variables and their derivatives and delayed versions. A survey of the source code library for existing *fREEDA*<sup>TM</sup> elements will reveal many instances of this sort of variable re-use in calls to `createTape( )`.

After the completion of the call to `createTape( )`, a vector variable `x[]` of type `adoublev&` will have been created. `x[]` is limited in scope to the instance of the model object. The number of elements in `x[]` will be equal to the number of unique mathematical objects requested in the call to `createTape( )`. For the

example here, there are 3 state variables, 3 first derivatives of state variables, 1 second derivative of a state variable, and 2 delayed state variables, so the total number of elements in `x[]` will be 9, with indices running from 0 to 8. State variables and their derivatives or delays are assigned indices in `x[]` consistent with the order of argument passage to `createTape()`. Table 3.3 summarizes the relationship between vector input variables to `createTape()` and the resulting state variables created by `createTape()` for the example call described here:

Table 3.3: State Variable representation in *f*FREEDA<sup>TM</sup>.

Physical to State Variable Mapping	
Physical Variable	X Variable
$x_0$	<code>x[0]</code>
$x_1$	<code>x[1]</code>
$x_2$	<code>x[2]</code>
$dx_0/dt$	<code>x[3]</code>
$dx_1/dt$	<code>x[4]</code>
$dx_2/dt$	<code>x[5]</code>
$d^2x_2/dt^2$	<code>x[6]</code>
$x_0(t - \tau_0)$	<code>x[7]</code>
$x_2(t - \tau_2)$	<code>x[8]</code>

One final note on constructing the arguments to `createTape()`: There are times when the implementation of a complex function may not require the creation of state variables corresponding to each argument in the most complex function prototype for `createTape()`. For example, suppose in the previous example, that we did not need the second derivative of the second state variable, but we still needed the time-delayed variables. Then instead of declaring an `IntVector` called `ivec_3` of non-zero length, we would declare vector without a declared length, or a null vector, and pass this as an argument to `createTape()` instead of `ivec_3`. It is common in many *f*FREEDA<sup>TM</sup> elements for this null vector to be called `novar`. For example, if indeed we wished to ignore second derivatives in the example in Listing 3.9, then we would alter lines 14 through 16 as follows in Listing 3.10 and then call `createTape()` as follows:

`createTape(i_vec1, i_vec2, novar, i_vec4, d_vec)`

Listing 3.10: Declaring a null vector argument for `createTape()`

---

```

14 // Argument vector for 2nd derivatives of state variables
15 IntVector novar           // Initialize a null vector (no elements)
16                          // Nothing to set!
```

---

*Notice: A limitation in the functionality of the Adol-C Library has recently been uncovered: The second derivative of state variables, normally invoked by using the most complex form of the `createTape` prototype, has been determined to be non-functional. Thus, the use of `novar` as noted in listing 3.10 is mandatory. The development of the `NAdolcElement` class is an attempt to work around this limitation. See section 3.3 for further information.*

Returning our attention to the `Tresistor` element, the function being implemented was `f(x0, x1)`, so that only one vector filled with 0, and 1 was needed. To better understand the process of creating the Adolc Tape, model writers are advised to read the implementation of the `createTape()` function which is a member function of the `AdolcElement` Class. Model writers are also advised to read the ADOL-C documentation [107].

### Constructor for Linear instance of Tresistor

The rest of the code in the `init()` function implements the linear part of the Constructor (invoked when `pdr == false` in the netlist instantiation) and is shown in Listing 3.11. Please refer to [106] for more details.

Listing 3.11: Init function of Tresistor.cc (continued)

---

```

16
17  else{ // pdr==false
18      // Set the number of terminals
19      setNumTerms(2);
20      // Set flags
21      setFlags(LINEAR | ONE_REF | TR_FREQ_DOMAIN);
22      // calculate the resistor value
23      if (isSet(&r0)){
24          r = r0*(1+tc1*(t-tnom)+tc2*(t-tnom)*(t-tnom));
25      }
26      else {
27          if (!isSet(&l))
28              throw("r0 or l must be specified for the resistance");
29          if (!isSet(&rsh))
30              throw("rsh must be specified for the resistance");
31          r = rsh*((1-narrow)/(w-narrow))*(1+tc1*(t-tnom)+tc2*(t-tnom)*(t-tnom));
32      }
33  }
34 }

```

---

### Local Reference Nodes and the `getLocalRefIdx( )` function

We will digress again from the `Tresistor` model for a moment to discuss the local reference node concept and some related modelling conventions inherent in `fREEDA`<sup>TM</sup>.

A spatially distributed circuit is a circuit which, by definition, is much larger than the wavelengths of the signals it processes. In such circuits, the notion of a single reference node or ground terminal is inapt, since currents passing through such a reference node from one piece of spatially distributed circuitry to another cannot physically do so instantaneously. To overcome this limitation and permit modelling of circuits which may be spatially distributed, the local reference node concept was developed. Put adroitly (but in a way that should be widely understood), local reference nodes allow the definition of multiple "ground" terminals so that spatially distinct circuitry can reference their local circuits to these physically isolated reference nodes. Readers interested in understanding more about the theory and the implementation of the local reference node concept should refer to [109] and [110].

The availability of local reference nodes in `fREEDA`<sup>TM</sup> enables the definition of spatially distributed circuit models. Before discussing the coding details, which are straightforward, some modelling conventions used in `fREEDA`<sup>TM</sup> will first be described. These conventions are familiar to researchers in the circuit simulation community, but may not be familiar to new `fREEDA`<sup>TM</sup> users who wish to write `fREEDA`<sup>TM</sup> models.

Consider the "Blob" circuit shown in Figure 3.1. Blob is a circuit that has 5 terminals and is spatially distributed. It has three circuit terminals with terminal numbers 0, 1, and 3, and two reference nodes which have terminal numbers 2 and 4, as shown in the figure. Circuit terminals 0 and 1 use terminal 2 as a local reference node and circuit terminal 3 uses terminal 4 as a local reference node. Although details of the internal operation of "Blob" may not be known, it is clear that the circuits associated with terminals 0 and 1 are spatially distinct from the circuits associated with terminal 3.

Note that in all `fREEDA`<sup>TM</sup> model *source code definitions*, terminal number 0 is *always* a circuit terminal. This may seem incorrect to SPICE users familiar with using node 0 as the ground terminal in netlist definitions. However, note that a `fREEDA`<sup>TM</sup> netlist still permits the use of node 0 in the netlist for the ground terminal. Bear in mind the distinction between coding a model definition, which we are doing here, and instantiating a model, which is done in a netlist.

