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# A Parallel Quantized State System Solver for ODEs

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## Abstract

This work introduces novel parallelization techniques for Quantized State System (QSS) simulation of continuous time and hybrid systems and their implementation on a multi-core architecture. Exploiting the asynchronous nature of QSS algorithms, the novel methodologies are based on the use of non-strict synchronization between logical processes. The fact that the synchronization is not strict allows to achieve large speedups at the cost of introducing additional numerical errors that, under certain assumptions, are bounded depending on some given parameters.

Besides introducing the parallelization techniques, the article describes their implementation on a software tool and it presents a theoretical analysis of the aforementioned additional numerical error. Finally, the performance of the novel methodology and its implementation is deeply evaluated on four large scale models.

*Keywords:* Parallel ODE Simulation, QSS, Hybrid Systems, Discrete Event Systems

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## 1. Introduction

The modeling and simulation (M&S) of continuous systems is at the core of scientific research and different engineering areas. Every year, M&S practitioners develop larger and more complex models and their simulations carry high computational costs. With the advent of multi-core processors and multi-node clusters of computers, the parallel simulation of continuous time systems became the usual way to reduce execution times of these simulations.

Continuous time models are usually expressed as (or transformed to) sets of Ordinary Differential Equations (ODE), where numerical integration algorithms must be applied in order to solve them. Most of these algorithms are based on time-discretization [1, 2], i.e. they compute the value of all state variables at some given time points.

There are certain problems in which the usage of these classic numerical algorithms yields huge computational costs. In particular, in very large systems (millions of state variables) the evaluations of the model functions may require millions of computations. In addition, if the system is stiff (i.e. in presence of fast and slow dynamics), implicit

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algorithms must be used that call for huge matrix inversions on several simulation steps. These problems are worsened in ODEs having frequent discontinuities, where the algorithms must detect their occurrence and restart after each of them.

Over the years, in order to mitigate the huge computational costs associated to these types of problems, various approaches for the parallel implementation of the numerical algorithms have been proposed [3, 4, 5, 6].

There is a newly developed family of ODE numerical integration methods called QSS that replace the time discretization of the classic algorithms by the quantization of the state variables [7, 1]. QSS integration methods have certain features (sparsity exploitation, efficient discontinuities handling [8], explicit stiff integration [9]) that reduce the computational costs in the simulation of large scale systems [10, 11, 12]. These facts motivated the usage of QSS methods in several applications, including Building and Power Systems simulation [13, 14] (where there are also plans to include these algorithm in the EnergyPlus software package [15]), as well as large biological models [16], water distribution models [17], wildfire propagation [18], and simulation of high energy particles [19] (where there is also a preliminary implementation of QSS algorithms in the software Geant4).

QSS methods are asynchronous in the sense that each state variable evolves at a different pace. Thus, if a loosely coupled large scale model is split in two or more parts, they only need to interact at the steps corresponding to changes in variables that are common to different sub-models. This fact enables to spread the simulation over different computing units so that each one integrates a different sub-model.

Based on this idea, a parallel implementation of QSS methods for a multi-core architecture was presented in [20]. That work used a discrete event implementation of the QSS algorithms on a software called PowerDEVS [21], and the synchronization between different sub-models was achieved using a real-time clock. A limitation of that approach was that discrete event implementations of QSS are inefficient [22] and the technique required the usage of a real time operating system, what restricted their usage on general purpose computers.

In this work, we present two novel techniques for parallel simulation with QSS algorithms that do not require the usage of real-time OS. Here, the synchronization between sub-models is non-strict, allowing a bounded difference between the local simulation times. We formally show that this bounded de-synchronization only introduces a bounded numerical error additional to that introduced by the QSS approximation.

We also describe the implementation of these techniques on the Stand Alone Quantized State System Solver [22], a tool implementing the whole family of QSS algorithms that is more than one order of magnitude faster than PowerDEVS. This implementation, based on native POSIX threads, targets shared memory architectures.

In order to evaluate the performance of the implementation, we perform a deep analysis on four large scale problems where it had been shown that the sequential implementation of QSS algorithms is very efficient.

The article is organized as follows. Section 2 introduces the main concepts used along the article and reviews the state of the art and related work in the area of parallel ODE simulation. Then, Section 3 presents a new parallelization technique and its implementation on the QSS Stand Alone Solver. Later, Section 4 analyzes the numerical error introduced by this technique and, based on this analysis, Section 5 proposes a second synchronization technique that performs an adaptive adjustment of the synchronization

parameter. Section 6 presents the application of these two techniques to four large scale models, and Section 7 concludes the article and discusses about future work.

## 2. Background

### 2.1. Quantized State System Methods

Quantized State System (QSS) methods replace the time discretization of classic numerical integration algorithms by the quantization of the state variables.

Given the ODE

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

with  $\mathbf{x}_a \in \mathbb{R}^n$ , the first order Quantized State System method (QSS1) [7] approximates it by

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

Here,  $\mathbf{x}$  is the state vector (of the numerical solution), and  $\mathbf{q}$  is the *quantized state vector*. Each entry  $q_j(t)$ , with  $j = 1, \dots, n$ , is related to that of the state vector  $x_j(t)$  by the following *hysteretic quantization function*:

$$q_j(t) = \begin{cases} x_j(t) & \text{if } |x_j(t) - q_j(t^-)| \geq \Delta Q_j \\ q_j(t^-) & \text{otherwise} \end{cases} \quad (3)$$

where  $\Delta Q_j$  is called *quantum* and  $q_j(t^-)$  denotes the left-sided limit of  $q_j(\cdot)$  at time  $t$ .

It can be easily seen that  $q_j(t)$  follows a piecewise constant trajectory that only changes when the difference between  $q_j(t)$  and  $x_j(t)$  becomes equal to the quantum. After each change in the quantized variable, it results that  $q_j(t) = x_j(t)$ .

Due to the particular form of the trajectories, the solution of Eq. (2) is straightforward and can be easily translated into a simple simulation algorithm.

For  $j = 1, \dots, n$ , let  $t_j$  denote the next time at which  $|q_j(t) - x_j(t)| = \Delta Q_j$ . Then, the QSS1 simulation algorithm works as follows:

Algorithm 1: QSS1.

```

1 while (t < t_f) // simulate until final time t_f
2   t = min(t_j), j ∈ [1, n] // advance simulation time
3   i = argmin(t_j), j ∈ [1, n] // the i-th quantized state changes first
4   e_xi = t - tx_i // elapsed time since last xi update
5   x_i = x_i + x_dot_i * e_xi // update i-th state value
6   q_i = x_i // update i-th quantized state
7   t_i = min(τ > t) subject to |q_i - x_i(τ)| = ΔQ_i // compute next i-th quantized
   state change. Here x_i(τ) = x_i + x_dot_i * (τ - t)
8   for each j ∈ [1, n] such that x_dot_j depends on q_i
9     e_xj = t - tx_j // elapsed time since last xj update
10    x_j = x_j + x_dot_j * e_xj // update j-th state value
11    tx_j = t // last xj update
12    x_dot_j = f_j(q, t) // recompute j-th state derivative
13    t_j = min(τ > t) subject to |q_j - x_j(τ)| = ΔQ_j // recompute j-th quantized
   state changing time. Here x_j(τ) = x_j + x_dot_j * (τ - t)
14   end for
15   tx_i = t // last xi update
16 end while

```

Notice that line 8 requires that the algorithm knows which state derivatives depend on each state variable, i.e., the implementation of a QSS1 solver needs structural information about the model.

The QSS1 method has the following features:

- The quantized states  $q_j(t)$  follow piecewise constant trajectories, and the state variables  $x_j(t)$  follow piecewise linear trajectories.
- The state and quantized variables never differ more than the quantum  $\Delta Q_j$ . This fact ensures stability and global error bound properties [7].
- The quantum  $\Delta Q_j$  of each state variable can be chosen to be proportional to the state magnitude, leading to an intrinsic relative error control [23].
- Each step is local to a state variable  $x_j$  (the one which reaches the quantum change), and it only provokes evaluations of the state derivatives that explicitly depend on it.
- The fact that the state variables follow piecewise linear trajectories makes very easy to detect discontinuities. Moreover, after a discontinuity is detected, its effects are not different to those of a normal step. Thus, QSS1 is very efficient to simulate discontinuous systems [8].

However, QSS1 has some limitations as it only performs a first order approximation, and it is not suitable to simulate stiff systems.

The first limitation was solved with the introduction of higher order QSS methods like the second order accurate QSS2 [24], where the quantized state follow piecewise linear trajectories, and the third order accurate QSS3 [25], where the quantized state follow piecewise parabolic trajectories.

Regarding stiff systems, a family of Linearly Implicit QSS (LIQSS) methods of order 1 to 3 was proposed in [9]. LIQSS methods are explicit algorithms that can efficiently integrate stiff systems provided that they have certain structure.

All QSS and LIQSS methods share the representation of Eq.(2). They only differ in the way that  $q_i$  is computed from  $x_i$ . Thus, their simulation algorithms are very similar to that of Algorithm 1.

## 2.2. The Stand-Alone QSS Solver

As mentioned above, QSS methods replace time discretization of classic integration algorithms by the quantization of the state variables leading to a discrete event approximation of the original model. As a consequence, most QSS implementations are based on DEVS (Discrete Event System Specification) [21]. Although these implementation are simple, the overhead imposed by the synchronization and event transmission mechanism of the DEVS simulation engine makes them inefficient. The stand-alone QSS solver presented in [22] implements the entire family of QSS methods in an efficient way without using a DEVS simulation engine improving simulation times in more than one order of magnitude.

