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PowerDEVS. A Tool for Hybrid System Modeling  
and Real Time Simulation.

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**Abstract**

This paper introduces a general purpose software tool for DEVS modeling and simulation oriented to the simulation of hybrid systems. The environment –called *PowerDEVS*– allows defining atomic DEVS models in C++ language that can be then graphically coupled in hierarchical block diagrams to create more complex systems. The environment automatically translates the graphically coupled models into a C++ code which executes the simulation.

A remarkable feature of *PowerDEVS* is the possibility of performing simulations under a real-time operating system (RTAI) synchronizing with a real time clock, which permits the design and automatic implementation of synchronous and asynchronous digital controllers. Combined with its continuous system simulation library, *PowerDEVS* is also an efficient tool for real time simulation of physical systems.

Another feature is the interconnection between *PowerDEVS* and the numerical package Scilab. *PowerDEVS* simulations can make use of Scilab workspace variables and functions, and the results can be sent back to Scilab for further processing and data analysis.

Besides describing the main features of the software tool, the article also illustrates its use with some examples which show its simplicity and efficiency.

**Keywords:** DEVS, Hybrid Systems, Real Time Simulation, Simulation Software

## 1 INTRODUCTION

DEVS [32] is the most general formalism for discrete event system modeling. It allows representing any system provided that it performs a finite number of changes in finite intervals of time. Thus, not only Petri-Nets, State-charts, Event-Graphs and other discrete event languages but also all discrete time systems can be seen as particular cases of DEVS [30, 23].

Taking into account that ordinary differential equations can be approximated by discrete time systems –using numerical integration methods– and that these systems are particular cases of DEVS, it results that DEVS can also approximate continuous systems. Moreover, there are numerical methods –like the Quantized State System (QSS) family [4]– which produce simulation models that cannot be represented in discrete time but only as DEVS models.

Thus, simulation tools based on DEVS are potentially more general than tools for different discrete formalisms, including the popular continuous time ones as Simulink [14] (Matlab), Scicos [3] (Scilab), etc.

Among the existing DEVS simulation tools we should mention DEVS–Java [31], DEVSsim++ [13], DEVS–C++ [6], CD++ [25] and JDEVS [7].

These software tools offer different features which include graphical interfaces and advanced simulation features for general purpose DEVS models and for some specific domains. However, they were developed before the mentioned discrete event methods for numerical integration of ordinary differential equations (ODEs).

These methods produce DEVS models with some very particular features on one hand. In fact, they are usually composed of several atomic DEVS models which belong to two basic classes: quantized integrators and static functions. The different quantized integrators differ from each other in only two parameters: the quantum and the initial state value. Similarly, different static functions only differ in some parameters as the gain, number of inputs, the calculated expression, etc.

On the other hand, these new numerical methods have many potential users outside the DEVS–working community. In strongly discontinuous systems the QSS methods offer solutions which are sensibly better than existing numerical algorithms [18] and they are starting to be used by continuous system simulation people. Unfortunately, most researchers and users of numerical ODE integration methods do not know about DEVS and they would appreciate to use the DEVS–based methods without learning DEVS. Moreover, they would be happier if the software they have to use looks similar to the software they use for conventional numerical methods (Simulink, Dymola, Scicos, etc.).

Taking into account these remarks, a DEVS simulation environment with library handling capabilities and a block–oriented graphical interface like Simulink where parameters can be changed without modifying the blocks and where the atomic DEVS definitions are *hidden* for non–DEVS–users appears as an appropriate solution for hybrid system simulation.

These ideas motivated the development of a general purpose DEVS simulation software oriented to hybrid system simulation. Here, we call *hybrid system* any system involving simultaneous continuous (represented by ODEs) and discrete (represented by DEVS) dynamics.

This software package, called *PowerDEVS* was conceived to be used by DEVS expert programmers as well as by end users who only want to connect predefined blocks and simulate. The tool, initially developed as a Diploma Work [22] in our University, was then re-written and is currently maintained by the Laboratory for System Dynamics and Signal Processing.

*PowerDEVS* is composed of various independent programs:

- The *Model Editor*, which contains the graphical interface allowing the hierarchical block-diagram construction, library managing, parameter se-

lection and other high level definitions as well as providing the linking with the other programs.

- The *Atomic Editor*, which permits editing DEVS atomic models of elementary blocks by defining transition functions, output function, time advance, etc.
- The *Preprocessor*, which translates the model editor files into structure files which contain the coupling structure and the information to build up the simulation code, links the code of the different atomic models according to the corresponding structure file and compiles it to produce a stand alone executable file which simulates the system.
- The *Simulation Interface*, which runs the stand alone executable and permits to vary simulation parameters such as final time, number of simulations to perform, and the simulation mode (normal simulation, timed simulation, step-by-step simulation, etc).
- A running instance of Scilab, which acts as a workspace, where simulation parameters can be read, and results can be exported to. This instance is a modification of Scilab 4.1.2 to support this type of operations.

All these applications were programmed in C++ with the graphical libraries QT, with the exception of the *Model Editor* that was programmed in Visual Basic. We provide compiled versions of these applications for Linux and Windows, and all the source code is available for download too.

*PowerDEVS* also runs under a real time operating system (RTAI[19]) synchronizing the events with a real-time clock with the capability of capturing interrupts at the atomic level. To this purpose, the DEVS simulation scheme of [32] was partially inverted so that simulators of atomic models can send messages to their parents informing about the next event time. In this way, *PowerDEVS* allows the direct implementation of asynchronous DEVS-based Quantized State Controllers [17] on a PC. Besides the Linux and Windows distributions, we developed a modified Kubuntu distribution that includes the RTAI kernel and *PowerDEVS*.

The paper is organized as follows: Section 2 introduces the main concepts used in the rest of the article. Then, Section 3 describes the main features of *PowerDEVS* and Section 4 analyzes its real time characteristics. Finally, Section 5 illustrates the usage of *PowerDEVS* in two examples (including a real time control application) and Section 6 concludes the article.

## 2 BACKGROUND

### 2.1 DEVS

DEVS stands for Discrete Event System specification, a formalism introduced first by Bernard Zeigler [29].

A DEVS model processes an input event trajectory and –according to that trajectory and its own initial conditions– it provokes an output event trajectory.

An *atomic* DEVS model is defined by the following structure:

$$M = (X, Y, S, \delta_{\text{int}}, \delta_{\text{ext}}, \lambda, ta)$$

where:

- $X$  is the set of input event values, i.e., the set of all possible values that an input event can adopt.
- $Y$  is the set of output event values.
- $S$  is the set of state values.
- $\delta_{\text{int}}$ ,  $\delta_{\text{ext}}$ ,  $\lambda$  and  $ta$  are functions which define the system dynamics.

Each possible state  $s$  ( $s \in S$ ) has an associated *Time Advance* computed by the *Time Advance Function*  $ta(s)$  ( $ta(s) : S \rightarrow \mathfrak{R}_0^+$ ). The *Time Advance* is a non-negative real number saying how long the system remains in a given state in absence of input events.

Thus, if the state adopts the value  $s_1$  at time  $t_1$ , after  $ta(s_1)$  units of time (i.e. at time  $ta(s_1) + t_1$ ) the system performs an *internal transition* going to a new state  $s_2$ . The new state is calculated as  $s_2 = \delta_{\text{int}}(s_1)$ . Function  $\delta_{\text{int}}$  ( $\delta_{\text{int}} : S \rightarrow S$ ) is called *Internal Transition Function*.

When the state goes from  $s_1$  to  $s_2$  an output event is produced with value  $y_1 = \lambda(s_1)$ . Function  $\lambda$  ( $\lambda : S \rightarrow Y$ ) is called *Output Function*. In that way, the functions  $ta$ ,  $\delta_{\text{int}}$  and  $\lambda$  define the autonomous behavior of a DEVS model.

When an input event arrives the state changes instantaneously. The new state value depends not only on the input event value but also on the previous state value and the elapsed time since the last transition. If the system arrived to the state  $s_2$  at time  $t_2$  and then an input event arrives at time  $t_2 + e$  with value  $x_1$ , the new state is calculated as  $s_3 = \delta_{\text{ext}}(s_2, e, x_1)$  (note that  $ta(s_2) > e$ ). In this case, we say that the system performs an *external transition*. Function  $\delta_{\text{ext}}$  ( $\delta_{\text{ext}} : S \times \mathfrak{R}_0^+ \times X \rightarrow S$ ) is called *External Transition Function*. No output event is produced during an external transition.

The formalism presented is also called *Classic DEVS* to distinguish it from *Parallel DEVS* [32], which consists in an extension of the previous one conceived to improve the treatment of simultaneous events.

Atomic DEVS models can be coupled. DEVS theory guarantees that the coupling of atomic DEVS models defines new DEVS models (i.e. DEVS is closed under coupling) and then complex systems can be represented by DEVS in a hierarchical way [32].