Figure 3.1 contains other information worth noting. First, note that each state variable is assigned, in order, to each *non-reference node* in the model. Second, notice the mention of the voltage variables `vp[]`

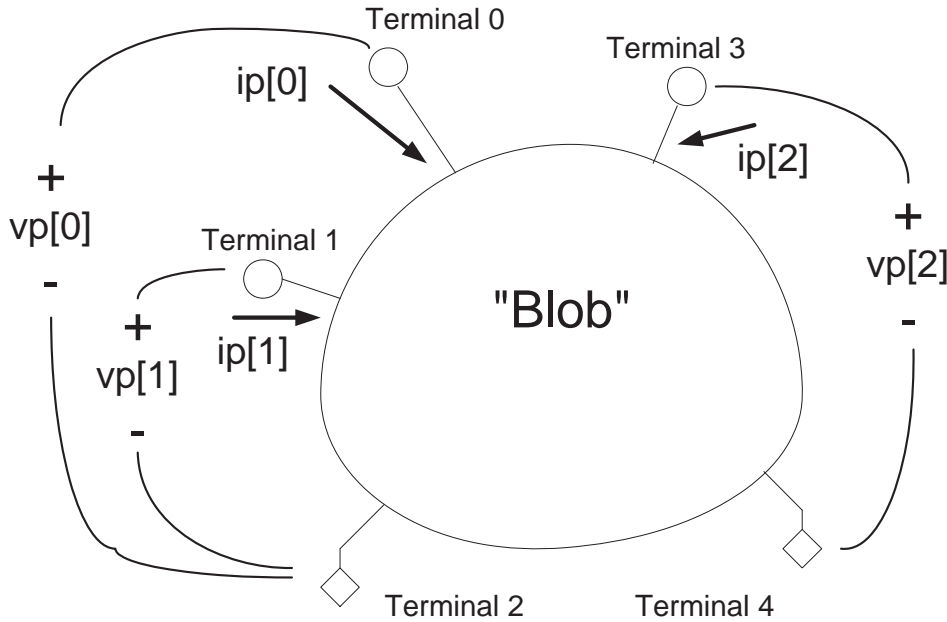


Figure 3.1: Spatially-distributed Blob circuit.

and the current variables `ip[]`. These variables are also initially created in the call to `createTape()`, and are made available to each model instance's `eval()` function. (The `eval()` function will be discussed in detail in section 3.2.4.) For each state variable created by the first argument in the call to `createTape()`, a corresponding `vp[]` and `ip[]` variable is created. Third, note that by convention, the direction of the current variable `ip[]` is into the terminal, and the voltage `vp[]` is defined positively at the circuit terminal and negatively at the corresponding reference node. Users wishing to define currents or voltages contrary to this convention may do so by simply negating the numerical expressions for `vp[]` and `ip[]` that appear in the model's `eval()` function. Finally, note that there are no state variables associated with the reference nodes, since voltages are defined relative to them and Kirchoff's current law requires that the sum of the non-reference terminal currents associated with a particular reference node must flow into that reference node.

Those accustomed to having a "port" perspective for models will notice that, in general, the number of ports is equal to the number of required state variables, even if local reference terminals are shared among one or more ports.

Returning now to the model code, implementation of the `getLocalRefIdx()` function is very simple and straight forward. Refer to Listing 3.12. The first two lines must always be included. Then, we push back each terminal in the terminal list by writing `term_list.push_back(getTerminal(n));` where "n" is the index of the terminal. When we push back a local reference node we must also push the node back into the local reference vector by writing `local_ref_vec.push_back(m);` where "m" is the index of the terminal that we want it to be a local reference.

The `getLocalRefIdx()` function definition follows in Listing 3.12.

Listing 3.12: `getLocalRefIdx` function of `Tresistor.cc`


---

```

122
123 void Tresistor::getLocalRefIdx(UnsignedVector& local_ref_vec ,
124                               TerminalVector& term_list)
125 {
126     // Make sure the vectors are empty
127     term_list.erase(term_list.begin(), term_list.end());

```

---

```

128 local_ref_vec.erase(local_ref_vec.begin(), local_ref_vec.end());
129
130 // Insert vector elements
131 term_list.push_back(getTerminal(0));
132 term_list.push_back(getTerminal(1)); // Local reference terminal
133 local_ref_vec.push_back(1); // Local reference index
134
135 if (pdr){
136     term_list.push_back(getTerminal(2));
137     term_list.push_back(getTerminal(3)); // Local reference terminal
138     local_ref_vec.push_back(3); // Local reference index
139 }
140
141 }

```

---

Finally, note that if a model has only 1 local reference node, then it is not necessary to declare the `getLocalRefIdx` function at all in the header file and it may be omitted from the source code file. Note that when `getLocalRefIdx` is not used, then the highest numbered terminal is defined as the local reference terminal. Referring to listings 3.8 and 3.11 where the call to `setNumTerms()` appears, if `n` is the argument to `setNumTerms()`, then there will be `n` terminals numbered `0,1,...,n-1`, and when `getLocalRefIdx` is not used, then `n-1` will be the terminal number of the reference node.

### The Eval function and using `condassign()`

The `eval()` function definition for `Tresistor` is shown in Listing 3.13.

Listing 3.13: Eval function of `Tresistor.cc`

---

```

100
101 void Tresistor::eval(adoublev& x,
102                     adoublev& vp, adoublev& ip)
103 {
104     // x[0]: resistor voltage
105     // x[1]: deltatemp in deg. Celsius
106
107     vp[0]=x[0];
108     vp[1]=x[1]+tnom+273; //vp[1]==tp[1] in Kelvin
109
110     adouble res;
111     if (isSet(&r0)){
112         res = r0 * (one + tc1 * x[1] + tc2 * x[1] * x[1]);
113     }
114     else {
115         res = rsh * ((l-narrow) / (w-narrow))
116             * (one + tc1*x[1] + tc2 * x[1] * x[1]);
117     }
118
119     ip[0] = x[0] / res;
120     ip[1] = - x[0] * ip[0]; //ip[1]==pp[0];
121 }

```

---

The `eval()` function is the implementation of the device equations. It is in here where the voltages and currents are related to the state variables. The `eval()` function takes three vectors of type `adoublev&`, the first one is the state variable vector, and it is considered as the input, the second and the third one are the voltages and the currents, and these two vectors are considered as output. The `eval()` function is called from the `createTape()` function. In this way, the Adol-C library keeps track of all operations that are performed on the `adouble` variables so that it can evaluate the derivatives whenever needed.

Due to encapsulation of most of the low-level handling of the state variables within `AdolcElement`, manipulation of the state variables within the `eval()` function for an element can be done in a straightforward

fashion using the same operators as that for floating-point arithmetic. These operators are all overloaded, but this is transparent in most cases. However, when conditional branches affecting the assignment of active variables are to be made within an `eval()` function, it is mandatory to use a routine with the Adol-C library designed to facilitate conditional branching. This routine is called `condassign()`.