The stand-alone QSS solver simulates models that can contain discontinuities. These models are represented as follows:

$$\dot{\mathbf{x}}(t) = \mathbf{f}(\mathbf{x}, \mathbf{d}, t) \quad (4)$$

where  $\mathbf{d}$  is a vector of discrete variables that can only change when a condition

$$ZC_i(\mathbf{x}, \mathbf{d}, t) = 0 \quad (5)$$

for some  $i \in \{1, \dots, z\}$  is met. The components  $ZC_i$  form a vector of zero-crossing functions  $\mathbf{ZC}(\mathbf{x}, \mathbf{d}, t)$ . When a zero-crossing condition of Eq.(5) is verified, the state and discrete variables can change according to the corresponding event handler:

$$(\mathbf{x}(t), \mathbf{d}(t)) = H_i(\mathbf{x}(t^-), \mathbf{d}(t^-), t) \quad (6)$$

These models are simulated using QSS methods that approximate Eq.(4) by

$$\dot{\mathbf{x}}(t) = \mathbf{f}(\mathbf{q}, \mathbf{d}, t) \quad (7)$$

where each component  $q_i(t)$  is a piecewise polynomial approximation of the corresponding component of the state  $x_i(t)$ .

The simulation is performed by three modules interacting at runtime:

1. The **Integrator**, that integrates Equation (7) assuming that the piecewise polynomial quantized state trajectory  $\mathbf{q}(t)$  is known.
2. The **Quantizer**, that computes  $\mathbf{q}(t)$  from  $\mathbf{x}(t)$  according to the QSS method in use and their tolerance settings (there is a different **Quantizer** for each QSS method). That way, it provides the polynomial coefficients of each quantized state  $q_i(t)$  and computes the next time at which a new polynomial section starts (i.e., when the condition  $|q_i(t) - x_i(t)| = \Delta Q_i$  is met).
3. The **Model**, that computes the scalar state derivatives  $\dot{x}_i = f_i(\mathbf{q}, \mathbf{d}, t)$ , the zero-crossing functions  $ZC_i(\mathbf{x}, \mathbf{d}, t)$ , and the corresponding event handlers  $H_i(\mathbf{q}, \mathbf{d}, t)$ . Besides, it provides the structural information required by the algorithms.

The structure information of the **Model** is automatically extracted at compile time by a **Model Generator** module. This module takes a standard model described in a subset of the Modelica language [26] and produces an instance of the **Model** module as required by the QSS solver including the structure information and the possibility of separately evaluating scalar state derivatives.

Figure 1 shows the basic interaction scheme between the three modules mentioned above.

The structural information is comprised in four binary incidence matrices:

- *SD* (states to derivatives) is such that  $SD_{i,j} = 1$  indicates that  $x_i$  is involved in the calculation of  $\dot{x}_j$ .
- *SZ* (states to zero-crossing functions) is such that  $SZ_{i,j} = 1$  indicates that  $x_i$  is involved in the calculation of  $ZC_j$ .
- *HD* (handlers to state derivatives) is such that  $HD_{i,j} = 1$  indicates that the execution of handler  $H_i$  changes some state or discrete variable involved in the calculation of  $\dot{x}_j$ .

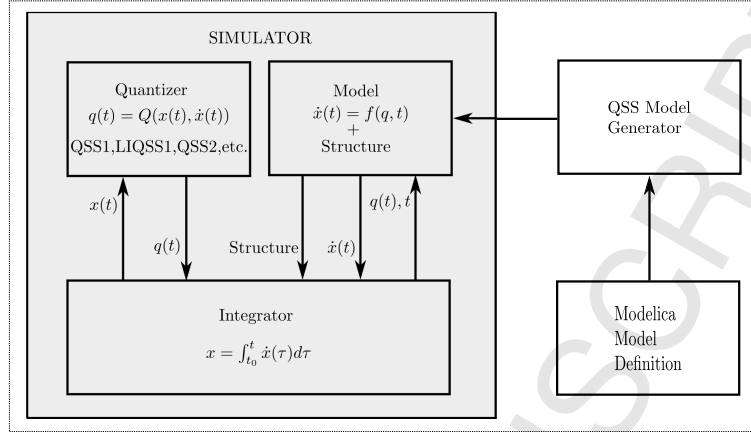


Figure 1: Stand Alone QSS Solver – Basic Interaction Scheme

- $HZ$  (handlers to zero-crossing functions) is such that  $HZ_{i,j} = 1$  indicates that the execution of handler  $H_i$  changes some state or discrete variable involved in the calculation of  $ZC_j$ .

Taking into account that most large systems are loosely coupled, the structure matrices are stored in a sparse form.

The simulation is carried on by the **Integrator** module, that advances the simulation time executing the simulation steps. Each simulation step may correspond to a change in a quantized variable  $q_i$  or to the execution of an event handler  $H_i$  triggered by a zero-crossing condition  $ZC_i(t) = 0$ . The integrator stores the state, quantized state and discrete state values  $x_i$ ,  $q_i$ , and  $d_i$ , respectively. It also stores the time of the next change in each quantized state  $tx_i$  and the time of the next crossing of each zero-crossing function  $tz_i$ .

The main simulation routine at the **Integrator** module looks as follows

Algorithm 2: QSS Integrator Module

```

1  while  $t < t_f$  //while simulation time t is less than the final time  $t_f$ 
2     $t_x = \min(tx_j)$  // time of the next change in a quantized variable
3     $t_z = \min(tz_j)$  // time of the next zero--crossing time
4     $t = \min(t_x, t_z)$  //advance simulation time
5    if  $t = t_x$  then //quantized state change
6       $i = \operatorname{argmin}(tx_j)$  // the i-th quantized state changes first
7      Quantized_State_Step(i) //execute a quantized state change on
      variable i
8    else //zero--crossing
9       $i = \operatorname{argmin}(tz_j)$  // the i-th event handler is executed first
10     Event_Handler_Step(i) //execute the procedure for the i-th event
      handler
11   end if
12 end while

```

When the next step corresponds to a change in a quantized variable  $q_i$  at time  $t$ , the **Integrator** proceeds as follows:

Algorithm 3: QSS Integrator Module - Quantized State Change

```

1 Quantized_State_Step(i)
2 {
3   integrateState( $x_i, \dot{x}_i, t$ ) // integrate i-th state up to time  $t$ .
4   Quantizer.update( $x_i, q_i$ ) // update i-th quantized state  $q_i$ 
5    $tx_i =$  Quantizer.nextTime( $x_i, q_i$ ) //compute next i-th quantized state change
      time
6   for each  $j$  such that  $SD_{i,j} = 1$ 
7     integrateState( $x_j, \dot{x}_j, t$ ) // integrate j-th state up to time  $t$ .
8      $\dot{x}_j =$  Model. $f_j(\mathbf{q}(t), \mathbf{d}(t), t)$  // recompute j-th state derivative
9      $tx_j =$  Quantizer.nextTime( $x_j(t), q_j(t)$ ) //recompute next j-th quantized
      state change time
10  end for
11  for each  $j$  such that  $SZ_{i,j} = 1$ 
12     $zc_j =$  Model. $zc_j(\mathbf{q}, \mathbf{d}(t), t)$  // recompute j-th zero--crossing function
13     $tz_j =$  nextEventTime( $zc_j$ ) //recompute next j-th zero--crossing time
14  end for
15 }
```

Similarly, when the next step corresponds to the execution of an event handler  $H_i$  at time  $t$ , the **Integrator** proceeds as follows:

Algorithm 4: QSS Integrator Module - Event Handler Execution

```

1 Event_Handler_Step(i)
2 {
3   Model. $H_i(\mathbf{q}(t), \mathbf{d}(t), t)$  //execute i-th event handler
4   for each  $j$  such that  $HD_{i,j} = 1$ 
5     integrateState( $x_j, \dot{x}_j, t$ ) // integrate j-th state up to time  $t$ .
6      $\dot{x}_j =$  Model. $f_j(\mathbf{q}(t), \mathbf{d}(t), t)$  // recompute j-th state derivative
7      $tx_j =$  Quantizer.nextTime( $x_j(t), q_j(t)$ ) //recompute next j-th quantized
      state change time
8   end for
9   for each  $j$  such that  $HZ_{i,j} = 1$ 
10     $zc_j =$  Model. $zc_j(\mathbf{q}, \mathbf{d}(t), t)$  // recompute j-th zero--crossing function
11     $tz_j =$  nextEventTime( $zc_j$ ) //recompute next j-th zero--crossing time
12  end for
13 }
```

### 2.3. Parallel Simulation

Simulations often require performing several expensive calculations. In order to obtain faster results, the usage of parallel computations is a common solution. As the current work deals with parallel simulation of ODEs using discrete event approximations, we shall mention some existing solutions in both fields: parallel simulation of ODEs and discrete event systems. Also, existing approaches for the parallel simulation with QSS methods are described as the most related results to our methodology.



### 2.3.1. Parallel ODE Simulation

Different strategies have been proposed over the years for parallel ODE simulation based on classic numerical integration algorithms. These strategies are characterized according to the computations that they parallelize. The main categories are:

- *parallelism across the model*: This technique is based on partitioning the mathematical model in a way that different components of function  $\mathbf{f}$  in Eq.(1) are computed in parallel.
- *parallelism across the method*: In this approach, parallelism is achieved by executing the intrinsic calculations of the numerical method in parallel. In most implementations, this approach is combined with the parallelization across the model.
- *parallelism across the time steps*: Here, the successive integration steps of the simulation are computed in parallel.