Coupling in DEVS is usually represented through the use of input and output ports. With these ports, the coupling of DEVS models becomes a simple block-diagram construction. Figure 1 shows a coupled DEVS model  $N$  which is the result of coupling the models  $M_a$  and  $M_b$ .

According to the closure property, the model  $N$  can be used itself as an atomic DEVS and it can be coupled with other atomic or coupled models.

## 2.2 SIMULATING A DEVS MODEL

One of the most important features of DEVS is that very complex models can be simulated in a very easy and efficient way.

The basic idea for the simulation of a coupled DEVS model can be described by the following steps:

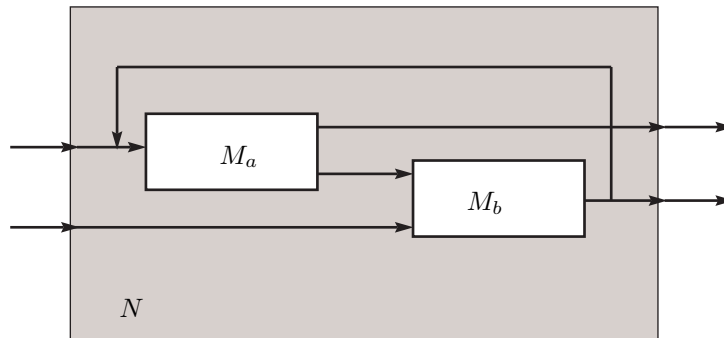


Figure 1: Coupled DEVS model

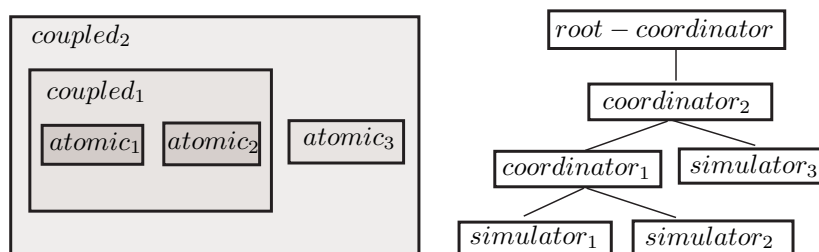


Figure 2: Hierarchical model and simulation scheme

1. Look for the atomic model that, according to its time advance and elapsed time, is the next to perform an internal transition. Call it  $d^*$  and let  $tn$  be the time of the mentioned transition.
2. Advance the simulation time  $t$  to  $t = tn$  and execute the internal transition function of  $d^*$ .
3. Propagate the output event produced by  $d^*$  to all the atomic models connected to it executing the corresponding external transition functions. Then, go back to step 1.

One of the simplest ways to implement these steps is writing a program with a hierarchical structure equivalent to the hierarchical structure of the model to be simulated. This is the method developed in [32] where a routine called *DEVSimulator* is associated to each *atomic DEVS model* and a different routine called *DEVScoordinator* is related to each *coupled DEVS model*. At the top of the hierarchy there is a routine called *DEVScoordinator-root* which manages the global simulation time.

Figure 2 illustrates this idea over a coupled DEVS model.

The simulators and coordinators of consecutive layers communicate with each other with messages. The coordinators send messages to their children so they execute the transition functions. When a simulator executes a transition, it calculates its next state and –when the transition is internal– it sends the

output value to its parent coordinator. In all the cases, the simulator state will coincide with its associated atomic DEVS model state.

When a coordinator executes a transition, it sends messages to some of its children so they execute their corresponding transition functions. When an output event produced by one of its children has to be propagated outside the coupled model, the coordinator sends a message to its own parent coordinator carrying the output value.

Each simulator or coordinator has a local variable  $tn$  which indicates the time when its next internal transition will occur. In the simulators, that variable is calculated using the time advance function of the corresponding atomic model. In the coordinators, it is calculated as the minimum  $tn$  of their children. Thus, the  $tn$  of the coordinator at the top is the time at which the next event of the entire system will occur. Then, the root coordinator only looks at this time, advances the global time  $t$  to this value and then it sends a message to its child so it performs the next transition, and then it repeats this cycle until the end of the simulation.

## 2.3 DEVS AND HYBRID SYSTEMS SIMULATION

Hybrid systems combine discrete and continuous dynamics. While discrete subsystems have a straightforward representation in DEVS, continuous submodels require some type of discretization.

Although DEVS can easily represent the discrete time approximations of continuous systems given by conventional numerical integration methods such as Euler, Runge Kutta, etc., it can also represent the approximations resulting from state quantization. Quantization based integration methods are noticeably efficient to simulate hybrid systems due to their ability to handle discontinuities.

### 2.3.1 QUANTIZATION BASED INTEGRATION

Continuous time systems can be written as set of ordinary differential equations (ODEs):

$$\dot{\mathbf{x}}(t) = \mathbf{f}(\mathbf{x}(t), \mathbf{u}(t)) \quad (1)$$

where  $\mathbf{x} \in \mathfrak{R}^n$  is the state vector and  $\mathbf{u} \in \mathfrak{R}^m$  is a vector of known input functions.

The simulation system (1) requires using numerical integration methods. While conventional integration algorithms are based on time discretization, a new family of numerical methods was developed based on state quantization [4]. The new algorithms, called Quantized State System methods (QSS methods), can approximate ODEs like that of Eq.(1) by DEVS models.

Formally, the first order accurate QSS method approximates Eq.(1) by

$$\dot{\mathbf{x}}(t) = \mathbf{f}(\mathbf{q}(t), \mathbf{v}(t)) \quad (2)$$

where each pair of variables  $q_j$  and  $x_j$  are related by a *hysteretic quantization function*.

The presence of a hysteretic quantization function relating  $q_j(t)$  and  $x_j(t)$  implies that  $q_j(t)$  follows a piecewise constant trajectory that only changes when the difference with  $x_j(t)$  becomes equal to a parameter  $\Delta q_j$  called *quantum*.

The variables  $q_j$  are called *quantized variables*, and can be seen as a piecewise constant approximation of the corresponding state variables  $x_j$ .

Similarly, the components of  $\mathbf{v}(t)$  are piecewise constant approximations of the corresponding components of  $\mathbf{u}(t)$ .

Since the components  $q_j(t)$  and  $v_j(t)$  follow piecewise constant trajectories, it results that the state derivatives  $\dot{x}_j(t)$  also follow piecewise constant trajectories. Then, the state variables  $x_j(t)$  have piecewise linear evolutions.

Each component of Eq.(2) can be thought of as the coupling of two elementary subsystems, a static one,

$$\dot{x}_j(t) = f_j(q_1, \dots, q_n, v_1, \dots, v_m) \quad (3)$$

and a dynamical one

$$q_j(t) = Q_j(x_j(\cdot)) = Q_j\left(\int \dot{x}_j(\tau) d\tau\right) \quad (4)$$

where  $Q_j$  is the hysteretic quantization function (it is not a function of the instantaneous value  $x_j(t)$ , but a functional of the trajectory  $x_j(\cdot)$ ).

Since the components  $v_j(t)$ ,  $q_j(t)$  and  $\dot{x}_j(t)$  are piecewise constant, both subsystems have piecewise constant input and output trajectories that can be represented by sequences of events.

Then, Subsystems (3) and (4) define a relation between their input and output sequences of events. Consequently, equivalent DEVS models can be found for these systems, called *static functions* and *quantized integrators*, respectively [4].

The piecewise constant input trajectories  $v_j(t)$  can be also represented by sequences of events, and *source* DEVS models that generate them can be easily obtained.

Then, the QSS approximation Eq.(2) can be exactly simulated by a DEVS model consisting in the coupling of  $n$  quantized integrators,  $n$  static functions and  $m$  signal sources. The resulting coupled DEVS model looks identical to the block diagram representation of the original system of Eq.(1).

Based on the idea of QSS, a second order accurate method was developed replacing the piecewise constant approximations by piecewise linear ones. The method, called QSS2, can be implemented using DEVS in the same way of QSS. However, the trajectories are now piecewise linear instead of piecewise constant. Thus, the events carry two numbers that indicate the initial value and the slope of each segment. Also, the static functions and quantized integrators are modified with respect to those of QSS so they can take into account the slopes.

Following the idea of QSS2, the third order accurate QSS3 method [16] uses piecewise parabolic trajectories. The family of QSS methods is completed with three methods for stiff systems (Backward QSS and Linearly Implicit QSS of order 1 and 2 [20]) and a method for marginally stable systems (Centered QSS [5]).

### 2.3.2 QSS and HYBRID SYSTEMS

The interaction of the continuous and discrete dynamics occurring in hybrid systems often implies that the right hand side of the system of Eq.(1) that models the continuous subsystems contains discontinuities.