The `condassign()` routine normally takes 4 arguments which can be of type adouble or scalars. For users familiar with C and C++, a function call such as `condassign(a,b,c,d)` corresponds with the syntax of the following conditional assignment:

$$\mathbf{a} = (\mathbf{b} > 0) ? \mathbf{c} : \mathbf{d}$$

For those familiar with Matlab, Listing 3.14 shows how the same conditional branching statements might be implemented in that language.

Listing 3.14: Matlab equivalent of `condassign()`

---

```

1 if (b > 0)
2     a = c
3 else
4     a = d
5 end

```

---

As an illustrative example, consider a rudimentary model for a semiconductor junction diode. Define the current as being 0 whenever the voltage across the junction is less than, say, 0.7 volts, and let the current be non-zero above the threshold. Let voltage be the parametric state variable, so  $v(t) = x(t)$ , and let the current be defined as follows:

$$i(t) = \begin{cases} I_s \left[ \exp\left(\frac{x(t)-0.7}{v_T}\right) - 1 \right] & \text{if } x(t) > 0.7 \text{ or } x(t) - 0.7 > 0 \\ 0 & \text{if } x(t) \leq 0.7 \end{cases}$$

In this diode equation,  $I_s$  is the reverse saturation current and  $v_T$  is the "temperature voltage." Both are constants. Letting `x[0]` be the state variable parameter  $x(t)$ , `vp[0]` be the junction voltage  $v(t)$ , and `ip[0]` be the current  $i(t)$  across the junction, Listing 3.15 shows how the conditional branch for the simple junction diode model would be implemented.

Listing 3.15: Example of branch for simple junction diode.

---

```

1 vp[0] = x[0];                                \\\ voltage equals parameter
2 condassign(ip[0], x[0]-0.7,                    \\\ a, b with b's condition shifted to 0.7
3             Is*(exp((x[0]-0.7)/vt)-1),         \\\ c == exponential characteristic
4             0);                               \\\ d == 0

```

---

## The fillMNAM function

The last two methods in the source file, in Listing 3.16, are the routines used to fill the MNAM of the circuit. Please refer to [106] for a detailed explanation on how to fill the MNAM.

Listing 3.16: fillMNAM function of `Tresistor.cc`

---

```

144
145 void Tresistor::fillMNAM(FreqMNAM* mnam)
146 {
147     // Ask my terminals the row numbers
148     mnam->setAdmittance(getTerminal(0)->getRC(), getTerminal(1)->getRC(),
149                        one/r);
150 }
151
152
153 void Tresistor::fillMNAM(TimeMNAM* mnam)
154 {

```

```

155 // Ask my terminals the row numbers
156 mnam->setMAdmittance(getTerminal(0)->getRC(), getTerminal(1)->getRC(),
157                      one/r);
158 }

```

---

### 3.2.5 Using the cout routine for debugging

In the course of developing models, debugging will inevitably be necessary. In this section we provide a few suggestions on using the C++ standard output routine `cout` to facilitate debugging. Often in model development, modelling equations are valid over restricted portions of the input domain. Considering the rudimentary p-n junction diode again, suppose the current is undefined for voltages below the threshold voltage, rather than set to zero. We dispense entirely with the `condassign()` statement, but now it would be useful to have some sort of error notification to the user if voltages below the threshold are present. Listing 3.17 shows an example of how the code for the rudimentary diode might be modified to use `cout` to print such a message.

Listing 3.17: Using `cout` to facilitate model debugging.

---

```

1 vp[0] = x[0];                                \\ voltage equals parameter
2 if (x[0] < 0.7)
3 {
4     cout << "x =" << x[0] << ".. Invalid input voltage." << endl;
5 }
6 ip[0] = Is*(exp((x[0]-0.7)/vt)-1);          \\ exponential characteristic

```

---

Another use for `cout` is to validate that model parameters are being passed as expected from the netlist into model instances. In these cases, `cout` statements would appear in the `init()` function call for the element and echo the parameter values of the model instance.

Note that when an active variable is streamed to the standard output, the real value of the variable is followed by an "(a)" to indicate that the variable is active. This is a result of the overloading of the "<<" operator by the Adol-C library.

### 3.2.6 Modifications to the rest of the FREEDA™ Source Files

To be completed.

### 3.2.7 Header File

The header file template is shown in Listing 3.18.

Listing 3.18: Element1.h

---

```

1
2 // This may look like C code, but it is really -*- C++ -*-
3 //
4 // This is a generic element template
5 //
6 //
7 //      +-----+
8 //      o-----+-----+-----o
9 //      +-----+
10
11 #ifndef Element1.h
12 #define Element1.h 1
13
14 class Element1 : public AdolcElement
15 {
16 public:
17

```

```

18 Element1(const string& iname);
19
20 ~Element1() {}
21
22 static const char* getNetlistName()
23 {
24     return einfo.name;
25 }
26
27 // Do some local initialization
28 virtual void init() throw(string&); // to perform initialization that
29                                     // cannot be done by the constructor
30                                     // and to create the Adolc Tape
31
32 // Implement the getLocalRefIdx function if you are creating
33 // an element with multiple reference nodes.
34 // Get a vector with the indexes of the local reference nodes.
35 // Get terminal pointers in term_list
36 // ordered by local reference node:
37 //
38 // t0 t1 t2 t3
39 //
40 //   ^   ^
41 //  LRN1 LRN2
42 //
43 // local_ref_vec contains: {1, 3}
44 //
45 virtual void getLocalRefIdx(UnsignedVector& local_ref_vec, // to implement a spatially
46                             TerminalVector& term_list); // distributed element, or an
47                                                         // electro-thermal one with
48                                                         // multiple reference nodes.
49
50 private:
51
52 // Implement the eval function. It is here where you write
53 // your device equations.
54 virtual void eval(adoublev& x, adoublev& vp, adoublev& ip);
55
56 // Parameter variables (replace by your parameter variables here)
57 double par1, par2, ..., parn;
58
59 // Element information
60 static ItemInfo einfo;
61
62 // Number of parameters of this element
63 static const unsigned n_par;
64
65 // Parameter information
66 static ParmInfo pinfo[];
67
68 };
69
70 #endif

```

---

### 3.2.8 Class Source File

The class source file template is shown in Listing 3.19.