A comparative analysis between the two first approaches is presented in [27].

An implementation that exploits efficient memory access using Iterated Runge Kutta methods in shared memory architectures can be found in [4, 5, 28]. In these works, parallelizing simultaneously across the methods, models and time steps, speedups of up to 300 times are achieved using 480 cores of a supercomputer in the simulation of ODEs with up to 8,000,000 state variables.

In [29] different integration methods are benchmarked on both CPU and GPU platforms. Similarly in [30] two Runge-Kutta methods are tested on CPU and GPU achieving speed-ups up to 115x for the GPU case.

Different applications of the parallelism across steps approach for classic numerical methods are presented and studied in [6, 31, 32, 33, 34].

### 2.3.2. Parallel Simulation of Discrete Event Systems

Parallelization of Discrete Event System simulations has been studied for some years now [35, 36]. The basic idea is to split the model into several sub-models and to simulate them concurrently on different *logical processors* (LPs), each one having its own logical time. As the sub-models are usually inter-dependent, the logical time of the different sub-simulations must be synchronized in order to satisfy the *causality constraint*, i.e., the preservation of the chronological order of the events.

The literature divides the different approaches into three categories:

- *Conservative* algorithms, like CMB [37], where there is a synchronization mechanism that forces each LP to wait until it is sure that it will not receive messages from “the past”. Conservative approaches usually achieve small speedups and suffer from certain issues (like possible deadlocks). They can be enhanced with the usage of *LookAhead* strategies [38].
- *Optimistic* algorithms, where each LP advances its logical time as much as it can and it rolls-back when an inconsistency among the interchanged messages is found. Optimistic approaches allow more parallel computations than conservative techniques at the expense of having to save intermediate simulation states and costly roll-back mechanisms. Algorithms like TimeWarp [39, 40] are examples of the optimistic approach. A recent work [41] shows results for TimeWarp running a

MPI based parallel simulation on almost two million cores obtaining super-linear speed-ups.

- Finally in [42], the idea of completely avoiding the synchronization is studied, i.e. each LP advances its logical time as fast as it can disregarding of what the other LP logical time is. The results show that this approach achieves larger speedups than the previous ones at the cost of introducing errors due to violation of the causality constraint.

### 2.3.3. Parallel Simulation with QSS methods

The asynchronous nature of QSS methods simplifies the parallelization of their computations. As it was mentioned in Section 2.2, QSS methods can be represented as discrete events system, thus the corresponding parallelization techniques of Section 2.3.2 could be used, in principle.

However, neither of these approaches fits well in the context of QSS algorithms. In this case, strict synchronization does not allow almost any concurrent computations as it was shown in [20]. Also, optimistic methods would require huge amounts of memory to implement the roll-back mechanisms on large systems. Finally totally unsynchronized techniques would introduce unacceptable numerical errors.

An initial study was done in [43] applying optimistic algorithm to parallelize QSS over a distributed-memory cluster showing a two-fold speedup simulating on four processors. Then, in [44] the authors present an implementation of QSS methods over shared-memory GPU architectures (Nvidia Tesla C1060, and Nvidia GeForce 8600). Preliminary results report speedups of up to eight times for models with 64 state variables. The results were not extended to larger models due to limitations on the GPU architecture (diverging branches, lack of synchronization, etc.).

Another related work was presented in [21] where two parallelization techniques, SRTS and ASRTS, are introduced for a shared-memory multi-core architecture. The synchronization amongst the different sub-simulations in this case is achieved using a real-time clock. Results show an almost linear scaling of speed-up with respect to the number of logical processors achieving speed-ups of up to 9 times for 12 logical processors. The cited work also analyzes the error introduced by these techniques showing that for a bounded difference between the logical time of each sub-simulation a bounded numerical error is introduced. This implementation was based on the PowerDEVS tool [21].

In conclusion, previous to the current work, the parallelization of QSS methods was limited to relatively small problems (a few thousands of state variables, at most) and implemented over different architectures using small numbers of processors.

## 3. A Novel Parallel QSS Simulation Methodology

In this section we describe a novel technique for parallel simulation with QSS algorithms and its implementation in the Stand-Alone QSS Solver. We first present the basic idea, and then we introduce the algorithms and implementation issues corresponding to the different components of the parallel solver.

### 3.1. Basic Idea

The presented parallelization technique is based on partitioning the model into  $P$  sub-models, so that each sub-model is simulated by a different logical process.

At each simulation step of every sub-model, the corresponding LP checks if the variable that changed must be communicated to other LPs using structural information. If so, the new values and the corresponding time-stamp are informed through an inter-process communication mechanism.

In order to avoid large errors introduced by new values of a quantized variable arriving at wrong time instants, the technique enforces the different processes to have a bounded difference between their logical simulation times. This difference is bounded by a user defined parameter  $\Delta t$ .

This non-strict synchronization is achieved by computing a global virtual time  $gvt$  (equal to the minimum logical time of all LPs) and not allowing the LPs to advance beyond  $gvt + \Delta t$ . For that goal, whenever an LP schedules its next simulation step beyond  $gvt + \Delta t$ , it enters a waiting routine until  $gvt$  advances or a change in a variable computed by another LP is detected.

### 3.2. Basic Structure of the Parallel Solver

The parallel extension of the QSS solver targets a MIMD (multiple instruction multiple data) multi-core architecture. It uses a partitioned model of Eqs.(4)–(6), where each LP simulates a different sub-model. Every LP is composed of an **Integrator** module, a **Quantizer** module, and a **Model** module, like in the sequential simulation described in Section 2.2.

Taking into account that the state derivatives computed in a LP may depend on the states computed in another LP, it is clear that the different LPs must communicate during the simulation. Furthermore, the computation of a state derivative  $\dot{x}_i$  at time  $t$  requires the knowledge of the quantized states involved in the calculation of  $f_i(\mathbf{q}, \mathbf{d}, t)$  at the same time  $t$ , thus, a synchronization mechanism between the different LPs is also necessary.

Towards this goal, the sequential **Integrator** module is modified with the addition of communication and synchronization mechanisms. The basic scheme of the parallel QSS solver is shown in Figure 2

Next, after discussing the model partitioning issues, we describe in detail the communication and synchronization mechanisms and the resulting parallel simulation algorithm and their implementation.

### 3.3. Model Partition

The QSS parallel simulation algorithm requires that the original model is split into  $p$  sub-models, so that each sub-model is simulated by a different LP.

In order to obtain an effective parallelization, the model partitioning should be balanced, and, at the same time, the communication between different LPs must be minimized. In some simple cases, when the system has a regular structure, this partitioning can be manually done.

In general cases, algorithms for automatic partitioning must be applied. A set of graph-theoretic sub-optimal partitioning algorithms for QSS parallel simulation was developed in [45].

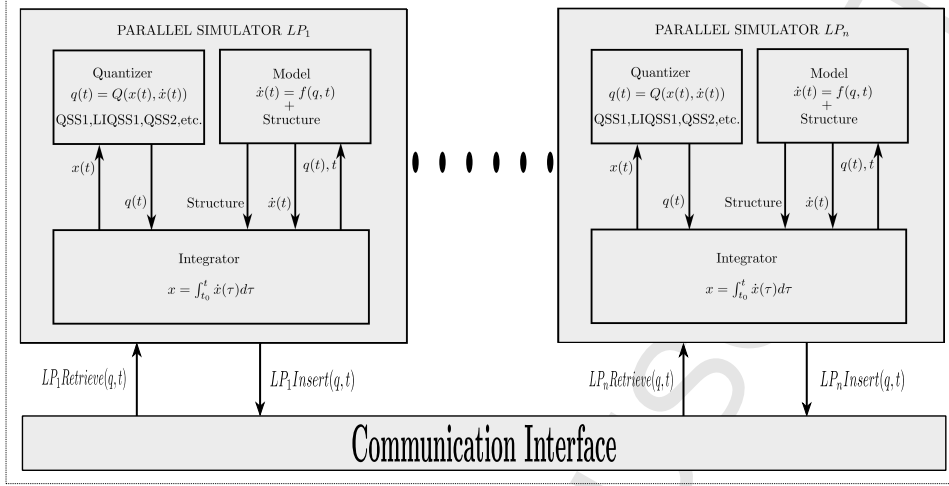


Figure 2: Parallel Stand Alone QSS Solver – Basic Interaction Scheme

In the context of this work, we shall assume that a *suitable* partition is provided.

The partitioning shall be represented by two arrays  $PX$ , and  $PZ$ , with their entries taking values in the set  $\{1, \dots, p\}$ . An entry  $PX_i = k$  says that the state  $x_i$  is computed by the  $k$ -th LP. Similarly, an entry  $PZ_i = k$  says that the zero-crossing function  $ZC_i$  and its corresponding event handler  $H_i$  are evaluated and executed by the  $k$ -th LP.

In the actual implementation of the parallel QSS solver, the end user can choose between the different automatic partitioning algorithms described in [45] or he can provide a manually generated partition.

### 3.4. Inter-Process Structure

As we explained in Section 2.2, the structure information of the model is comprised in matrices  $SD$ ,  $SZ$ ,  $HD$ , and  $HZ$ . These structural matrices represent the direct influences of states and event handlers in state derivatives and zero-crossing functions.