If a numerical integration method performs an integration step that crosses through a discontinuity, the result will have an unacceptable error.

Conventional numerical methods solve this problem by finding the instant of time at which the discontinuity occurs (this is usually called *zero crossing*). Then, they advance the simulation up to that point, and they restart the integration from the new condition (after the event).

Although this idea works fine, it adds some computational cost: zero crossing detection implies performing some iterations and the simulation restart can be also quite expensive. Due to the presence of unbounded iterations, this solution is usually unacceptable in the context of real time simulation [4].

Besides these difficulties, the simulation of a hybrid system requires also the representation and simulation of the remaining discrete subsystems, which in turn calls for the use of a common scheduling algorithm.

The usage of QSS methods on a DEVS simulation engine solves all the mentioned problems. On one hand, DEVS provides the unified framework to represent the discrete and the continuous (quantized) dynamics and to couple them on a single model.

Also, according to the order of the QSS method used, the trajectories are piecewise linear, parabolic or cubic. Thus, the zero crossing detection can be analytically solved, without performing iterations at all.

Although the event associated to a discontinuity must occur at the right time, the methods do not need to restart after that. After all, discontinuities in QSS occur all the time, as the trajectories of  $q_j(t)$  are discontinuous. Thus, for the QSS approximations, a discontinuity has the same effect of a normal step.

Moreover, when a component  $f_j$  of the system (1) contains a discontinuity, the DEVS static function computing  $\dot{x}_j(t)$  will be in charge of detecting the discontinuity and provoking the right event trajectory for the state derivative. In other words, discontinuities are detected and handled locally, without any additional computational cost for the rest of the simulation.

These advantages result in a noticeable simulation speedup with respect to conventional numerical algorithms. In models with rapidly occurring discontinuities such as power electronic systems, the high order QSS2 and QSS3 and LIQSS2 can perform simulations up to 20 times faster than all existing conventional methods [18].

## 2.4 SIMILAR TOOLS

There is a great variety of tools in the field of DEVS simulation. Some are plain DEVS simulators, without real time support, and others are general continuous system simulation tools. Here we describe a few of these tools.

- ADEVS [21] is a C++ library to simulate DEVS based models. Unlike *PowerDEVS* it is based on two DEVS formalism extensions, called *Parallel DEVS* and *Dynamic DEVS* which treat simultaneous events in a different way (*confluent transition*). ADEVS is a library, but it does not have a GUI for model coupling and the model has to be described alphanumerically.
- CD++[24] is a DEVS simulation tool developed by Gabriel Wainer's group. It is based on another DEVS extension, called *Cell-DEVS*, which merges DEVS and cellular automata. It has support for real time simulation.



- DEVSJava[32, 11] is a DEVS simulation tool developed by Bernard Zeigler and Hessam Sarjoughian. It has interesting features such as changing the model structure at run time (or simulation time). Also, it offers a hierarchical view of the model structure. It has support for real time simulation in a *best shot* way, because it is written in Java and runs under non-real-time Operating Systems.
- Matlab Real Time Workshop[12] is a toolbox of Simulink. Simulink is a tool for modeling and simulation of continuous systems. It has a GUI in which models can be described in a Block Diagram fashion. It supports generating C++ code for running the simulation under a variety of RTOS. The Real Time Workshop (being a part of Matlab) is a proprietary software of *The MathWorks* and its use and distribution is restricted by its license.

### 3 POWERDEVS

In this section we describe the main components and features of the software developed.

#### 3.1 POWERDEVS COMPONENTS

As we mentioned, *PowerDEVS* is composed of various independent programs: the *model editor*, the *atomic editor*, the *preprocessor* and the *simulation interface* and a workspace corresponding to a *Scilab* instance.

In this section we describe each of this tools and how they interact with each other.

##### 3.1.1 THE MODEL EDITOR

The Model Editor is –from a user point of view– the *main* program of PowerDEVS as it provides the graphical interface and the link with the rest of the applications.

Besides building and managing models and libraries, it permits launching a simulation (by invoking the Preprocessor) and editing elementary blocks up to the atomic model definitions (by invoking the Atomic Editor).

The Model Editor main window (Fig. 3) allows the user to create and open models and libraries. It also permits exploring the libraries and dragging blocks from the libraries to the models.

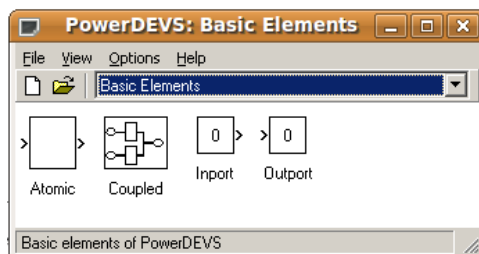


Figure 3: Model Editor main window.

There are also some advanced features which can be managed from the main window like setting which are the *active* libraries (i.e. which libraries are shown when exploring), and configuring the tool bars and menu to invoke new external applications.

Models and libraries can be edited in a model window with the *open* and *new model* commands. Figure 4 shows a model window with a model composed by five sub-models.

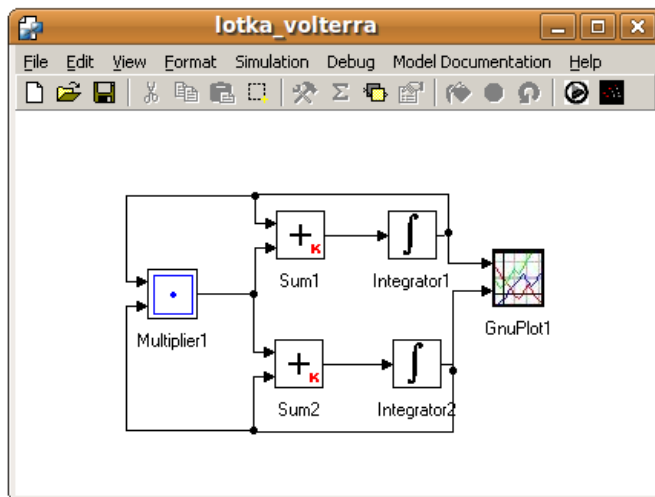


Figure 4: Model Window.

The Model Windows provide all the typical graphical edition facilities so that blocks can be copied, resized, rotated, etc. while the connections can be directly drawn between different ports.

From the edit menu (or with the right button) it is possible to edit the features of each block, no matter if it corresponds to a coupled or an atomic model.

The Block Edition Window (Fig. 5) allows to configure the graphic appearance of the block, to choose the block parameters and –in the case of atomic models– to select the file which contains the associated code with the DEVS model definitions.

The block parameters are defined and selected in the block edition windows. After being defined, their values can be changed by double-clicking on the block (Fig. 6). Thus, when we take predefined blocks from the libraries, we can change the parameter values without editing them. As we shall see in Section 3.1.3, the values of these parameters are passed to the corresponding DEVS atomic or coupled models.

Coupled models do not have an associated code, but they have some extra features which can be modified from the block edition window and the edit menu (the internal priorities and the order of the input and output ports, for instance).

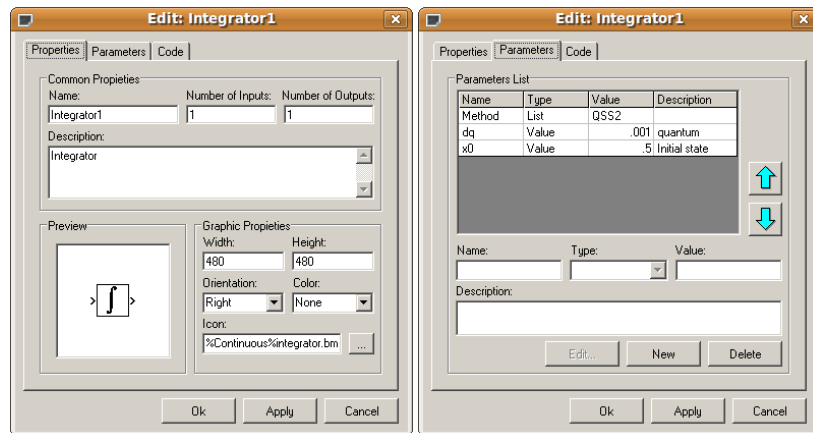


Figure 5: Block Edition Window.

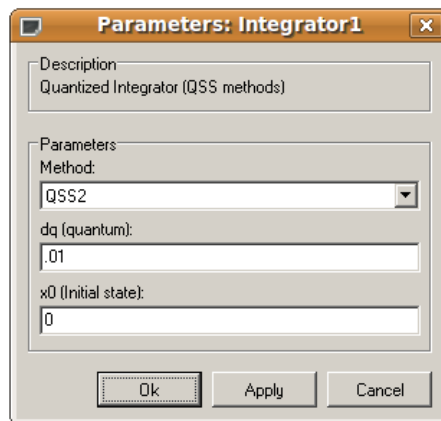


Figure 6: Changing parameter values.