Listing 3.19: Element1.cc

---

```

1
2 #include " ../network/ElementManager.h"

```



```

3 #include "../network/AdolcElement.h"
4 #include "../analysis/FreqMNAM.h"
5 #include "../analysis/TimeMNAM.h"
6 #include "Element1.h"
7
8 // Static members (set to the number of parameters of your element)
9 const unsigned Element1::n_par = ;
10
11 // Element information
12 ItemInfo Element1::einfo = {
13     "tres",
14     "Element1",
15     "Your Name",
16     "Reserved for documentation use"
17 };
18
19 // Parameter information
20 ParmInfo Element1::pinfo[] = {
21     {"par1", "Parameter 1 (Ohms)", TR_DOUBLE, true},
22     {"par2", "Parameter 2 (The Unit)", TR_DOUBLE, false}
23     .
24     .
25     .
26     {"parn", "Parameter n (The Unit)", TR_BOOLEAN, false}
27 };
28
29 Element1::Element1(const string& iname) : AdolcElement(&einfo, pinfo, n_par, iname)
30 {
31     // Set parameters
32     paramvalue[0] = &par1;
33     paramvalue[1] = &(par2 = 1e-12); // Example of how to set a default value.
34     .
35     .
36     .
37     paramvalue[n] = &(parn = false);
38 }
39
40 void Element1::init() throw(string&)
41 {
42     // Set the number of terminals (Set to the number of terminals of
43     // your element)
44     setNumTerms(4);
45
46     // Set flags. If your element has multiple reference nodes,
47     // set the second flag to MULTLREF
48     setFlags(NONLINEAR | ONE_REF | TR_TIME_DOMAIN);
49
50     // Set number of states
51     setNumberOfStates(2);
52
53     // create the tape
54     IntVector var(2,0);
55     var[1] = 1;
56     createTape(var);
57 }
58
59 void Element1::eval(adoublev& x,
60                    adoublev& vp, adoublev& ip)
61 {
62     // Implement the eval function
63     // write the device equations
64 }

```

```

65
66 void Element1::getLocalRefIdx(UnsignedVector& local_ref_vec,
67                               TerminalVector& term_list)
68 {
69     // Make sure the vectors are empty
70     // always write these functions as they are
71     term_list.erase(term_list.begin(), term_list.end());
72     local_ref_vec.erase(local_ref_vec.begin(), local_ref_vec.end());
73
74     // Insert vector elements
75     // push_back all the terminals in the term_list
76     // push_back all the local reference terminal in the local_ref_vec
77     term_list.push_back(getTerminal(0));
78     term_list.push_back(getTerminal(1)); // Local reference terminal
79     local_ref_vec.push_back(1); // Local reference index
80     term_list.push_back(getTerminal(2));
81     term_list.push_back(getTerminal(3)); // Local reference terminal
82     local_ref_vec.push_back(3); // Local reference index
83 }

```

---

### 3.3 A Note on the Eval Routines in fFREEDA<sup>TM</sup>

Most nonlinear computations require the evaluation of first and higher derivatives of vector functions with  $m$  components in  $n$  real or complex variables. fFREEDA<sup>TM</sup> uses the package ADOL-C which automatically computes the derivatives of non-linear functions by successive implementation of the chain rule. These calculation occur at a fraction of the time and is relatively free of truncation errors. The interface details to the ADOL-C library from within fFREEDA<sup>TM</sup> can be found in path\_to\_freeda/simulator/network/NAdolcElement.cc.

Every nonlinear element in fFREEDA<sup>TM</sup> has an eval function that describes the constitutive equations for the particular element at any instant. This function is called from within NAdolcElement.cc during evaluation of the tape. For more details on the tape, refer to ADOL-C documentation.

#### 3.3.1 Parameterized Device Models

A non-linear device model can be described with the following set of equations

$$\mathbf{v}(t) = \mathbf{v}(\mathbf{x}(t), d\mathbf{x}/dt, \dots, d^m \mathbf{x}/dt^m, \mathbf{x}_D(t)) \quad (3.1)$$

$$\mathbf{i}(t) = \mathbf{i}(\mathbf{x}(t), d\mathbf{x}/dt, \dots, d^m \mathbf{x}/dt^m, \mathbf{x}_D(t)) \quad (3.2)$$

where  $\mathbf{v}(t)$  and  $\mathbf{i}(t)$  are vectors of voltages and currents at the ports of the non-linear device,  $\mathbf{x}(t)$  is a vector of parameters or state variables and  $\mathbf{x}_D(t)$  is a vector of time delayed state variables. There are several models like the microwave diode, the Gummel-Poon BJT and Berkeley's BSIM4 which for accurate modeling, require charge to be defined as a state variable. In some cases it may not be possible to find the voltage as a function of charge and the only alternative is to treat charge as a state variable.

Consider a simplified model for the microwave diode shown in Figure 3.2. The corresponding equations for this model are

$$i(v) = I_s(\exp(\alpha v) - 1) \quad (3.3)$$

and

$$c_j(v) = C_{t0}(1 - v/\phi)^{-\gamma} + C_{d0} \exp(\alpha' v) \quad \text{if } v \leq .8\phi \quad (3.4)$$

$$C_{t0}(.2)^{-\gamma} + C_{d0} \exp(\alpha' v) \quad \text{if } v \geq .8\phi \quad (3.5)$$

where  $v$  is the junction voltage. The capacitor voltage  $q_j$  is given by

$$q_j(v) = \int_0^v c_j(u) du \quad (3.6)$$

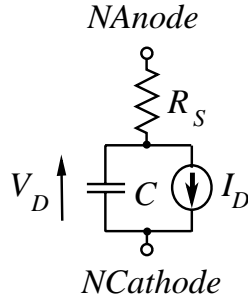


Figure 3.2: Simplified schematic for a diode model

Accurate transient analysis requires  $q_j$  to be chosen as a state variable. We need  $v$  to calculate  $i_1(v)$ . Since it is not possible to solve analytically for  $v(q)$ , the alternative is to model the diode with two state variables, namely  $v$  and  $q$ . The diode equations can now be formulated in two stages. Hence for this case, we will have two *eval* functions; *eval1* and *eval2*. In *eval1*, we will calculate  $i(v)$  and  $q(v)$ , both are which are functions of voltage  $v$ . In this case, the values of  $q(v)$  are passed to *eval2*, which automatically calculates  $dq/dt$ . Hence we now have the current through the capacitor as

$$i_c = \frac{dq}{dt} \quad (3.7)$$

All the code that performs the actual derivatives is outside the nonlinear model and is handled automatically. Currently in *fREEDA*<sup>TM</sup>, only three levels of eval are supported. This means that the highest order of derivatives possible is two. Most nonlinear elements do not require more than two levels of derivatives. The entire formulation can be generalized as follows:

$$\begin{aligned} \text{stage 1} & : \quad f_1(x, x_D) \\ & \quad g_1(x, x_D) \end{aligned} \quad (3.8)$$

$$\begin{aligned} \text{stage 2} & : \quad f_2(f_1, dg_1/dt) \\ & \quad g_2(f_1, dg_1/dt) \end{aligned} \quad (3.9)$$

$$\vdots$$

$$\begin{aligned} \text{stage n} & : \quad v(f_{n-1}, dg_{n-1}/dt) \\ & \quad i(f_{n-1}, dg_{n-1}/dt) \end{aligned} \quad (3.10)$$

### 3.4 Contributors

The following contributed to this chapter:

Frank P. Hart

Houssam S. Kanj

Nikhil Kriplani

Carlos Christoffersen

Michael Steer.