Once the model is partitioned into  $p$  sub-models (according to arrays  $PX$  and  $PZ$ ), it can happen that a change in a state variable –or the execution of an event handler– computed at the  $k$ -th sub-model has a direct influence on some state derivatives or zero-crossing functions computed at different sub-models.

Thus, an inter-process communication mechanism is necessary, which in turn requires the knowledge of inter-process structure information.

In our implementation, this information is provided by two incidence matrices at each LP:

- $SO^k$  is such that  $SO_{i,l}^k = 1$  indicates that the  $i$ -th state of the  $k$ -th sub-model influences on some state derivative or zero-crossing function computed by the  $l$ -th sub-model.
- $HO^k$  is such that  $HO_{i,l}^k = 1$  indicates that the variables changed by the  $i$ -th event handler execution at the  $k$ -th sub-model influences on some state derivative or zero-crossing function computed by the  $l$ -th sub-model.

Matrices  $SO^k$  and  $HO^k$  are computed at initialization time based on the structure matrices  $SD$ ,  $SZ$ ,  $HD$ ,  $HZ$ , and the partition arrays  $PX$  and  $PZ$ . Like the remaining structure matrices,  $SO^k$  and  $HO^k$  are stored in a sparse format.

### 3.5. Simulation Algorithm

The parallel simulation algorithm is locally implemented at each logical process. As we already explained, each LP contains an **Integrator**, a **Quantizer**, and a **Model** module. The **Quantizer** and **Model** modules are identical to their sequential counterparts, while the **Integrator** now includes the communication and synchronization mechanisms.

The communication involves informing the new values of states and discrete variables to other LPs after state changes or event handler executions that, according to the inter-process structure, affect state derivatives or zero-crossing functions calculated at other LPs. On the other side, the LPs must check if those new values were changed

As we explained before, the synchronization mechanism limits the advance of the simulation at each process in order to keep a bounded difference between the logical simulation times of the different LPs.

Later on, we shall explain in more detail the communication and synchronization procedures.

Taking into account these modifications, the parallel  $k$ -th **Integrator** algorithm works as follows:

Algorithm 5: QSS Parallel Integrator Module

```

1  while  $t^k < t_f$  //while simulation time of the k--th LP is less than the
   final time  $t_f$ 
2   $t_x^k = \min(t_{x_j} | PX_j = k)$  // time of the next change in a quantized variable
   computed by the k-th LP
3   $t_z^k = \min(t_{z_j} | PZ_j = k)$  // time of the next zero--crossing time computed by
   the k--th LP
4   $t_m^k = \text{ChangesList.First().time}()$  // timestamp of the first informed
   change
5   $t^k = \min(t_x^k, t_z^k, t_m^k)$  // attempt to advance LP simulation time
6   $gvt = \min(t^i)$  // recompute global virtual time
7  Synchronize() // call synchronization procedure
8  if  $t^k = t_x^k$  then //state change
9     $i = \text{argmin}(t_{x_j})$  // the i-th quantized state changes first
10   Parallel_Quantized_State_Step(i) //execute a quantized state change
   on variable i
11  elseif  $t^k = t_z^k$  then //zero--crossing
12    $i = \text{argmin}(t_{z_j})$  // the i-th event handler is executed first
13   Parallel_Event_Handler_Step(i) //execute the procedure for the i-th
   event handler
14  else
15   Process_External_Change() //process the first enqueued change
16  end if
17  end while

```

This algorithm is similar to its sequential counterpart (Algorithm 2). It adds the calculation of the global virtual time  $gvt$  as the minimum between the local simulation

times of all LPs and a synchronization routine to avoid that the local time  $t^k$  advances much beyond  $gvt$ . Additionally, it takes into account the incoming changes in the input variables informed by different LPs.

When the next step in the  $k$ -th LP corresponds to a change in a quantized variable  $q_i$  at time  $t$ , the **Integrator** proceeds as follows:

Algorithm 6: QSS Parallel Integrator Module - Quantized State Change

```

1 Parallel_Quantized_State_Step(i)
2 {
3   integrateState( $x_i, \dot{x}_i, t^k$ ) // integrate i-th state up to time  $t^k$ .
4   Quantizer.update( $x_i, q_i$ ) // update i-th quantized state  $q_i$ 
5    $tx_i = \text{Quantizer.nextTime}(x_i, q_i)$  //compute next i-th quantized state change
      time
6   for each  $l$  such that  $SO_{i,l}^k = 1$ 
7     ChangesList.InsertState(1,i, $q_i, t_k$ ) //inform new quantized state and
      time stamp to  $l$ --th LP
8   end for
9   for each  $j$  such that  $SD_{i,j} = 1$  and  $PX_j = k$ 
10    integrateState( $x_j, \dot{x}_j, t$ ) // integrate j-th state up to time  $t$ .
11     $\dot{x}_j = \text{Model}.f_j(\mathbf{q}^k(t), \mathbf{d}^k(t), t)$  // recompute j-th state derivative
12     $tx_j = \text{Quantizer.nextTime}(x_j(t), q_j(t))$  //recompute next j-th quantized
      state change time
13  end for
14  for each  $j$  such that  $SZ_{i,j} = 1$  and  $PZ_j = k$ 
15     $zc_j = \text{Model}.zc_j(\mathbf{q}^k, \mathbf{d}^k(t), t)$  // recompute j-th zero--crossing function
16     $tz_j = \text{nextEventTime}(zc_j)$  //recompute next j-th zero--crossing time
17  end for
18 }
```

This algorithm is very similar to the sequential one (Algorithm 3). The difference is that it only updates the state derivatives that are calculated by the current LP, and, additionally, it informs the values and the corresponding time-stamp to the other LPs that compute the remaining state derivatives affected by the quantized state change.

Notice that the entire quantized state vector  $\mathbf{q}(t)$  used to calculate state derivatives and zero-crossing functions can contain components  $q_j(t)$  computed at different LPs. In that case, the  $k$ -th LP has a *local* copy (possibly outdated) of its actual value. Thus, the vector is denoted as  $\mathbf{q}^k(t)$ , comprising the components calculated at the LP and the local copies of the remaining entries. A similar remark can be done regarding the discrete state  $\mathbf{d}(t)$ .

When the next step corresponds to the execution of an event handler  $H_i$  at time  $t$ , the **Integrator** proceeds like Algorithm 4 with identical modifications.

As we mentioned above, when a Logical Process informs changes in quantized states or discrete variables, the new values are stored in a time-stamp-sorted list. Additionally, on each simulation step, every LP checks the time-stamp of the first element of the list and, in case the time-stamp ( $m_1.t$ ) is less than the time of the scheduled next local change, the **Integrator** processes the external change as follows (assuming that the incoming new values are provoked by a quantized state change):

Algorithm 7: QSS Parallel Integrator Module - External Quantized State Change

```

1 Process_External_Change ()
2 {
3   m1=ChangesList.RetrieveState() //get the first element of the list.
4   i=m1.index() //index of the quantized variable that changed
5   q_i^k = m1.q //update local copy of quantized state q_i.
6   for each j such that SD_{i,j}=1 and PX_j=k
7     integrateState(x_j, x_dot_j, t) // integrate j-th state up to time t.
8     x_dot_j = Model.f_j(q(t), d(t), t) // recompute j-th state derivative
9     tx_j = Quantizer.nextTime(x_j(t), q_j(t)) //recompute next j-th quantized
      state change time
10  end for
11  for each j such that SZ_{i,j}=1 and PZ_j=k
12    zc_j = Model.zc_j(q, d(t), t) // recompute j-th zero--crossing function
13    tz_j = nextEventTime(zc_j) //recompute next j-th zero--crossing time
14  end for
15 }

```

When the incoming variables that changed corresponds to a handler execution, the procedure is similar to the previous one.

### 3.6. Inter-Process Synchronization

As we mentioned before, the parallel implementation of the QSS solver uses a non-strict synchronization mechanism that is necessary to ensure that calculations are actually performed in parallel.

This synchronization mechanism requires that, before each simulation step, the LPs calculate

$$gvt = \min_{1 \leq l \leq P} (t^l) \quad (8)$$

as the minimum logical simulation time of all LPs. Then, each LP limits the advance of its local simulation time  $t^k$  to the value  $gvt + \Delta t$ , where  $\Delta t$  is a user defined parameter. Notice that  $\Delta t$  is an upper bound for the difference between the local simulation times of all the LPs and it defines how strict the synchronization is.

The synchronization procedure of each LPs implements a waiting routine while the condition

$$t^k > gvt + \Delta t \quad (9)$$

is verified. When  $gvt$  advances (because other LPs update their local simulation time) or when  $t^k$  goes back (because the  $k$ -th LP detected an external change), and the condition of Eq.(9) is no longer accomplished, the simulation at the  $k$ -th LP continues. That way, the synchronization routine works as follows:

Algorithm 8: LP Synchronization

```

1 Synchronize()
2 {
3   while ( $t^k - gvt > \Delta t$ )
4      $rcv = \text{ChangesList.First().time()}$  // timestamp of the first informed
       change
5     if ( $rcv < t^k$ )
6        $t^k = rcv$ 
7      $gvt = \min(t^l)$  // recompute the minimum global virtual time
8   end while
9 }

```

Notice that the line 7 of this code explicitly computes the minimum global virtual time depending on the local logical times  $t^l$  of all processes. This calculation is the reason that restricts the usage of the whole strategy to shared memory architectures. Any attempt to use this algorithm on a distributed memory architecture would result in a huge traffic through the network communicating the logical times of the different processors at each step.