### 3.1.2 THE ATOMIC EDITOR

The *Atomic Editor* facilitates the edition of the C++ code corresponding to each atomic DEVS model.

It can be invoked from the Block Edition Window to edit an existing code or to create a new one. It can be also run directly from the OS since it is a stand alone application. The *Atomic Editor* main window is shown in Figure 7.

Using the atomic editor, the user only has to define the variables which form the state and the output of the DEVS model and the variables which represent the parameters of the system. After that, the C++ code of the time advance, transition and output functions must be placed in the corresponding windows. There are two additional windows (init and exit) where the user can also add a piece of code that is executed before the simulation starts (to set initial states and parameters, for instance) and a piece of code that is executed at the end of the simulation (to close some open files, for instance). When the model is saved, the code is automatically completed and stored in the corresponding .cpp and

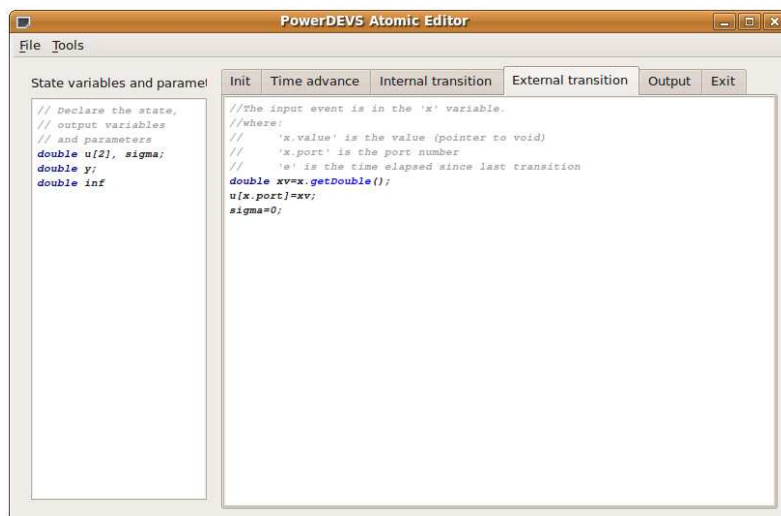


Figure 7: Atomic Editor main window.

.h files.

Besides facilitating the programming, the Atomic Editor was designed to give the user the possibility of writing a code which is very similar to the DEVS model definition. All the rest of the job –related to simulation and implementation issues– is automatically performed by the program.

Let us illustrate this fact with a simple DEVS model. Consider for instance a system which calculates a static function  $f(u_0, u_1) = u_0 - u_1$  with  $u_0$  and  $u_1$  being real-valued piecewise constant trajectories. If we represent those trajectories by sequences of events –as it is done in QSS–methods– we can build the following atomic DEVS model:

$$\begin{aligned}
 M &= (X, Y, S, \delta_{\text{int}}, \delta_{\text{ext}}, \lambda, ta), \text{ where} \\
 S &= \mathfrak{R}^2 \times \mathfrak{R}^+ \\
 \delta_{\text{int}}(s) &= \delta_{\text{int}}(u_0, u_1, \sigma) = (u_0, u_1, \infty) \\
 \delta_{\text{ext}}(s, e, x) &= \delta_{\text{ext}}(u_0, u_1, \sigma, e, x_v, p) = \tilde{s} \\
 \lambda(s) &= \lambda(u_0, u_1, \sigma) = (u_0 - u_1, 0) \\
 ta(s) &= ta(u_0, u_1, \sigma) = \sigma
 \end{aligned}$$

with

$$\tilde{s} = \begin{cases} (x_v, u_1, 0) & \text{if } p = 0 \\ (u_0, x_v, 0) & \text{otherwise} \end{cases}$$

We use integer numbers from 0 to  $n - 1$  to denote the input and output ports (because PowerDEVS does so).

This DEVS model translated into PowerDEVS has the following code (at the Atomic Editor Fig. 7):

**ATOMIC MODEL STATIC1**  
**State Variables and Parameters:**

```

float u[2],sigma; //states
float y; //output
float inf; //parameter

```

**Init Function:**

```

inf = 1e10;
u[0] = 0;
u[1] = 0;
sigma = inf;
y = 0;

```

**Time Advance Function:**

```

return sigma;

```

**Internal Transition Function:**

```

sigma=inf;

```

**External Transition Function:**

```

float xv;
xv=(float*)(x.value);
u[x.port] = xv;
sigma = 0;

```

**Output Function:**

```

y = u[0] - u[1];
return Event(&y,0);

```

It is clear that the translation from the atomic DEVS into the PowerDEVS code is almost straightforward.

With that code, the Atomic Editor automatically produces the .cpp and .h files which are then used by the Preprocessor to generate the simulation of the whole system.

### 3.1.3 THE PREPROCESSOR

The *Preprocessor* takes a .pdm (or .pds) file produced by the *model editor* and produces the simulation program.

It basically translates the .pdm file into a header .h file (called *model.h*) which binds the simulators and coordinators according to the coupling structure passing also the block parameters.

The preprocessor also produces a makefile (*Makefile.include*) which is then invoked to compile and generate the program which implements the simulation (that program is called *model*).

As we mentioned before, the *Preprocessor* can be invoked in a transparent way using the *Quick Simulation* command. Anyway, it can be also called from the command line (it is also a stand alone application).

### 3.1.4 THE SIMULATION INTERFACE

The generated program *model*, when executed, simulates the associated DEVS model.

*PowerDEVS* provides a graphical interface for running the simulation (Fig. 8).

The interface also allows to change some parameters to set up the experiment:

- Final Time: tells for how long to simulate the model.

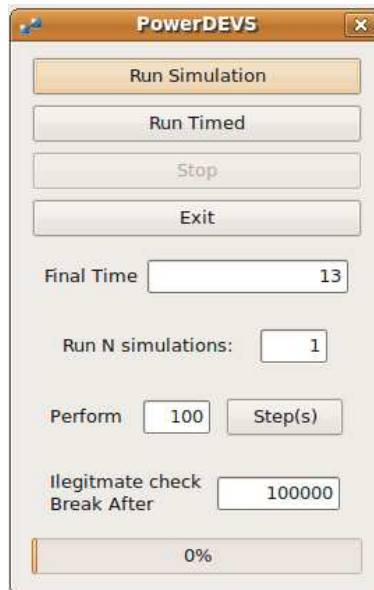


Figure 8: Simulation Interface

- Simulations to run: multiple simulation runs can be executed at once. This can be useful when statistics from simulation results are to be calculated.
- Illegitimate check break: *PowerDEVS* stops the simulation if the time does not advance after a selected number of steps. This avoids hang-ups in illegitimate models.
- Simulate step-by-step: the simulation can be advanced performing one step (or many) at the time, and results can be analyzed in between.
- Synchronize time: The simulation can be run synchronized with the real clock (with the precision of the underlying OS).

Also, the *model* program can be invoked from the command line, and executed in an interactive shell. All the simulation parameters can be changed in the same way they are changed from the graphical interface.

## 3.2 INTERNAL IMPLEMENTATION

Having described the main functional aspects of *PowerDEVS*, we shall now explain the way in which the simulation algorithm described in Section 2.2 is implemented inside the program that executes the simulation.

As *PowerDEVS* performs an object oriented simulation, we shall start describing the internal class structure.

Atomic models with the same associated code belong to a particular class (defined by that code). For instance, the *Integrator* atomic models in the model of Fig. 4 belong to the *Integrator* class, defined in the files *integrator.h* and *integrator.cpp* which contain the code associated to that atomic model.

When the code of an atomic model is written using the Atomic Editor, the code related to the class definition is automatically generated.

All the atomic classes are derived from the *simulator* class. The simulator class is an abstract class which acts as an interface to deal with different atomic model implementations. The variables representing the state of the model and the functions that operate on it (time advance, transition and output functions) are member variables and methods.

The external transition and output functions of the simulator class receive and return respectively objects belonging to the *Event* class. Events have the following properties:

- *Event.port*: is an integer number indicating the input or output port where the event is received or sent.
- *Event.value*: is a pointer to void. That way, the values carried by the events can belong to arbitrary types.
- *Event.realTimeMode*: is an integer that can take the following values: 0 (indicating that the event is not synchronized with the real time), 1 (with normal synchronization) and 2 (with precise synchronization). When an event has its mode set to 1 or 2 (the difference between them will be explained in Section 4.2.1), the simulation engine waits until the physical time reaches the simulation time to propagate it.