## Appendix A

# Object Oriented Programming Basics

Object oriented programming (OOP) [60] provides a means for abstraction in both programming and design. OOP does not deal with programming in the sense of developing algorithms or data structures but it must be studied as a means for the organization of programs and, more generally, techniques for designing programs.

As the primary means for structuring a program or design, OOP provides *objects*. Objects may model real life entities, may function to capture abstractions of arbitrary complex phenomena, or may represent system artifacts such as stacks or graphics. Operationally, objects control the computation. From the perspective of program development, however, the most important characteristic of objects is not their behavior as such, but the fact that the behavior of an object may be described by an abstract characterization of its interface. Having such a characterization suffices for the design. The actual behavior of the object may be implemented later and refined according to the need. A *class* specifies the behavior of the objects which are its instances. Also, classes act as templates from which actual objects may be created. An *instance* of a class is an object belonging to that class. A procedure (or function) inside an object is called a *method*. A *message* to an object is a request to execute a method.

*Inheritance* is the mechanism which allows the reuse of class specifications. The use of inheritance results in a class hierarchy that, from an operational point of view, decides what is the method that will be selected in response to a message.

Finally, an important feature of OO languages is their support for *polymorphism*. This makes it possible to hide different implementations behind a common interface.

The notion of flow diagram in procedural programming is replaced in OOP by a set of objects which interact by sending messages to each other.

We will briefly review what are traditionally considered to be features and benefits of OOP. Both *information hiding* (also known as *encapsulation*) and *data abstraction* relieve the task of the programmer using existing OO code, since with these mechanisms the programmer's attention is no longer distracted by irrelevant implementation details. The flexible dispatching behavior of objects that lends them their polymorphic behavior is due to the dynamic binding of methods to messages. For the C++ language, polymorphic object behavior is effected by using *virtual functions*, for which in contrast to ordinary functions, the binding to an actual function takes place at run time and not at compile time.

Encapsulation promotes *modularity*, meaning that objects may be regarded as the building blocks of a complex system. Another advantage often attributed to the OOP is code reuse. Inheritance is an invaluable mechanism in this respect, since it enables the programmer to modify the behavior of a class of objects without requiring access to the source code.

Although an object oriented approach to program development indeed offers great flexibility, some of the problems it addresses are intrinsically difficult and cannot really be solved by mechanisms alone. For example, it is more likely to achieve a stable modularization when shifting focus from programming to design.

C++ virtual functions [70] can have big performance penalties such as extra memory accesses or the

possibility of an unpredictable branch so pipelines can grind to a halt (note however, that some architectures have branch caches which can avoid this problem). There are several research projects which have demonstrated success at replacing virtual function calls with direct dispatches. Note however, that virtual functions are not always bad when it comes to performance. The important questions are: How much code is inside the virtual function? How often is it used? If there is a lot of code (*i.e.* more than 25 flops), then the overhead of the virtual function will be insignificant. But if there is a small amount of code and the function is called very often (*e.g.* inside a loop), then the overhead can be critical.

## A.1 UML diagrams

The Unified Modeling Language (UML) [83, 65] is a language for specifying, visualizing, and constructing the artifacts of software systems as well as for business modeling. The goal of UML is to become a common language for creating models of object oriented computer software. It is used here to graphically illustrate the relationship of classes using what is called a class diagram. A class diagram is a graph of Classifier elements with connections indicating by their various static relationships (Figure 1.1). (Note that a “class” diagram may also contain interfaces, packages, relationships, and even instances, such as objects and links. Perhaps a better name would be “static structural diagram” but “class diagram” is shorter and its use is well established.) A class is drawn as a solid-outline rectangle with 3 compartments separated by horizontal lines. The top name compartment holds the class name and other general properties of the class (including stereotype); the middle list compartment holds a list of attributes; the bottom list compartment holds a list of operations. Either or both of the attribute and operation compartments may be suppressed. A separator line is not drawn for a missing compartment. If a compartment is suppressed, no inference can be drawn about the presence or absence of elements in it.

Each instance of type **Element**, for example, seems to contain an instance of type **ElementData**. This class relationship is indicated by the joining line. The relationship is composition — indicated by the solid diamond symbol. The arrowhead denotes that the relationship is navigable in only one direction, *i.e.*, **ElementData** does not know anything about **Element**. The inheritance relationship in UML is depicted by the triangular arrowhead and points to the base class. A line from the base of the arrowhead connects it to the derived classes, *e.g.* **Element** is derived from **NetListItem**.

Other forms of containment do not have whole/part implications and are called association relationships indicated by a line drawn between the participating classes. (This relationship will almost certainly be implemented using pointers unless it is very weak.) If the relationship between two classes is very weak (*i.e.* very little data is shared) then a dashed line is used. For example, in Figure 1.1, **ElementManager** somehow depends upon **Diode**. (In C++ the weak relationship is almost always implemented using an `#include`.)

An illustration showing examples for the notation is given in Figure A.1.

### A.1.1 Collaboration Diagrams

The collaboration diagram [83] is part of the Unified Modeling Language (UML), a language for specifying, visualizing, and constructing the artifacts of software systems as well as for business modeling. The UML represents a collection of ‘best engineering practices’ that have proven successful in the modeling of large and complex systems. Behavior is implemented by sets of objects that exchange messages within an overall interaction to accomplish a purpose. To understand the mechanisms used in a design, it is important to see only the objects and the messages involved in accomplishing a purpose or a related set of purposes, projected from the larger system of which they are part for other purposes. Such a static construct is called a collaboration. A collaboration is a set of participants and relationships that are meaningful for a given set of purposes. The identification of participants and their relationships does not have global meaning. In the collaboration diagram, each box represents an object (in this case, either a terminal or an element). The lines between boxes represent associations and the arrows are messages. The order in which messages are sent is shown by the numbers.