Figure 3 illustrates the synchronization mechanism during a parallel simulation with two LPs.

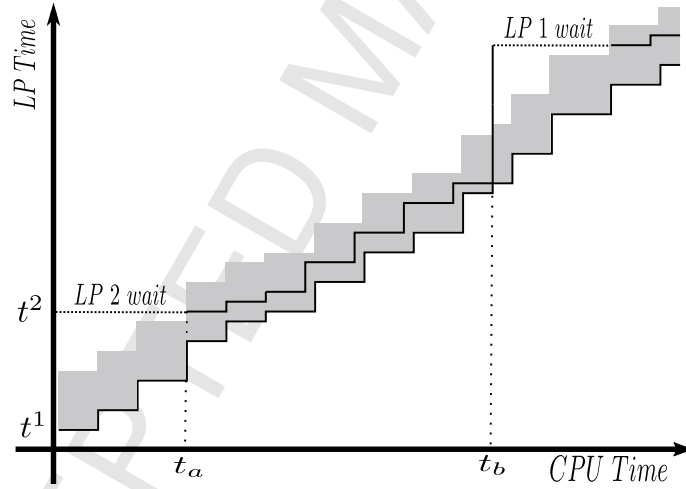


Figure 3: Parallel QSS solver synchronization scheme example

There, the maximum simulation time to which a LP can advance is limited by  $gvt + \Delta t$ , represented by the gray region above  $gvt$ .

At the beginning of the simulation, the second logical process (LP2) schedules its next change  $t^2$  to a value that is larger than  $gvt + \Delta t$ , so it must wait until  $gvt$  advances (i.e., the simulation time  $t^1$  of LP1 advances).

Then, at the instant of CPU time  $t_a$ , the simulation time  $t^1$  (and also  $gvt$ ) is such that  $t^2 < gvt + \Delta t$  and LP2 can resume the simulation steps. During the interval



$(t_a, t_b)$ , the distance between  $t^1$  and  $t^2$  is less than  $\Delta t$  and so both processes simulate in parallel without waiting for synchronization. Then, at time  $t_b$ , the first process LP1 schedules its next simulation step for time  $t^1 > gvt + \Delta t$ , and now it must wait until the synchronization point is reached again.

Notice that if we had taken  $\Delta t = 0$ , then each LP can only advance when  $gvt = t^k$ , leading to a sequential simulation and there is no parallelization at all. On the contrary, if we set  $\Delta t \geq t_f$  (where  $t_f$  is the final simulation time) there is no synchronization at all and each LP simulates as fast as it can, like in a *No-Time* strategy, but the numerical error introduced would possibly result unacceptable.

Thus, a good value for  $\Delta t$  should be such that the error is acceptable and enough parallelization is achieved. We shall deeply analyze this problem later, proposing also an adaptive strategy for finding an adequate value for  $\Delta t$ .

### 3.7. Inter-Process Communication

As we mention above, the different LPs informs changes in state and discrete variables during the simulation. For that purpose, each logical process has a list of the changes occurred in the variables computed at the other LPs that are needed for the local computations. This list is sorted by their corresponding time-stamps and it provides the following services:

- **InsertState**( $l, i, q_i, t_k$ ): Inserts the new value of the  $i$ -th quantized state  $q_i$  that changed at time  $t_k$  computed by the  $k$ -th LP into the  $l$ -th LP list of changes.
- **InsertEvent**( $l, i, \{d_j\}, \{x_j\}, t_k$ ), Inserts the new values of the set of discrete states  $\{d_j\}$  and the set of continuous states  $\{x_j\}$  that changed after the execution of the  $i$ -th event handler at time  $t_k$  computed by the  $k$ -th LP into the  $l$ -th LP list of changes.
- **RetrieveState**( $l$ ), Gets the first element of the list of changes caused by an external quantized state update.
- **RetrieveEvent**( $l$ ), Gets the first element of the list of changes caused by an external handler execution.

If a change with a time-stamp  $m.t$  less than the time of the last step performed by the LP  $t^{k-}$  is detected, then the time-stamp is modified to  $t^{k-}$ . That way, we avoid that the simulation goes back in time.

Notice that the modification of this time-stamp is the only effect that synchronization errors have on the simulation results. Anyway, the difference between  $m.t$  and  $t^{k-}$  is always bounded by  $\Delta t$ .

The communication mechanism between threads is hybrid:

- State changes are asynchronously informed using the previously mentioned lists of changes.
- Discrete changes are also informed using lists of changes, but in a synchronous way where the LP that communicates the new values must wait until all the receiver LPs process them.

The later case ensures that the new values of all the discrete variables are processed by all the affected LPs before the LP that is responsible for the change continues with the simulation. Taking into account that discrete changes can provoke large changes in discrete and state variables, this policy prevents the introduction of large numerical errors due to lack of synchronization after discontinuities.

It is worth mentioning that in most models containing discontinuities the number of continuous changes is significantly larger than the number of discrete changes. Therefore, in practice, the communication is mostly asynchronous.

### 3.8. Some Implementation Details

The execution model for each logical processor is implemented using native Linux POSIX threads. Threads are pinned to specific CPUs using `sched_setaffinity`. The assignment of threads to CPUs is given by the partition matrices  $PX$  and  $PZ$ .

Initially, the main thread reads the partition structure and computes the interprocess structure matrices. Then, it creates  $P$  threads where each of them locally implements an instance of a **Parallel Simulator** module containing a **Parallel Integrator**, a **Quantizer**, and a **Model** modules.

At this point, there are two alternative implementations. The first one, uses a **Model** module identical to that of the sequential implementation. Since the sequential model computes the state derivatives, handlers and zero crossing functions depending on the full state, this implementation requires working with arrays containing the full state size.

In this first implementation, the **Parallel Integrator** module has a data structure containing the following local data:

- The local logical simulation time  $t^k$  and a local copy of the global virtual time  $gvt$ .
- The corresponding inter-process structure matrices  $SO^k$  and  $HO^k$ .
- Two arrays containing the indexes of the state derivatives and zero-crossing functions evaluated at the current LP.
- A sorted list containing changes corresponding to quantized state variables.
- A sorted list containing changes corresponding to event handler executions.

Additionally, a compilation flag allows to keep local copies of

- The full state and quantized state arrays  $\mathbf{x}$ , and  $\mathbf{q}$ .
- The full array of zero-crossing function values.
- The full discrete state array  $\mathbf{d}$ .

Otherwise, those arrays are shared by all the LPs together with the remaining global data, consisting of:

- An array containing the local simulation times of all LPs  $t^l$ .
- The four incidence matrices  $SD$ ,  $HD$ ,  $SZ$ , and  $HZ$ .

The optional usage of local copies for the full state and quantized state allows to target Non-Uniform Memory Access (NUMA) platforms. Here, the memory latency for a given address depends on each processor. Thus, by allocating local copies we enforce that each processor accesses its own memory bank. Independently on this, having local copies of the full discrete array can help to avoid concurrency problems due to the fact that each discrete variable can be computed by more than one LP in certain models.

The price paid for using local full array copies is a large increase in the memory consumption. However, it simplifies the implementation, and, in cases where the memory usage is not critical, acceptable results can be obtained.

A second implementation uses a different model instance for each LP, so that the functions defining each sub-models depend only on the local variables of each LP. Also, the structure information provided by each sub-model is local to that sub-model. Thus, in this implementation, each LP contains only local copies of the corresponding sub-arrays, whose size is proportional to the size of the sub-model. That way, as we increase the number of partitions, the total memory consumption is almost unaffected.

The drawback of this implementation is that it requires the generation of a different model instance for each LP. Starting from the original model and structure matrices, the construction of the partitioned sub-models requires remapping the state variable indexes and to generate the corresponding code for the resulting functions and structure. In theory, this problem has a straightforward solution, but its automatic implementation requires to modify the **Model Generator** module of the QSS Solver.

We are currently working on this last issue. So far, the current version of the parallel QSS Solver supports the usage of partitioned models, but the sub-model instances must be manually generated.

The performance of the alternative implementations shall be compared later on an example in Sec.6.1.

For both implementations, in spite of the fact that all the data is allocated in shared memory, the only data that is actually updated by the different processes concurrently are the arrays containing the local simulation times of the LPs (for synchronization purposes) and the changes lists used for communication.

The array with the local simulation times is accessed for computing the global virtual time  $gvt$ . Anyway, in the implementation,  $gvt$  is only updated at the  $k$ -th LP when the local time  $t^k$  becomes greater than  $gvt + \Delta t$ . That way, in most steps, there is no actual access to that shared array.

#### 4. Synchronization and Numerical Errors

The non strict synchronization mechanism adopted can provoke that when the  $k$ -th LP informs a change in a quantized state  $q_i$  at time  $t^k$ , this change is detected late by the  $l$ -th LP, i.e., when the  $l$ -th LP has already performed one or more steps at time  $t^l > t^k$ . In that case, as there is no roll-back mechanism, the  $l$ -th LP will just modify the time-stamp of the incoming values pretending that the change in  $q_i$  occurred at time  $t^l$ .

This implies that, during the interval  $(t^k, t^l)$ , the  $l$ -th LP integrates using an incorrect (outdated) local copy of the quantized state  $q_i$ . Anyway, taking into account that the difference between  $t^l$  and  $t^k$  is bounded by  $\Delta t$ , and knowing that  $q_i$  is an approximation of

a continuously varying signal  $x_i(t)$ , we can expect that the difference between the actual value of  $q_i$  (computed at the  $k$ -th LP) and the local copy at the  $l$ -th LP is not significant and that it will introduce a bounded numerical error in the subsequent calculations.