To provide the capability of initializing and stopping devices that interact with the hardware, the *simulator* class has two extra methods which are not included in the definition of [32]. As we already mentioned, for real-time simulation purposes *PowerDEVS* also inverts the time managing features. Thus, when an atomic model receives an interrupt request from an external device, it informs the change in its time to next event (*tn*) to its parent.

The hierarchical coupling structure is implemented by the *coupling* class. This class is similar to the coordinator in Fig. 2. Each object of this class is associated to a coupled DEVS model and it contains a list of references to the corresponding connections, atomic and coupled models.

The difference with the coordinator is that, following the *simulator* class behavior, the *coupling* objects are able to receive the messages coming from their children notifying changes in their time to next event. Similarly, the *coupling* objects have also the possibility of informing their own parent about changes in their *tn*. Coherently with the closure property, the *coupling* class is derived from the *simulator* class.

The *coupling* and *simulator* objects also contain objects belonging to the classes *connection* and *event* (the first one is only in the *coupling* class). The *connection* class is formed by four integer numbers representing two pairs of models and ports. The *event* objects are formed by a pointer to void and an integer (identifying the input or output port). Thus *PowerDEVS* models can produce output events which belong to different types.

Having defined the model structure in terms of components and functionality, we shall now describe the framework developed to actually execute the simulation.

The *root-simulator* class is in charge of running the simulation. Basically, this class manages the simulation advance interacting with the object represent-

ing the coupling at the top of the structure. Thus, the *root-simulator* plays the role of the root-coordinator in Fig. 2.

Before following the description we should mention that the behavior of the top coupling object has a small difference with the others. In terms of the simulation execution, there is no coupling above to notify the timing changes but these changes should be notified directly to the *root-simulator*. Thus, a new class denominated *root-coupling* has been introduced. This class, derived from the *coupling* class, has the same functionality as the latter but it differs in the hierarchical superior object.

Finally, the *main* class of the simulation program is called *model*. This class provides the interactive shell for interacting with the user (or with the graphical simulation interface of Fig.8 ) and running the simulation. This class talks directly with the *root-simulator* allowing the user to start the simulation in different modes.

The objects which form the coupling structure are instantiated at the *model.h* file. This file is automatically generated by the Preprocessor and it declares the simulators, couplings and connections according to the structure file.

In fact, the only difference between the codes that execute the simulation of two different models is in that file (*model.h*). Obviously, the fact that both files are different also implies that they may include different *simulators* corresponding to different atomic models.

### 3.3 INTERCONNECTION WITH SCILAB

Scilab [8] is a numerical computational package developed by the Institut National de Recherche en Informatique et Automatique (INRIA) and it is released under the GPL license. Scilab is an Open Source alternative to the widely used software Matlab.

Scilab contains an interactive interface where the user can define variables and matrices and perform complex operations between them. It also has a programming language that can be used to define new functions and algorithms, and some graphical tools to plot data. The Scilab distribution includes several toolboxes (i.e., set of functions developed by different people) to solve problems related to Linear Algebra, Signal Processing, Automatic Control, Optimization, Filtering Design, etc.

To make the communication between *PowerDEVS* and Scilab, some modifications to the source code of Scilab were made. These modifications open a "back door" into Scilab's workspace, where variables can be read or written from *PowerDEVS*. The communication is performed through UDP messages over the port 27015 in a client-server model, with Scilab acting as server and *PowerDEVS* as client.

#### 3.3.1 THE SCILAB SIDE

As Scilab is an open-source tool, all its source code is available online to view, edit and enhance. We used Scilab 4.1.2 version and made modifications in both, Linux and Windows source versions. These modifications were made in the files *routines/wsci/wScilex/wScilex.c* (for Windows) and *routines/xsci/x\_main.c* (for Linux).



As the idea was to use Scilab as *PowerDEVS workspace* –that is, *PowerDEVS* runs the simulation and Scilab responds to *PowerDEVS* commands– it was necessary to modify the normal behavior of Scilab in order to establish the communication.

To keep running the Scilab GUI, while communicating with *PowerDEVS* a new thread is created in Scilab address space. This new thread is in charge of receiving, executing, and responding to *PowerDEVS* requests.

The actual communication between *PowerDEVS* and this Scilab thread was implemented in a Client-Server model over UDP. Under this architecture, Scilab (acting as the server) waits for requests on the UDP port 27015 and *PowerDEVS* sends requests to that port (see Fig. 9).

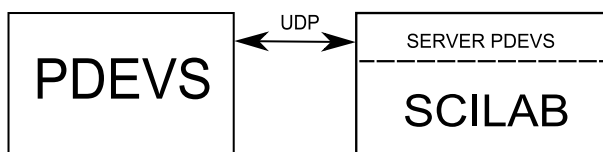


Figure 9: *PowerDEVS* –Scilab communication

The *PowerDEVS* requests sent to Scilab consist of string representations of the commands to execute in Scilab workspace (for example “freq=pi\*345”). Each request starts with a character indicating if Scilab must notify *PowerDEVS* once the command execution is finished.

The Scilab notification to *PowerDEVS* (when the first character indicates so) consists of a UDP message containing the value of the “ans” variable in Scilab workspace. This permits –indirectly– to evaluate any Scilab expression or variable from *PowerDEVS* sending the correct command sequence.

### 3.3.2 THE *PowerDEVS* SIDE

The *PowerDEVS* simulation engine has a set of services to communicate with Scilab. These services can be called from any atomic model written with the atomic editor:

```
void  getAns(double *ans, int r, int c);
void  putScilabVar(char *varname, double v);
double  getScilabVar(char *varname);
double  executeScilabJob(char *job, bool blocking);
```

The function `getAns(double *ans, int r, int c)` returns the answer of the last calculation in the Scilab workspace (i.e., the “ans” variable). The result is put in the double **pointed** by `ans`. The `r` and `c` arguments indicate the size of the result to be retrieved (Scilab works with matrices and so does the *PowerDEVS* interface). If a scalar value is to be retrieved `r` and `c` must be 1.

The functions `putScilabVar(char *varname, double v)` and `getScilabVar(char *varname)` are the basic methods to interact with Scilab variables. `putScilabVar` creates (or updates) the Scilab variables named `varname` with value `v`. On the other hand, `getScilabVar` reads the variable named `varname` from Scilab workspace. If the variable does not exist this function returns 0.0 and writes a warning message in the simulation log.

Finally, the function `executeScilabJob(char *job, bool blocking)` runs the command `job` in the Scilab workspace. It works just like when one types the sentence in the Scilab command window. The second argument `blocking` indicates if the function should return immediately or should wait for the command to finish. The function returns the “ans” variable (like `getAns`) which –according to the command– can represent the result of the executed command.

### 3.4 LIBRARY

The *PowerDEVS* distribution comes with a set of atomic and coupled models already programmed, that can be used to build new models. These basic models form the *PowerDEVS* library (any advanced user can easily extend the library including his own developed models).

The library is divided into the following categories:

**Basic Elements:** This is the only *PowerDEVS* core library. Based on these models, all the other libraries were developed. It contains four basic models, an atomic (a skeleton for all the atomics), a coupled (an empty coupled model), and two special objects called `inport` and `outport`. The last two objects represent external input and output interfaces for coupled models.

**Continuous:** This library contains models that process continuous signals based on the QSS methods to simulate continuous systems<sup>1</sup>. It includes integrators, gains, non-linear static functions, multipliers, etc.

**Discrete:** This library contains models for simulating discrete time systems.

**Hybrid:** Under this category, a set of models are included combining continuous and discrete features to simulate hybrid systems: quantizers, switches, comparators, samplers, etc.

**Real Time:** This library contains specific blocks to make use of different real time features of *PowerDEVS*. They are described in Section 4.3.

**Source:** Here we can find several sources for continuous time signals approximated by QSS methods, like sine waves, ramps, pulses, square waves, etc.

**Sink:** This library contains different sink models where simulation results are sent for visualization or future processing. Some models included in this library are: `GnuPlot` (which plots its input signals), `ToDisk` (which writes its input signals to a CVS file), `toWorkspace` (that writes in Scilab workspace the received signal), etc.

All the models in the library accept Scilab expressions as parameter values. For instance, one could use “`freq*2+0.5`” as the gain parameter in a gain block, and this expression will be evaluated in Scilab workspace (in this example `freq` must be a defined Scilab variable).

---

<sup>1</sup>The Continuous library was conceived to simulate Ordinary Differential Equations. Differential Algebraic Equations can be simulated with the use of the Implicit Function Block, that implements the methodology developed in [15]. If instead of using that block an algebraic loop is included in the model, the simulation will result illegitimate.

## 4 POWERDEVS IN REALTIME

As we anticipated earlier, *PowerDEVS* was extended to run on a real time operating system. For some reasons that will be explained soon in this section, we choose Linux RTAI as the RTOS for *PowerDEVS*.