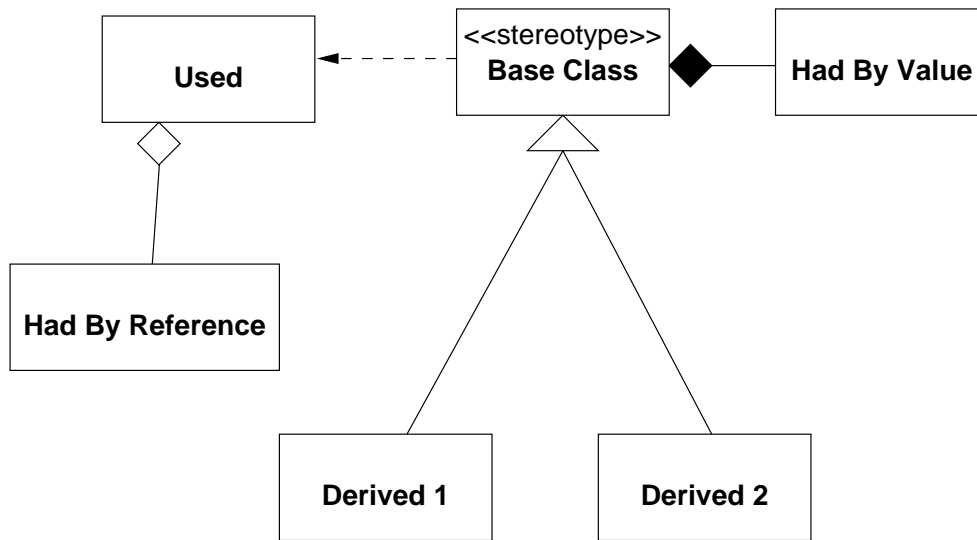


Figure A.1: Notation for a class diagram.

## A.2 Contributors

The following contributed to this chapter

Carlos Christoffersen

Michael Steer.





# Appendix B

## Support Libraries

A large number of support libraries are available (many of them freely) and some of these are used in *fREEDA*<sup>TM</sup>. The various libraries, which should be of general interest to the microwave modeling community, are described below.

### B.1 Solution of Sparse Linear Systems (Sparse, SuperLU)

*Sparse 1.3*<sup>1</sup> [73] is a flexible package of subroutines written in C used to numerically solve large sparse systems of linear equations. The package is able to handle arbitrary real and complex square matrix equations. Besides being able to solve linear systems, it is also able to quickly solve transposed systems, find determinants, and estimate errors due to ill-conditioning in the system of equations and instability in the computations. Sparse also provides a test program that is able to read matrix equation from a file, solve it, and print useful information (such as condition number of the matrix) about the equation and its solution. Sparse was originally written for use in circuit simulators and is well adapted to handling nodal- and modified-nodal admittance matrices.

*SuperLU*<sup>2</sup> is used in the wavelet and time marching transient analyses. It contains a set of subroutines to numerically solve a sparse linear system  $\mathbf{Ax} = \mathbf{b}$ . It uses Gaussian elimination with partial pivoting (GEPP). The columns of  $\mathbf{A}$  may be preordered before factorization; the reordering for sparsity is completely separate from the factorization. SuperLU is implemented in ANSI C. It provides support for both real and complex matrices, in both single and double precision.

### B.2 Vectors and Matrices (MV++, MTL)

Most of the vector and matrix handling in *fREEDA*<sup>TM</sup> uses *MV++*<sup>3</sup> [72]. This is a small set of vector and simple matrix classes for numerical computing written in C++. It is not intended as a general vector container class but rather designed specifically for optimized numerical computations on RISC and pipelined architectures which are used in most new computer architectures. The various MV++ classes form the building blocks of larger user-level libraries. The MV++ package includes interfaces to the computational kernels of the Basic Linear Algebra Subprograms package (BLAS) which includes scalar updates, vector sums, and dot products. The idea is to utilize vendor-supplied, or optimized BLAS routines that are fine-tuned for particular platforms.

The *Matrix Template Library* (MTL)<sup>4</sup> is a high-performance generic component library that provides comprehensive linear algebra functionality for a wide variety of matrix formats. It is used in the wavelet

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<sup>1</sup><http://www.netlib.org/sparse/>

<sup>2</sup><http://www.nersc.gov/xiaoye/SuperLU/>

<sup>3</sup><http://math.nist.gov/mv++/>

<sup>4</sup><http://www.lsc.nd.edu/research/mtl/>

and time marching transient analyses.

As with STL, MTL uses a five-fold approach, consisting of generic functions, containers, iterators, adaptors, and function objects, all developed specifically for high performance numerical linear algebra. Within this framework, MTL provides generic algorithms corresponding to the mathematical operations that define linear algebra. Similarly, the containers, adaptors, and iterators are used to represent and to manipulate matrices and vectors.

### B.3 Solution of Nonlinear Systems (NNES)

Nonlinear systems of equations in *fREEDA*<sup>TM</sup> are solved using the *NNES*<sup>5</sup> [58] library. This package is written in Fortran and provides Newton and quasi-Newton methods with many options including the use of analytic Jacobian or forward, backwards or central differences to approximate it, different quasi-Newton Jacobian updates, or two globally convergent methods, etc. This library is used through an interface class (**NLSInterface**), so it is possible to install a different routine to solve nonlinear systems if desired by just replacing the interface (four different nonlinear solvers have already been used). The Fortran routine *NLEQ1* (Numerical solution of nonlinear (NL) equations (EQ)<sup>6</sup>) can also be used as a compile option.

### B.4 Fourier Transform (FFTW)

Fourier transformation is implemented in *fREEDA*<sup>TM</sup> using the *FFTW*<sup>7</sup> library [51]. FFTW is a C sub-routine library for computing the Discrete Fourier Transform (DFT) in one or more dimensions, of both real and complex data, and of arbitrary input size. Benchmarks, performed on a variety of platforms show that FFTW's performance is typically superior to that of other publicly available FFT software. Moreover, FFTW's performance is portable: the program performs well on most computer architectures without modification.

### B.5 Automatic Differentiation (Adol-C)

Most nonlinear computations require the evaluation of first and higher derivatives of vector functions with  $m$  components in  $n$  real or complex variables [71]. Often these functions are defined by sequential evaluation procedures involving many intermediate variables. By eliminating the intermediate variables symbolically, it is theoretically always possible to express the  $m$  dependent variables directly in terms of the  $n$  independent variables. Typically, however, the attempt results in unwieldy algebraic formulae, if it can be completed at all. Symbolic differentiation of the resulting formulae will usually exacerbate this problem of *expression swell* and often entails the repeated evaluation of common expressions.