The following analysis formally shows this fact.

Consider the QSS approximation of an ODE, given by

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

and suppose that this system is partitioned in two parts as follows:

$$\begin{aligned}\dot{\mathbf{x}}_a(t) &= \mathbf{f}_a(\mathbf{q}_a(t), \mathbf{q}_b(t), \mathbf{v}(t)) \\ \dot{\mathbf{x}}_b(t) &= \mathbf{f}_b(\mathbf{q}_a(t), \mathbf{q}_b(t), \mathbf{v}(t))\end{aligned}\quad (10)$$

where:

$$\mathbf{x} = [\mathbf{x}_a \ \mathbf{x}_b]^T; \quad \mathbf{q} = [\mathbf{q}_a \ \mathbf{q}_b]^T; \quad \mathbf{f} = [\mathbf{f}_a \ \mathbf{f}_b]^T$$

Let us consider that we implement the simulation of this system in parallel, such that  $\mathbf{x}_a$  is computed in one LP and  $\mathbf{x}_b$  is computed in a second LP according to the mechanism explained in the previous section.

Then, the non strict synchronization implies that the first LP can have a local copy  $\mathbf{q}_b^c(t)$ , which is an outdated version of the components of  $\mathbf{q}_b(t)$ . Similarly, the second LP can have a local copy  $\mathbf{q}_a^c(t)$ , containing outdated versions of the components of  $\mathbf{q}_a$ .

Thus, the parallel algorithm simulates the following approximation of Eq.(10):

$$\begin{aligned}\dot{\mathbf{x}}_a(t) &= \mathbf{f}_a(\mathbf{q}_a(t), \mathbf{q}_b^c(t), \mathbf{v}(t)) \\ \dot{\mathbf{x}}_b(t) &= \mathbf{f}_b(\mathbf{q}_a^c(t), \mathbf{q}_b(t), \mathbf{v}(t))\end{aligned}\quad (11)$$

Let us define  $\Delta_a(t) \triangleq \mathbf{q}_a^c(t) - \mathbf{q}_a(t)$ , and  $\Delta_b(t) \triangleq \mathbf{q}_b^c(t) - \mathbf{q}_b(t)$ . Then, Eq.(11) can be rewritten as

$$\begin{aligned}\dot{\mathbf{x}}_a(t) &= \mathbf{f}_a(\mathbf{q}_a(t), \mathbf{q}_b(t) + \Delta_b(t), \mathbf{v}(t)) \\ \dot{\mathbf{x}}_b(t) &= \mathbf{f}_b(\mathbf{q}_a(t) + \Delta_a(t), \mathbf{q}_b(t), \mathbf{v}(t))\end{aligned}\quad (12)$$

that constitutes a *perturbed* version of the original system of Eq.(10), with perturbation terms  $\Delta_a(t)$  and  $\Delta_b(t)$ .

Notice that a component  $q_i(t)$  computed at the first LP and its local copy  $q_i^c(t)$  at the second LP can have a maximum delay in time given by the parameter  $\Delta t$ , i.e.,  $q_i^c(t) = q_i(t - \tau)$  with  $0 \leq \tau < \Delta t$ . Then, it results that

$$|q_i^c(t) - q_i(t)| = |q_i(t - \tau) - q_i(t)| \leq |x_i(t - \tau) - x_i(t)| + 2\Delta Q_i$$

where we used the fact that the difference between  $x_i$  and  $q_i$  is always bounded by the quantum  $\Delta Q_i$ . Then, assuming that the state derivative  $\dot{x}_i = f_i(\mathbf{q}, t)$  is bounded by a constant  $M_i$  while the quantized states  $\mathbf{q}(t)$  remains in certain bounded region, it results that

$$|x_i(t - \tau) - x_i(t)| < M_i \cdot \tau \leq M_i \Delta t$$

and then

$$|q_i^c(t) - q_i(t)| \leq M_i \cdot \Delta t + 2\Delta Q_i \quad (13)$$

Applying the same analysis to all the components of the array  $\mathbf{q}_a$  and  $\mathbf{q}_b$ , it results that all the components of the perturbation terms  $\Delta_a(t)$  and  $\Delta_b(t)$  are bounded in their absolute value for a constant depending on the parameter  $\Delta t$  and the quantum  $\Delta Q$ .

When the original system of Eq.(1) is linear time invariant (LTI) and asymptotically stable, it can be easily proved that the presence of the bounded perturbations  $\Delta_a(t)$  and  $\Delta_b(t)$  only adds a bounded numerical error proportional to the perturbation bound [24, 1]. In nonlinear cases, assuming that the perturbation bound is small enough, a similar property can be derived [7].

If we had more than two processors, the analysis can be easily extended arriving to the same results.

In conclusion, the parallelization introduces an additional error to that introduced by the QSS methods. This additional error can be bounded by a quantity that increases proportional to the parameter  $\Delta t$ .

Although this conclusion is of qualitative nature (i.e., it does not provide a quantitative bound for the additional numerical error), in the next Section we shall derive an algorithm for adaptive computation of  $\Delta t$  in order to ensure a quantitative error bound.

## 5. Adaptive $\Delta t$ Adjustment

The election of the parameter  $\Delta t$  imposes a trade-off between numerical errors and the actual parallelization achieved. Obtaining a suitable value for  $\Delta t$  usually involves running a set of experiments in order to observe the error introduced by the parameter and the speedup obtained. Also, a value for  $\Delta t$  that is suitable at the beginning of the simulation may be not optimal after some time due to changes in the dynamics of the model.

In order to overcome these problems, we developed an adaptive algorithm that allows the dynamic calculation of  $\Delta t$  in order to keep a bounded numerical error.

As we analyzed in the previous section, the parallel implementation with non strict synchronization provokes a difference between the quantized states computed in one LP and their local copies used in other LPs. This difference can be seen as a bounded perturbation that introduces a bounded numerical error.

The difference between a quantized state  $q_i(t)$  and a local copy  $q_i^c(t)$  is bounded according to Eq.(13). Thus, if we want to bound this perturbation to be proportional to the prescribed tolerance, it should be proportional to the quantum  $\Delta Q_i$ :

$$|q_i^c(t) - q_i(t)| \leq M_i \cdot \Delta t + 2\Delta Q_i \leq \alpha \cdot \Delta Q_i$$

Here,  $M_i$  is an upper bound for the state derivative  $\dot{x}_i(t)$  and  $\alpha$  is a user defined parameter. From this last inequality we obtain

$$\Delta t \leq \frac{(\alpha - 2) \cdot \Delta Q_i}{M_i} \quad (14)$$

That limit on  $\Delta t$  must be accomplished for all the quantized states that have local copies in other LPs, i.e, for all the *output* quantized states.

The set of output quantized states of the  $k$ -th LP can be formally defined as  $O_k \triangleq \{i | \exists l \text{ such that } SO_{i,l}^k = 1\}$ . Then, the set of all output quantized states results  $O \triangleq \bigcup_k O_k$ .

As we want that Eq.(14) is satisfied for all the output quantized states, the adaptive algorithm takes  $\Delta t$  as

$$\Delta t = \min_{i \in O} \frac{(\alpha - 2) \cdot \Delta Q_i}{M_i} \quad (15)$$

For that goal, each LP computes its minimum  $\Delta t^k$  as

$$\Delta t^k = \min_{i \in O_k} \frac{(\alpha - 2) \cdot \Delta Q_i}{M_i} \quad (16)$$

and then  $\Delta t$  is computed as the minimum of all  $\Delta t^k$ .

The bounds on the state derivatives  $M_i$  are numerically estimated as<sup>1</sup>

$$M_i \approx \frac{|x_i(t_i) - x_i(t_i^{prev})|}{t_i - t_i^{prev}} \quad (17)$$

where  $t_i$  and  $t_i^{prev}$  are the two last times of change in the quantized state  $q_i$ .

Notice that the estimate  $M_i$  is updated each time the output quantized state  $q_i$  changes. That change in  $M_i$  may modify the LP minimum  $\Delta t^k$  computed in Eq.(16) that can in turn change the global  $\Delta t$ . Thus,  $\Delta t$  is asynchronously updated. This fact in turn implies the need of an additional synchronization mechanism that must be introduced in the **Parallel Integrator** module.

To achieve this goal, the **Parallel Integrator** Algorithm 5 must be modified so that when a simulation step changes an output quantized variable, the local minimum  $\Delta t^k$  can be updated. If so, the algorithm checks if this update changes the global value  $\Delta t$ . If the global parameter is affected, a global synchronization routine is called where all LPs obtain the new value for  $\Delta t$ .

For that purpose, the data structure of each LP is extended with the addition of a local copy of the global  $\Delta t$  parameter. Also, a global array shared by all LPs contains the local minimum  $\Delta t^k$ .

Notice that the usage of this strategy only requires to choose the *error bound coefficient*  $\alpha$ , and then the parameter  $\Delta t$  is automatically adapted. In principle, the choice  $\alpha \approx 10$  seems reasonable, as the error introduced by the lack of synchronization would be at most one order of magnitude larger than the error introduced by the QSS approximation itself.

Notice that, while the analysis of Section 4 provides a qualitative error bound, the strategy for adaptation of  $\Delta t$  provides a quantitative bound for the numerical error, where we ensure that it cannot become larger than  $\alpha$  times the error introduced by the sequential QSS algorithm. Thus, provided that the quantum  $\Delta Q$  was adequately chosen by the user for the sequential case, the parallel implementation will work within a prescribed tolerance.