The simulation engine of *PowerDEVS* is an implementation in C++ of the simulator described in Section 2.2. As RTAI is an extension of Linux, in principle, the *PowerDEVS* simulations (that run fine under Linux) should also run under RTAI.

Although this is true, the goal of running a simulation under RTAI is making use of its services to ensure real time performance. So several modifications were made in the *PowerDEVS* simulation engine which enable the user to easily access basic services like synchronizing events, capturing and handling hardware interrupts, obtaining file support and performing real time measurement.

Also, the *PowerDEVS* library was extended with a set of blocks to make use of these services in a simple drag-and-drop way.

This section, after introducing some concepts related to real time systems and real time simulation, describes the extensions made to the simulation engine, the new library blocks and some experiments done to measure different performance indicators of *PowerDEVS* in real time.

### 4.1 REAL-TIME SYSTEMS & RTOS

#### 4.1.1 REAL-TIME SYSTEMS

A real-time system is a system in which computations are subject to *real-time constraints*. That is, responses to stimuli have a dead-line that must be met, regardless of system load, in order for the system to be considered correct.

Real-Time systems can be classified in two categories:

**Hard real time systems:** in which the completion of a computation after its deadline is considered useless, and may cause critical failures in the system.

**Soft real-time systems:** in which a missed deadline can be accepted, or ignored or even corrected.

#### 4.1.2 REAL-TIME OPERATING SYSTEMS (RTOS)

Real-Time Operating Systems (RTOS) are a special class of Operating Systems (OS) which provide the basic tools and services to implement systems with time constraints.

This kind of OS should be expressive enough to represent the time constraints of each task, the way these tasks communicate, and must have some control over the low-level hardware of the computer (memory, interrupts, ports).

Some commonly used RTOS are QNX [10], VxWorks [26], Ddrops [9], RT-Linux [1], RTAI [19], etc..

Among them, we decided working using RTAI (RealTime Application Interface) because it consists of an extension of the Linux Kernel [2] that permits running all the software that runs on Linux. This was a crucial requirement in the context of this work since we wanted to run not only *PowerDEVS*, but also the external applications that communicate with it (GNUPlot and Scilab).

Also, the fact that RTAI is distributed under the GPL License allows us to freely distribute the real time version of PowerDEVS included with the OS installer.

### 4.1.3 RTAI

RTAI is a RTOS that supports several architectures (i386, PowerPC, ARM). RTAI is an extension of the Linux kernel to support real time tasks to run concurrently with Linux processes. This extension uses a method to enable this, first used in RT-Linux [1].

Linux kernel is not itself a RTOS. It does not provide real time services, and in some parts of the kernel, interrupts are disabled as a method of synchronization (to update internal structures in an atomic way). These periods of time where interrupts are disabled, lead to a scenario where the response time of the system is unknown, and time deadlines could be missed.

To avoid this problem RTAI inserts an abstraction layer beneath the Linux kernel. In this way, Linux never disables the *real* hardware interrupts. The Linux kernel is run under another micro-kernel (RTAI + Adeos <sup>2</sup>) as user processes. All hardware interrupts are captured by this micro-kernel and forwarded to Linux (if Linux has interrupts enabled).

Another problem running real time tasks under Linux is that the Linux scheduler can take control of the processor from any running process without restrictions. This is unacceptable in a real time system. Thus, in RTAI real time tasks are run in the micro-kernel, without any supervision of Linux. Moreover, these processes are not seen by the Linux scheduler.

Real time tasks are managed by the RTAI scheduler. Clearly, there are two different kinds of running processes, Linux processes and RTAI processes. RTAI processes cannot make use of Linux services (such as File system) and vice versa. To avoid this problem, RTAI offers various IPC mechanisms (Pipes, Mailboxes, Shared Memory, etc.).

RTAI provides the user with some basic services to implement real-time systems:

**Deterministic interrupt response time:** An interrupt is an external event to the system. The response time to this event varies from computer to computer, but RTAI guarantees a boundary (on each computer), which is necessary to communicate with external hardware, for example a data acquisition board.

**Inter Process Communication (IPC):** RTAI offers various methods of IPC, such as semaphore, pipes, mailbox and shared memory. These IPC mechanisms can be used to connect processes running in real time with normal Linux processes or vice versa.

**High precision timers:** When developing real time systems, the time handling accuracy is very important. RTAI offers clocks and timers with ns ( $1e^{-9}$  s) precision. On i386 architectures, these timers use the processor's time stamp, therefore, they are very precise.

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<sup>2</sup>Adeos [27, 28] is the abstraction layer used in RTAI. Adeos implements the concept of *interrupt domain*. <http://home.gna.org/adeos/>

**Interrupt Handling** RTAI allows the user to capture hardware interrupts and treat them with custom user handlers. Normal Operating Systems (such as Linux, Windows) hide hardware interrupts from the user, making the development of communication with external hardware a bit more difficult (you have to write a kernel driver). As said before, the response time is bounded.

RTAI tasks can be run in two different modes

**Loadable kernel module:** in this mode executable files are not stand alone programs but loadable kernel modules. The RTAI process is *inserted* in the running Linux kernel and from then on it switches to the RTAI scheduler. Running in kernel mode (as modules do) gives the process certain privileges and rights that normal user processes do not have (accessing kernel objects, writing/reading ports, etc).

**LXRT:** RTAI offers a library called LXRT for developing real time systems in user space. LXRT is a “bridge” between Linux user mode and RTAI. LXRT brings to the user space all RTAI services. LXRT works with a concept called *angel process*. LXRT creates a RTAI counterpart (the angel process) in charge of executing all the real time services on behalf of the user space process.

Having described the main features of RTAI, we shall now focus on the extensions made to *PowerDEVS* .

## 4.2 *PowerDEVS* ENGINE SUPPORT

Four new modules were added to the simulation engine to make use of the real time services provided by RTAI. The modules consist in new functions that can be used by any atomic model.

### 4.2.1 SYNCHRONIZATION MODULE

This module is the cornerstone of *PowerDEVS* in real time. It allows the simulation engine to synchronize the events with the RTAI real clock.

RTAI has two methods of synchronizing, both with ns precision:

- *rt\_sleep*: waits for an amount of time, releasing the processor.
- *rt\_busy\_sleep*: waits for an amount of time by busy-waiting, that is, burning CPU cycles until the wait is over. In this mode, a more precise synchronization is obtained.

Based on these two methods, the *PowerDEVS* simulation engine implements the function `waitFor(unsigned long int nanoseconds, int mode)` which waits the specified time in one of two modes: *Normal* or *Precise*.

This function has the following logic:

- If the amount of time to wait is *too* small (smaller than a constant that defines the synchronization accuracy), the sentence is ignored.
- If the function is called in *Normal* mode, the non-busy *rt\_sleep* is invoked (where the CPU is released).

- If the function is called in *Precise* mode, the behavior depends on the time to wait:
  - If the period is less than a given constant (which defines the accuracy of the non-busy *rt\_sleep*), a busy waiting is performed invoking *rt\_busy\_sleep*.
  - Otherwise, the waiting is split in two periods. The first period is computed as the whole period minus twice the non-busy accuracy, and it is performed in a non-busy way invoking *rt\_sleep*. After finishing this period, the time is measured again (to eliminate the error introduced by the non-busy waiting) and the remaining waiting time is done with a busy-waiting. This way, the synchronization has the accuracy of a busy waiting, but the processor is released most of the time.

The function `waitFor` can be invoked by any atomic model, but it is automatically called by the simulation engine when an output event has its `realTimeMode` property (see Section 3.2) set to 1 or 2.

This way, in the same model, some events can be propagated immediately (when their `realTimeMode` is set to 0) and others can be synchronized either in normal or precise mode. This gives the user the possibility of choosing which events must be synchronized (typically those that involve communication with external hardware) and which should be computed as fast as possible (those involving intermediate calculations).

#### 4.2.2 INTERRUPT HANDLING MODULE

This module is in charge of managing the interaction between the *PowerDEVS* simulation engine and the hardware interrupts.

Any atomic can express interest in a specific hardware interrupt by calling:

```
requestIRQ(int irq);
```

When a hardware interrupt `irq` occurs, the simulation engine sends an external transition message to all the atomic models that have expressed interest in that interrupt. The message arrives to a non-existing port (denoted -1).

Since these atomic models will possibly change their time advances in the external transition, the engine automatically transmits from the bottom to the top of the hierarchy about these changes. This is the mechanism that we mentioned in Section 2.2, and it is the only modification of the abstract simulator of [32] introduced by *PowerDEVS*.