An obvious way to avoid such redundant calculations is to apply an optimizing compiler to the source code that can be generated from the symbolic representation of the derivatives in question. Exactly this approach was investigated by Speelpenning during his Ph.D. research [74] at the University of Illinois from 1977 to 1980. Eventually he realized that at least in the cases  $n = 1$  and  $m = 1$ , the most efficient code for the evaluation of derivatives can be obtained directly from the evaluation of the underlying vector function. In other words, he advocated the differentiation of evaluation algorithms rather than formulae. In his dissertation he made the particularly striking observation that the gradient of a scalar-valued function (*i.e.*  $m = 1$ ) can always be obtained with no more than five times the operations count of evaluating the function itself. This bound is completely independent of  $n$ , the number of independent variables, and allows the row-wise computation of Jacobians for at most  $5m$  times the effort of evaluating the underlying vector function.

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<sup>5</sup><http://www.netlib.org/opt/>

<sup>6</sup>Konrad-Zuse-Zentrum für Informationstechnik Berlin (ZIB). Contact: Lutz Weimann, ZIB, Division Scientific Computing, Department Scientific Software, e-mail: [weimann@zib.de](mailto:weimann@zib.de)

<sup>7</sup><http://www.fftw.org>

Given a code for a function  $F : \mathbb{R}^n \rightarrow \mathbb{R}^m$ , automatic differentiation (AD) uses the chain rule successively to compute the derivative matrix. AD has two basic modes, forward mode and reverse mode [75]. The difference between these two is the way the chain rule is used to propagate the derivatives.

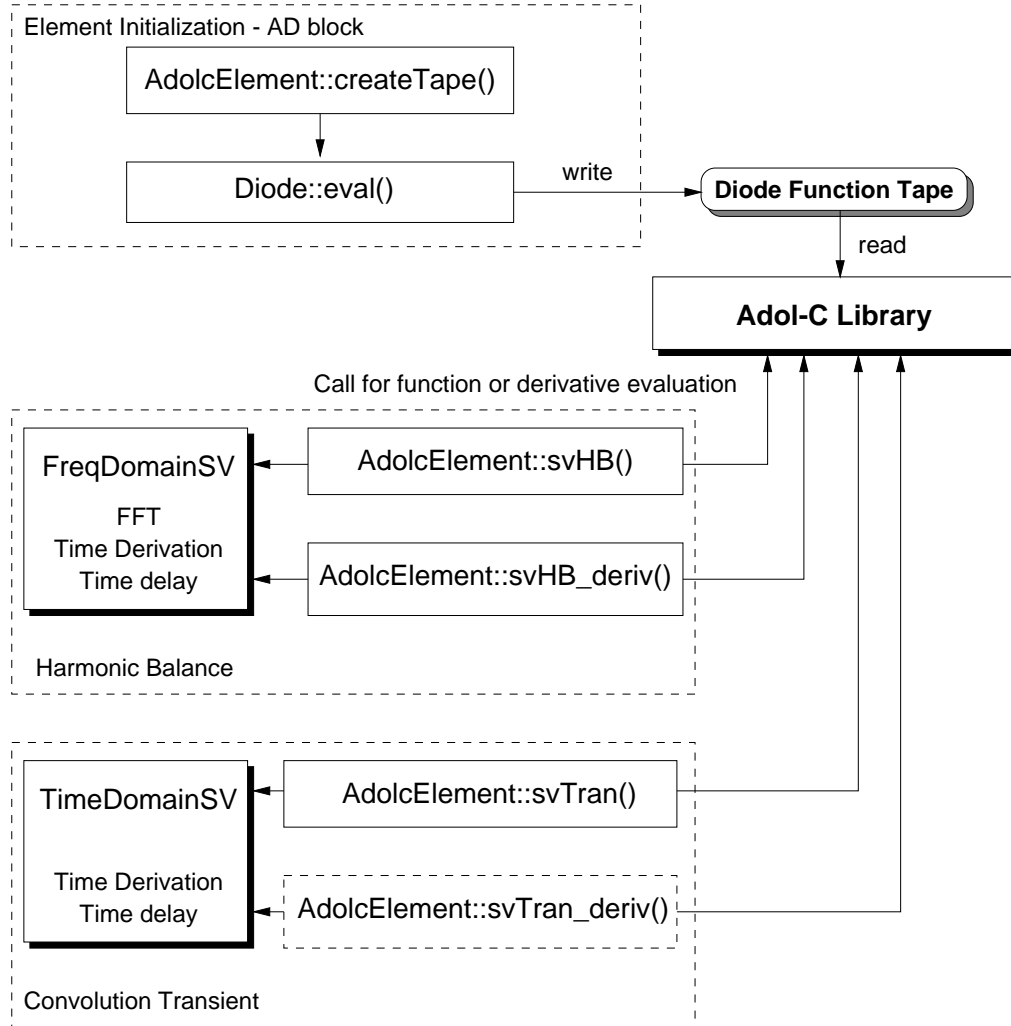


Figure B.1: Implementation of automatic differentiation.

A versatile implementation of the AD technique is *Adol-C*<sup>8</sup> [71], a software package written in C and C++. The numerical values of derivative vectors (for example, required to fill a Jacobian in Harmonic Balance analysis [2], see Figure B.1) are obtained free of truncation errors at a small multiple of the run time required to evaluate the original function with little additional memory required. It is important to note that AD is not numerical differentiation and the same accuracy achieved by evaluating analytically developed derivatives is obtained.

The `eval()` method of the nonlinear element class is executed at initialization time and so the operations to calculate the currents and voltages of each element are recorded by Adol-C in a *tape* which is actually an internal buffer. After that, each time that the values or the derivatives of the nonlinear elements are required, an Adol-C function is called and the values are calculated using the tapes. This implementation

<sup>8</sup><http://www.math.tu-dresden.de/~adol-c/>

is efficient because the taping process is done only once (this almost doubles the speed of the calculation compared to the case where the functions are taped each time they are needed). When the Jacobian is needed, the corresponding Adol-C function is called using the same tape. We have tested the program with large circuits with many tones, and the function or Jacobian evaluation times are always very small compared with the time required to solve the matrix equation (typically some form of Newton's method) that uses the Jacobian. The conclusion is that there is little detriment to the performance of the program introduced by using automatic differentiation. However the advantage in terms of rapid model development is significant. The majority of the development time in implementing models in simulators, particularly harmonic balance simulators, is in the manual development of the derivative equations. Unfortunately the determination of derivatives using numerical differences is not sufficiently accurate for any but the simplest circuits. With Adol-C full 'analytic' accuracy is obtained and the implementation of nonlinear device models is dramatically simplified. From experience the average time to develop and implement a transistor model is an order of magnitude less than deriving and coding the derivatives manually. Note that time differentiation, time delay and transformations are left outside the automatic differentiation block. The calculation speed achieved is approximately ten times faster than the speed achieved by including time differentiation, time delay and transformations inside the block.