---

<sup>1</sup> $M_i$  can be directly obtained as  $M_i = |\dot{x}_i|$ , as  $\dot{x}_i$  is stored by the QSS algorithms. However, when a solution oscillates around an equilibrium point (as it occurs often in QSS solutions)  $\dot{x}_i$  can be larger than the actual variation of the state, and a numerical approximation can provide a better result.

## 6. Results

In this section we present simulation results on four large-scale models. In these models, we analyze and compare the performance of the parallel QSS Solver using the proposed synchronization strategies with different parameter settings.

The examples are ordered by their complexity. The first one, corresponding to a large population of air conditioner units, is a hybrid system that does not require communication between LPs so it is useful to measure the overhead introduced by the Parallel QSS Solver mechanism. The second model is a two dimensional Advection-Reaction partial differential equation (PDE) semi-discretized with the Method of Lines [1]. This is a purely continuous system that allows also to validate the results regarding the numerical error introduced by the parallelization strategies. The third model is a modification of the first one with the addition of a centralized power control, that introduces inter-process communication of discontinuous changes. The last one, corresponding to a network of spiking neurons, has a complex connection structure with events that provoke instantaneous changes on continuous states.

### *Simulation Framework*

The results presented below were obtained using a 64-core server with 32 GB of RAM running a 64 bit Linux OS (Ubuntu 14.04). The server is composed of 4 AMD Opteron 6272 processors, with 16 cores (in 8 physical modules) each<sup>2</sup> and a Non-uniform memory access (NUMA) architecture. We used the Parallel QSS Solver implementation version 3.1 (git commit number [c10a14]).

In order to evaluate the performance of the simulations we took into account the following metrics:

- Initialization Time: It corresponds to the time spent at the initialization routines of the Parallel QSS Solver.
- Simulation Time: It is the CPU time taken by the simulation at the slowest logical process. It does not take into account the Initialization Time.
- Memory Usage: We report the total amount of memory allocated by the simulation.
- Simulation Speedup:  $Speedup(P) = \frac{T_1}{T_P}$ , where  $T_1$  is the simulation time spent using a single core and  $T_P$  is the simulation time spent using  $P$  cores.
- Normalized Mean Error<sup>3</sup>:

$$error = \frac{mean(|\mathbf{y}^P - \mathbf{y}^{ref}|)}{mean(\mathbf{y}^{ref})} \quad (18)$$

<sup>2</sup>The AMD Opteron 6272 processors are based on the AMD Bulldozer micro-architecture. They have two integer units but only one floating point (FP) scheduler per module, thus, depending on the application, each processor can be considered to have 8 or 16 cores. As the QSS solver is always combining FP with integer operations (as the structure plays a fundamental role in the computations), it can be expected in principle that the 16 threads can efficiently run in parallel.

<sup>3</sup>In this work we compute the mean error rather than the maximum global error because in the examples the states are either discontinuous or have very steep trajectories. That way, a very small delay between two trajectories provokes that the maximum error is equal to the trajectory amplitude (independently on the delay value). Thus, the mean error captures better the actual difference between the simulated trajectories.

where the reference trajectory  $\mathbf{y}^{ref}$  is the one obtained using a sequential simulation whereas  $\mathbf{y}^P$  is the trajectory obtained with  $P$  cores. Both trajectories are evaluated at 5000 equidistant time points.

Notice that this error does not include the numerical error introduced by the QSS approximation. It only evaluates the additional numerical error introduced by the lack of synchronization between processes.

For the different models, we performed the following experiments:

1. In order to analyze the effects of using different values of the synchronization parameter  $\Delta t$ , we run several simulations varying that parameter. These simulations were performed using 62 cores<sup>4</sup>.
2. After selecting a suitable value for  $\Delta t$  (i.e., a value that achieves a good trade-off between speedup and error) we run simulations varying the number of cores in order to characterize the speedup ratio w.r.t. the number of cores. We
3. For the adaptive synchronization strategy, we run several simulations varying the error bound parameter  $\alpha$  on a 62 cores configuration.
4. Then, using the error bound parameter  $\alpha = 10$  (which is a reasonable choice, as analyzed in the previous section), we run simulations varying the number of cores in order to characterize the achieved speedup.

In all cases, the models were manually partitioned.

### 6.1. Air Conditioners Population

This model, taken from [11], studies the dynamics of a large population of air conditioners (ACs) when they follow the same reference temperature. The  $i$ -th air conditioner is used to control the temperature of the  $i$ -th room, modeled by:

$$\dot{\theta}_i(t) = -\frac{1}{C_i \cdot R_i} [\theta_i(t) - \theta_a + R_i \cdot P_i \cdot m_i + w_i(t)] \quad (19)$$

where  $R_i$  represents the thermal resistance,  $C_i$  is the thermal capacity,  $P_i$  is the power of the AC unit in *on* state,  $\theta_a$  is the outside ambient temperature (common to all rooms), and  $w_i(t)$  is a noise signal representing thermal disturbances.

The variable  $m_i(t)$  is the state of the  $i$ -th AC unit that takes the value 1 in *on* state and 0 otherwise. This state follows a hysteretic on-off control law:

$$m_i(t^+) = \begin{cases} 0 & \text{if } \theta_i(t) \leq \theta_r^i(t) - 0.5 \text{ and } m_i(t) = 1 \\ 1 & \text{if } \theta_i(t) \leq \theta_r^i(t) + 0.5 \text{ and } m_i(t) = 0 \\ m_i(t) & \text{otherwise} \end{cases} \quad (20)$$

where  $\theta_r^i(t)$  is the reference temperature, that obeys the following profile:

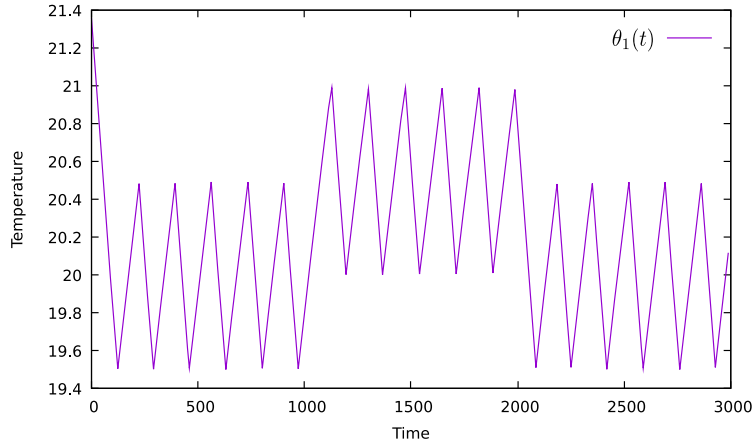
$$\theta_r^i(t) = \begin{cases} 20 & \text{if } 0 \leq t < 1000 \\ 20.5 & \text{if } 1000 \leq t < 2000 \\ 20 & \text{if } 2000 \leq t \leq 3000 \end{cases} \quad (21)$$

<sup>4</sup>We used a maximum of 62 cores leaving 2 cores free to attend the OS tasks.



Finally, the thermal disturbances  $w(t)$  is updated once every minute, taking pseudo-random values uniformly distributed in the interval  $(-1, 1)$ . The temperature of the first AC unit is depicted in Fig.4.

Figure 4: Air Conditioners Population Model - Output Trajectory



For this experiment, we simulated a population of 248000 AC units and set their parameters according to [11].

Notice that each air conditioner is modeled by one differential equation (Eq.(19)) and three zero crossing functions: one associated to the hysteretic control law of Eq.(20), another corresponding to the evolution of the reference temperature of Eq.(21), and the last one associated to the update of the noise signal  $w_i(t)$ . Thus, the model contains a set of 248000 differential equations and 992000 zero-crossing functions.

Observe also that the dynamics of each air conditioner unit is independent on the evolution of the remaining air conditioners. Therefore, the model can be partitioned in a way such that no communication between the different sub-models is required and hence, the lack of synchronization in the parallel simulation does not introduce errors.

This model was simulated using the QSS2 method with a quantization of  $\Delta Q_{rel} = \Delta Q_{min} = 1e - 3$  and the final simulation time  $t_f$  was set to 3000 minutes.

In this case, the Parallel QSS solver automatically detects that there is no communication between the different LPs and it automatically sets the parameter  $\Delta t$  equal to the final simulation time 3000. Thus, in this introductory case, we did not run experiments for different values of  $\Delta t$  or  $\alpha$  as these parameters are meaningless here.

Also, we do not report errors as they were null in all cases (in absence of communication, sequential and parallel results are identical).

The goal of this example is to measure the overhead introduced by the parallel simulation when the synchronization mechanism is not needed. This overhead includes the access to the structural matrices (that are shared by the different LPs), the additional calls to the synchronization routines and the access to the inter-process communication structures. Additionally, in this experiment, we include the results obtained using a the model partitioning strategy described in Section 3.8 to improve the memory con-

sumption. In this simple case, we were able to generate the different **Model** instances manually.

Then, we run simulations for a different number of cores ranging from 1 to 62. The results are reported in Table 1.