#### 4.2.3 FILE SUPPORT MODULE

When a task is running under RTAI, it can not make use of Linux services (such as Linux File System). This is because a system call to the Linux kernel does not have a deterministic response time, so, a system call could lead to missing a deadline in the real time system. In fact, RTAI tasks are not scheduled by the Linux scheduler which does not know they exist. Thus, to enable the simulations to use files, a File Support Module was developed. This module has two parts:

**Real Time Interface:** It is a real time task (running under RTAI) that accepts the requests of the real time simulation (like open, read, write files). It communicates with a program running under Linux user space (which has access to the Linux File System) using a Real Time FIFO (one of the IPC provided by RTAI).

**User space Angel:** It is a normal Linux program that accepts commands from the Real Time Interface. This program (which is launched together with the simulation) makes all the system calls on behalf of the simulation.

The interface provides similar functions to those of the c library (stdio) that can be directly used by any atomic model.

#### 4.2.4 REALTIME MEASUREMENT MODULE

This module allows the atomics models to ask for the value of the real time clock. This is useful, for instance, when a user wants to change the behaviour of an atomic model depending on the difference (in either way) between the real time and the simulated time (to handle overruns or to compute more precisely if enough time is available).

### 4.3 REALTIME LIBRARY

To make use of all the real time features in a simple way, the *PowerDEVS* library was extended to include several atomic blocks that can be included directly in the models.

**RTWait:** This atomic block receives events and outputs them synchronized with the real time clock. In this way, a normal model (i.e. non-real time) can be transformed into a real time model by inserting these blocks on the connections carrying events that need to be synchronized. The block has a parameter to choose what kind of synchronization must be made, *precise* or *normal* (see 4.2.1).

**RTClock:** When this block receives an event (no matter what the value of the event is), it emits an event whose value is equal to the real time clock value.

**ToDisk:** This model writes the input signal in a CSV(Comma Separated Values) file. It makes use of the File Support Module to run in real time.

**IRQDetector:** This model is the user interface for the Interrupt Handling Module. It emits an event whenever the hardware interrupt occurs. From the point of view of the DEVS formalism (and from *PowerDEVS* too) this is a passive model (its time advance is  $\infty$ ), but when the associated interrupt is triggered, the model changes its own time advance to 0 (and the engine automatically notifies its parent of this change). The block has a parameter to indicate which interrupt it waits for.

#### 4.4 REAL TIME PERFORMANCE

One of the most important features to characterize a real time system is the *error* (or latency) of synchronization.

We conducted several experiments to see how the system responds (in relationship to the load of the system). All the experiments were run on a PC AMD Athlon 1.8 *GHz*.

First, we did the following experiment: we simulated a source block which generates a QSS-approximation of a sine wave at 440Hz (this model generates 20 events per cycle, to a total of 8800 events per second). The model was simulated in real time measuring the maximum and average latency. Also the two different modes (see Sect. 4.2.1) of waiting were tested. The following results were obtained:

Table 1: Synchronization error with low load

Wait	Maximum	Average	Overruns
Normal	4800 <i>ns</i>	1500 <i>ns</i>	-
Precise	450 <i>ns</i>	180 <i>ns</i>	-

Next, a simulation of a more complex model was performed: a PI control of a DC motor using Pulse With Modulation (PWM). In this example, several simulations were run varying the PWM carrier from 1000 to 20000 Hz:

Table 2: Synchronization error with varying load

f ( <i>Hz</i> )	Wait Mode	Maximum	Average.	Overruns
1000	Normal	5786 <i>ns</i>	1512 <i>ns</i>	-
	Precise	1330 <i>ns</i>	180 <i>ns</i>	-
15000	Normal	5622 <i>ns</i>	1622 <i>ns</i>	372 / 19905 ev.
	Precise	1000 <i>ns</i>	512 <i>ns</i>	305 / 19905 ev.
17000	Normal	4547 <i>ns</i>	1648 <i>ns</i>	6119 / 17616 ev.
	Precise	973 <i>ns</i>	454 <i>ns</i>	5924 / 17616 ev.
20000	Normal	0 <i>ns</i>	0 <i>ns</i>	18292 / 18292 ev.
	Precise	0 <i>ns</i>	0 <i>ns</i>	18292 / 18292 ev.

As can be seen in table 2, using a frequency of 1000 *Hz* there are no *overruns*. With frequencies between 15000 *Hz* and 17000 *Hz*, there are more *overruns* and the Linux system (not the real-time system) experiences some delays, which indicates that the system load is high.

With a frequency of 20000 *Hz* the system stops responding (the Linux part) and we see that all events are emitted after their time deadline.

This last case reflects a limitation of the hardware platform, not a limitation of the system synchronization. What happens with a frequency of 20000Hz is that the hardware cannot complete the calculations in time.



## 4.5 INTERRUPT LATENCY

To measure the interrupt latency the following procedure was used.

A *PowerDEVS* model was created (Fig. 10) which generates hardware interrupts and then captures back these interrupts, measuring the period of time between the generation of the signal provoking the interrupt and the capture of the interrupt by the handler.

For that purpose, we used the PC Parallel port. One of the pins on the parallel port (*STO*) is in charge of generating interrupts. This pin was wired to a data pin, thus generating a hardware interrupt every time that the data pin goes from a low state (0 V) to a high state (5V). The model stores the time at which a 1 is written to the data pin (high state), and the time at which the interrupt handler of the parallel port is run. The difference between these two times is an upper bound of the interrupt latency of the system (in fact, that difference includes also the time needed to write on the parallel port and the electrical transients).

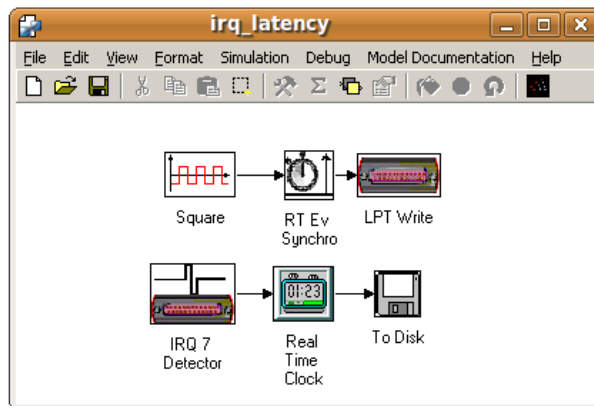


Figure 10: Interrupt Latency tester model

Running this experiment on the same hardware (PC AMD 1.8Ghz) we obtained an upper bound on the interrupt latency of about  $20\mu\text{s}$ .

## 5 EXAMPLES

This section describes two examples that show different features of *PowerDEVS*.

### 5.1 RIPPLE VS FREQUENCY IN A BUCK CIRCUIT

Figure 11 represents a DC-DC converter circuit known as *Buck Circuit*.

The presence of the switch introduces hybrid behavior to the system.

The goal of the experiments is to analyze the dependence of the ripple amplitude on the switching frequency. In this example we used the following parameters  $L_1 = L_2 = 0.1\text{mH}$ ,  $C = 20\mu\text{F}$ ,  $Rl = 10\Omega$  and  $U = 12\text{V}$ .

Figure 12 shows the *PowerDEVS* model of the circuit, made entirely with atomic blocks from the *PowerDEVS* library. The switch is commanded by a

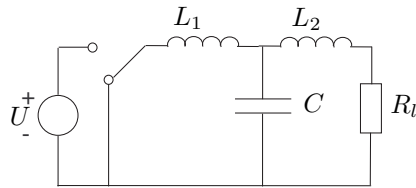


Figure 11: Buck Circuit

PWM signal. The remaining blocks implement source, static functions and integrators based on QSS methods.

The frequency of the triangular carrier was chosen in a way that increments 2000Hz in each simulation. The “RunScilab Job” block increments the Scilab variable  $n$  after each simulation and calculates the ripple amplitude in steady state. This amplitude is saved in the array  $u(n)$ .

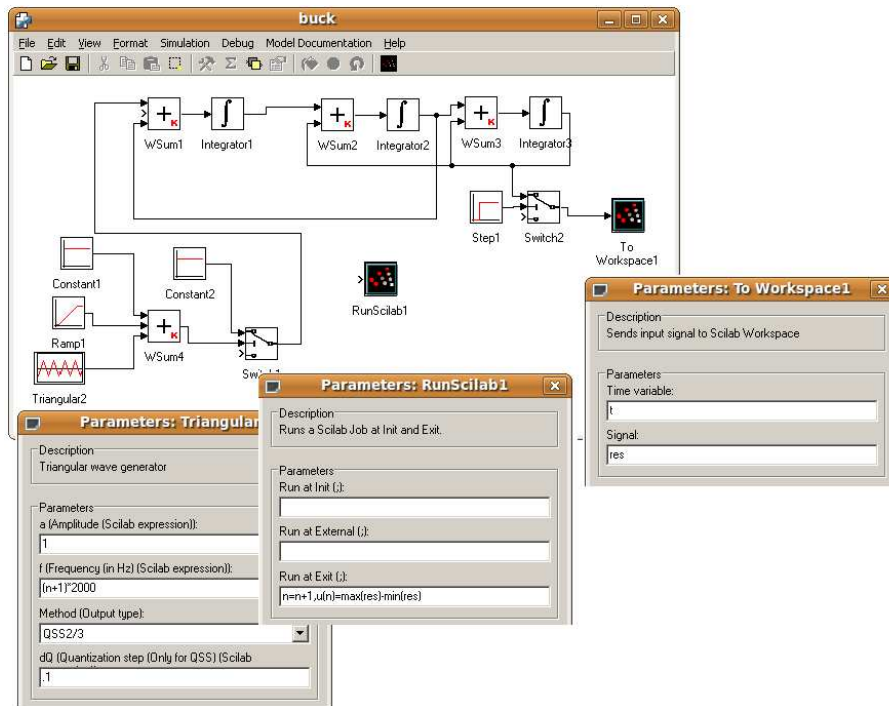


Figure 12: Buck Circuit Model

After 100 simulations (in which the frequency goes from 2000 to 200000 Hz), we can plot the results directly in Scilab (see Fig. 13).