## B.6 Contributors

The following contributed to this chapter

Carlos Christoffersen

Michael Steer.

## Appendix C

# Thermal Resistor

Breakout of the code for the electro-thermal resistor model, resistorT.

```
#include "../network/ElementManager.h"
#include "../analysis/FreqMNAM.h"
#include "../analysis/TimeMNAM.h"
#include "Tresistor.h"
```

Part 1 : Include Files.

```
// Static members
const unsigned int n_par = 11;
```

Part 2 : Number of Parameters.

```
// Element information
ItemInfo Tresistor::einfo = {
    "tr",
    "Tresistor",
    "Houssam S. Kanj",
    DEFAULT_ADDRESS"elements/resistor.html"
};
```

Part 3 : Device instance name  
and Author information.

```
// Parameter information
ParmInfo Tresistor::pinfo[] = {
    {"r0", "Resistance value (Ohms)", TR_DOUBLE, false},
    {"l", "length (meters)", TR_DOUBLE, false},
    {"w", "width (meters)", TR_DOUBLE, false},
    {"t", "system Temperature (Celsius)", TR_DOUBLE, false},
    {"rsh", "sheat resistance (Ohms/sq)", TR_DOUBLE, false},
    {"defw", "default device width (meters)", TR_DOUBLE, false},
    {"narrow", "narrowing due to side etching (meters)", TR_DOUBLE, false},
    {"tnom", "initial Temperature (Celsius)", TR_DOUBLE, false},
    {"tc1", "Temperature Coefficient (1/Celsius)", TR_DOUBLE, false},
    {"tc2", "Temperature Coefficient (1/Celsius)", TR_DOUBLE, false},
    {"pdr", "Power Depenent Resistor", TR_BOOLEAN, false}
};
```

Part 4 :  
Parameter  
Definition.

```
Tresistor::Tresistor(const string& iname) : AdolcElement(&einfo, pinfo, n_par, iname)
{
    // Set parameters
    paramvalue[0] = &(r0);
    paramvalue[1] = &(l);
    paramvalue[2] = &(w);
    paramvalue[3] = &(t);
    paramvalue[4] = &(rsh);
    paramvalue[5] = &(defw = 1e-6);
    paramvalue[6] = &(narrow = zero);
    paramvalue[7] = &(tnom = 27);
    paramvalue[8] = &(tc1 = zero);
    paramvalue[9] = &(tc2 = zero);
    paramvalue[10] = &(pdr=false);
}
```

Part 5 :  
Parameter  
Initialization.

This documentation is correct as of fREEDA version 1.2 . In version 1.3 the use of `DEFAULT_ADDRESS"elements/resistorTH.html` is obsolete. The element is now called `resistorT` and not `tr` .

```

void resistor::init() throw(string&)
{
    if (!isSet(&w))
        w = defw;
    if (!isSet(&t))
        t = tnom;

    if (pdr) {
        // Set the number of terminals
        setNumTerms(4);
        // Set flags
        setFlags(NONLINEAR | MULTI_REF | TR_TIME_DOMAIN);
        // Set number of states
        setNumberOfStates(2);

        if (!isSet(&r0)) {
            if (!isSet(&l))
                throw("r0 or l must be specified for the resistance");
            if (!isSet(&rsh))
                throw("rsh must be specified for the resistance");
        }
        // create tape
        IntVector var(2,0);
        var[1] = 1;
        createTape(var);
    }
    else { // pdr==false
        // Set the number of terminals
        setNumTerms(2);
        // Set flags
        setFlags(LINEAR | ONE_REF | TR_FREQ_DOMAIN);
        // calculate the resistor value
        if (isSet(&r0)) {
            r = r0*(1+tc1*(t-tnom)+tc2*(t-tnom)*(t-tnom));
        }

        else {
            if (!isSet(&l))
                throw("r0 or l must be specified for the resistance");
            if (!isSet(&rsh))
                throw("rsh must be specified for the resistance");
            r = rsh*((l-narrow)/(w-narrow))*(1+tc1*(t-tnom)+tc2*(t-tnom)*(t-tnom));
        }
    }
}

```

Part 6 : State  
Variable definition  
for a power  
dissipating resistor  
(pdr).

Part 7 : State  
Variable  
definition for a  
linear resistor.

```

void resistor::eval(adoublev& x, adoublev& vp, adoublev& ip)
{
    // x[0]: resistor voltage
    // x[1]: deltatemp in deg. Celsius

    vp[0]=x[0];
    vp[1]=x[1]+tnom+273; //vp[1]==tp[1] in Kelvin
}

```

Part 8 : Output vector  
assignment : voltage  
and temperature  
outputs.



```

void resistorT::init() throw(string&)
{
    if (!isSet(&w))
        w = defw;
    if (!isSet(&t))
        t = tnom;

    if (pdr) {
        // Set the number of terminals
        setNumTerms(4);
        // Set flags
        setFlags(NONLINEAR | MULTI_REF | TR_TIME_DOMAIN);
        // Set number of states
        setNumberOfStates(2);

        if (!isSet(&r0)) {
            if (!isSet(&l))
                throw("r0 or l must be specified for the resistance");
            if (!isSet(&rsh))
                throw("rsh must be specified for the resistance");
        }
        // create tape
        IntVector var(2,0);
        var[1] = 1;
        createTape(var);
    }
    else { // pdr==false
        // Set the number of terminals
        setNumTerms(2);
        // Set flags
        setFlags(LINEAR | ONE_REF | TR_FREQ_DOMAIN);
        // calculate the resistor value
        if (isSet(&r0)) {
            r = r0*(1+tc1*(t-tnom)+tc2*(t-tnom)*(t-tnom));
        }

        else {
            if (!isSet(&l))
                throw("r0 or l must be specified for the resistance");
            if (!isSet(&rsh))
                throw("rsh must be specified for the resistance");
            r = rsh*((l-narrow)/(w-narrow))*(1+tc1*(t-tnom)+tc2*(t-tnom)*(t-tnom));
        }
    }
}

```

Part 6 : State Variable definition for a power dissipating resistor (pdr).

Part 7 : State Variable definition for a linear resistor.

```

void resistorT::eval(adoublev& x, adoublev& vp, adoublev& ip)
{
    // x[0]: resistor voltage
    // x[1]: deltatemp in deg. Celsius

    vp[0]=x[0];
    vp[1]=x[1]+tnom+273; //vp[1]==tp[1] in Kelvin
}

```

Part 8 : Output vector assignment : voltage and temperature outputs.



# Bibliography

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