It must be noticed that 100 simulations of this hybrid system (with very quick commutations) were run, and thanks to the efficiency of the QSS methods to treat this kind of problems, the experiment only took about 13 seconds (while in Matlab/Simulink it takes about a minute).

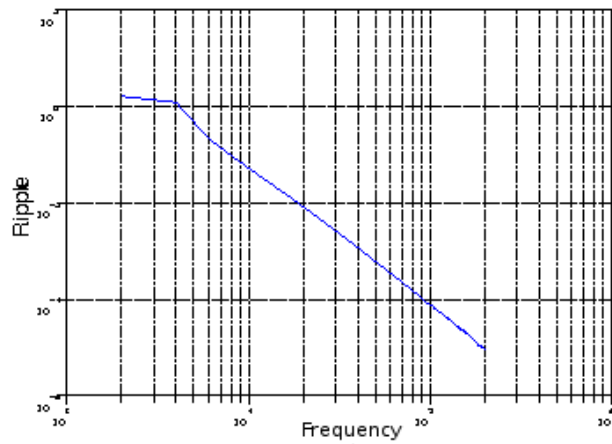


Figure 13: Ripple vs. Frequency

## 5.2 DC MOTOR CONTROL IN REALTIME

This example shows the real time asynchronous control of a DC Motor. For this purpose, we used a small DC drive moving an old PC mouse wheel acting as an encoder (see Fig.14). Each time the wheel moves a small angle, a pulse is sent to the bit *STO* of the parallel port, which triggers an interrupt in the PC. The motor is fed by an amplifier, implemented with a transistor and a filter taking the on-off voltage from a data pin of the parallel port.

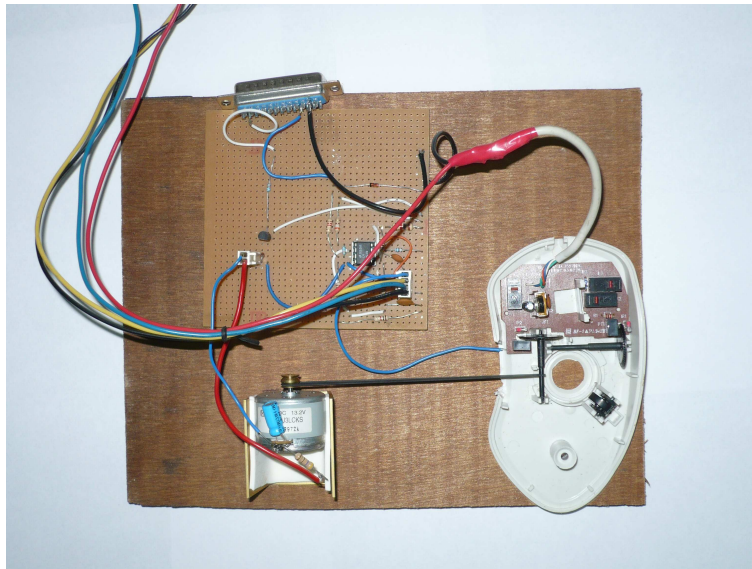


Figure 14: Motor and mouse (encoder)

The control system is composed of the following subsystems (see Fig. 15):

- The *Motor Speed* block detects and counts the interrupts generated by the encoder to estimate the motor speed.

- This estimation is compared to the reference speed (the block *WSum4* calculates the difference or error).
- In base of this error a Proportional-Integral (PI) Control is applied, using QSS discretization.
- The control signal (previously saturated to avoid over-modulation) is pulse width modulated.
- The PWM signal (calculated in the *Comparator1* block), is sent *synchronized* to the parallel port.

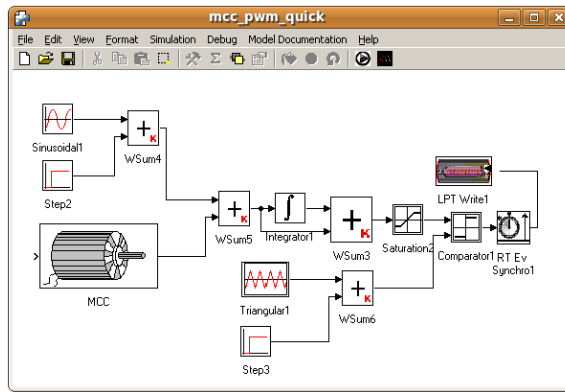


Figure 15: Proportional Control Model

The whole control system of Fig.15 was built using blocks from the *PowerDEVS* library, except the model that estimates the motor speed based on the number of interrupts it receives per second.

Figure 16 shows the reference speed signal while Fig. 17 shows the speed measured by the control system.

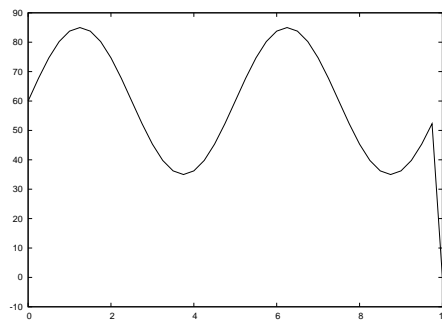


Figure 16: Reference speed

## 6 CONCLUSIONS AND FUTURE WORK

We introduced and described a general purpose tool for DEVS simulation. We illustrated its use in hybrid system simulation, where it shows the most impor-

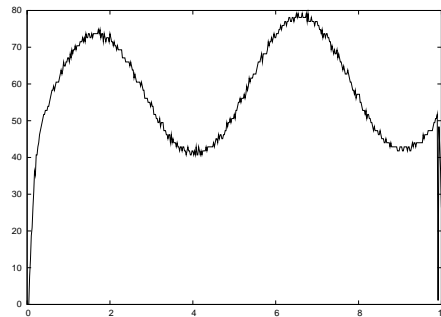


Figure 17: Measured speed

tant facilities and advantages compared with existing simulation software.

Besides the user friendly environment and the simplicity it offers to different kinds of users, *PowerDEVS* has a new way of managing the simulation time advance which allows to implement simulations synchronized with a real-time clock and with the possibility of handling interrupts in an easy fashion. In that way, it can directly implement hybrid asynchronous QSC controllers (as it was done in the example of the DC motor control).

However, *PowerDEVS* is not only a tool for hybrid system simulation and control. As we mentioned before, it is a general purpose DEVS simulator and taking into account that its use is very simple, it results appropriated for education.

Compared with other existing DEVS simulation environments, the main advantage of *PowerDEVS* is the convenient user interface and the simplicity to implement continuous and hybrid system simulations based on the family of QSS methods. Also the possibility to running simulations under a real time operating system (RTAI) and the communication with Scilab are remarkable features.

When it comes to future work, an immediate goal is to finish the migration of *PowerDEVS* source code from Visual Basic to C++ with QT libraries. At the moment, only the model editor needs to be translated. After that, the complete environment would run under different platforms without the need of Windows emulators (the current version of the *PowerDEVS* model editor uses the Wine Windows Emulator to run under Linux and RTAI).

The incorporation of visual programming tools can also improve the atomic model edition. At the moment, users can build complex models by coupling simpler ones (unless they are good programmers). An alternative way to build complex atomic models is with the help of graphical tools, like Simulink Stateflows. A tool that translates graphical formalisms into atomic *PowerDEVS* C++ code might facilitate the generation of new atomic blocks. If such a tool is developed (as a separated program) it can be integrated with *PowerDEVS* in a straightforward manner because of its user-configurable menu features.

Finally, a project is worked on at the ETH Zurich to automatically translate Modelica models into *PowerDEVS* models (based on QSS approximations). The completion of that goal shall permit to drastically increase the modeling capability of *PowerDEVS*, also giving Modelica users the possibility of implementing QSS simulations.

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