Table 1: Air Conditioners Population Model - Simulation Results  
Speedup and memory consumption comparison between the solver implementation and the model partition prototype (MPP)

<b>Cores</b>	<b>Init Time (ms)</b>	<b>Simulation Time (seconds)</b>	<b>Memory (MBytes)</b>	<b>Speedup</b>	<b>MPP Simulation Time (seconds)</b>	<b>MPP Memory (MBytes)</b>	<b>MPP Seedup</b>
1	580	1132	360	1.00	1132	360	1.00
2	894	668	655	1.69	644	388	1.75
4	948	357	1081	3.17	341	501	3.31
8	969	192	1835	5.89	185	544	6.11
16	1085	109	3342	10.38	96	628	11.79
32	1165	52	6356	21.76	50	804	22.64
48	1281	32	9371	35.37	31	976	36.51
62	1406	25	10223	45.28	23	1057	49.21

As we can see in Table 1, the achieved speedups for this model are linear. Running on 62 cores speeds up the simulation by a factor of 45 using full state copies and 49 using the model partition strategy. The speed difference between these strategies can be explained by the additional memory accesses needed by the non partitioned implementation. Initialization times are increased with the number of cores (the initialization involves the computation and allocation of interprocess communication matrices, that grow in complexity with the number of processes). Anyway, the initialization times are neglectable in all cases.

The memory consumption, in turn, grows almost linearly with the number of cores in the non partitioned implementation, increasing the memory consumption more than 28 times for 62 cores. This is due to the fact that each LP has local copies of the states, quantize states, etc., and, additionally, they allocate communication data structures and the interprocess communication matrices. However, when using the model partitioning strategy, we can observe that the memory consumption for 62 cores is less than 3 times the memory needed for running the simulation with 1 core. This is due to the fact that each sub-model contains only the sub-array of the state, quantized state and zero-crossing functions that each LP needs, together with the corresponding structural information which is also partitioned. Also, as a consequence, there is no need to store the additional arrays containing the indexes of the state derivatives and zero-crossing functions evaluated at each LP.

As an additional test, we measured the simulation time spent by the sequential implementation of the QSS Solver. This experiment took 1050 seconds, which is about 8% faster than the parallel implementation running on a single core. This overhead is due to the extra control mechanisms of the parallel solver (for communication, synchronization, and access to inter-process structures). These mechanisms are not disabled when

running on a single core although they are not actually necessary. Similar tests were run on the different examples arriving always to very similar overhead figures.

A final remark is that, in this case, the simulation using one core takes less than 20 minutes so the large-scale nature of the model can be questioned. Anyway, this was just an illustrative case, where we can easily simulate the system until a final time ten times larger arriving to identical conclusions.

## 6.2. Advection Reaction Equation

The following model, taken from [12], represents a 2D Advection–Reaction equation. This equation can describe, for instance, a river transporting a substance experiencing a chemical reaction.

After applying the method of lines on this PDE, the following set of ODEs is obtained:

$$\dot{u}_{i,j} = -a_x \frac{u_{i,j} - u_{i,j-1}}{\Delta x} - a_y \frac{u_{i,j} - u_{i-1,j}}{\Delta y} + ru_{i,j}(u_{i,j} - \alpha)(u_{i,j} - 1) \quad (22)$$

for  $i = 2, \dots, N$ ,  $j = 2, \dots, M$ , where  $u_{i,j}(t)$  is the concentration of the transported substance at the  $i, j$  grid point of the spatial domain. The parameters  $a_x$  and  $a_y$  represent the speed of the transporting flow in the  $x$  and  $y$  coordinates respectively, and  $r$  is the rate of the chemical reaction. Finally,  $\Delta x$  and  $\Delta y$  are the width of each grid section.

At the borders, the dynamics is defined by

$$\dot{u}_{i,1} = -a_x \frac{u_{i,1}}{\Delta x} - a_y \frac{u_{i,1} - u_{i-1,1}}{\Delta y} + ru_{i,1}(u_{i,1} - \alpha)(u_{i,1} - 1) \quad (23)$$

for  $i = 2, \dots, N$

$$\dot{u}_{1,j} = -a_x \frac{u_{1,j} - u_{1,j-1}}{\Delta x} - a_y \frac{u_{1,j}}{\Delta y} + ru_{1,j}(u_{1,j} - \alpha)(u_{1,j} - 1) \quad (24)$$

for  $j = 2, \dots, M$ , and,

$$\dot{u}_{1,1} = -a_x \frac{u_{1,1}}{\Delta x} - a_y \frac{u_{1,1}}{\Delta y} + ru_{1,1}(u_{1,1} - \alpha)(u_{1,1} - 1) \quad (25)$$

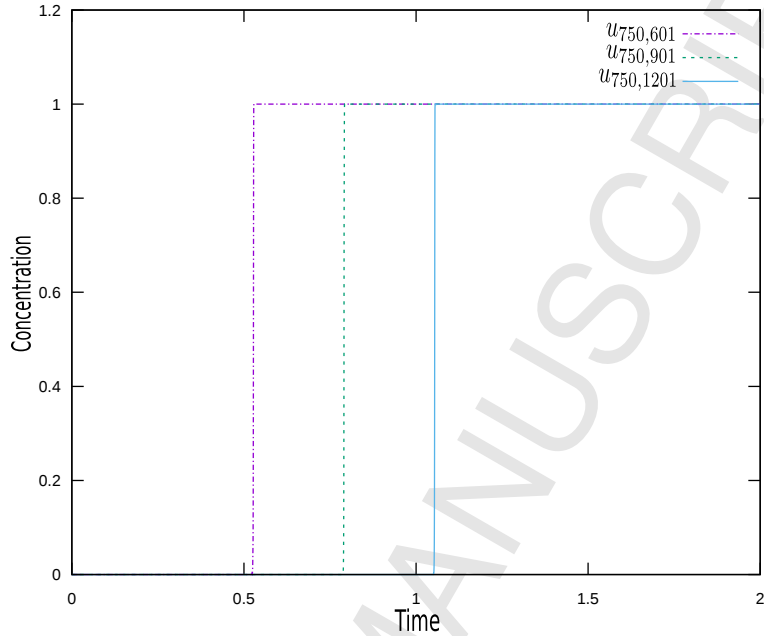
The initial conditions are set to

$$u_{i,1} = 1 \quad (26)$$

for  $i = 1, \dots, N$  and  $u_{i,j} = 0$  otherwise.

The concentration of three output variables in the middle of the grid, namely  $u_{750,301}$ ,  $u_{750,901}$  and  $u_{750,1201}$  are illustrated in Fig.5.

Figure 5: Advection Reaction Model - Output Trajectories



Taking into account that the model is *stiff* (the chemical reactions are much faster than the transport dynamics), the linearly implicit LIQSS2 algorithm was used, setting the tolerance to  $\Delta Q_{rel} = \Delta Q_{abs} = 10^{-3}$ . The model parameters were  $r = 10000$ ,  $a_x = 1$ ,  $a_y = 0.1$ ,  $\Delta x = 1/N$ ,  $\Delta y = 1/M$  with  $N = M = 1500$ . That way, we have a spatial domain of size  $1 \times 1$  with  $1500 \times 1500 = 2250000$  grid points, where each grid point represents a state equation.

Table 2 reports the simulation results for different values of the  $\Delta t$  parameter on 62 cores. Here, the error reported is the mean error on 20 state variables  $u_{i,j}(t)$  taken over a line at the middle of the spatial domain. Each error was computed using Eq.(18).

Table 2: Advection Reaction Model - Fixed  $\Delta t$  - Using 62 Cores

$\Delta t$	Simulation Time (seconds)	Speedup	Error
1e-07	1104	2.40	3.61e-06
1e-06	288	9.20	3.45e-06
1e-05	145	18.28	3.66e-06
1e-04	119	22.27	6.22e-06
1e-03	121	21.90	1.38e-05
1e-02	115	23.05	2.12e-05
1e-01	112	23.66	2.39e-04

As expected, for increasing values of  $\Delta t$  the speedup and the error increase. From these results, evidently, the best performance is obtained with  $\Delta t = 1e - 4$ . Thus, fixing  $\Delta t$  to this value, we computed the speedup and errors for different number of cores (Table 3).

Table 3: Advection Reaction Model - Fixed  $\Delta t = 1e - 04$

Cores	Init Time (ms)	Simulation Time (seconds)	Memory (MBytes)	Speedup	Error
1	174	2650	1058	1.00	0
2	699	2248	1626	1.18	3.63e-06
4	666	1216	2763	2.18	3.66e-06
8	748	715	5035	3.71	3.33e-06
16	850	375	9580	7.07	3.14e-06
32	1100	203	14573	13.05	5.44e-06
48	1300	153	23647	17.32	4.83e-06
62	1460	119	31614	22.27	6.22e-06

We can notice that the error has always the same order of magnitude. As before, the memory consumption and the initialization times grow with the number of cores (the initialization time is again negligible). Finally, the speedup reaches a reasonable value of 22 for 62 cores, growing almost linearly with the number of cores.

We also simulated the model using the adaptive  $\Delta t$  strategy, varying the error bound parameter  $\alpha$ , obtaining the results reported in Table 4.

Table 4: Advection Reaction Model - Adaptive  $\Delta t$  - Using 62 Cores

$\alpha$	Simulation Time (seconds)	Speedup	Error
2	169	15.68	2.96e-06
10	135	19.63	4.50e-06
20	124	21.37	4.57e-06
50	122	21.72	6.71e-06
100	127	20.87	9.33e-06

As predicted in Section 5, the best results are obtained around  $\alpha = 10$ , achieving a similar speedup to that obtained using  $\Delta t = 1e - 4$ , but with less error. Notice that using this strategy the speedup and errors are robust with the choice of the parameter  $\alpha$ .

Using the error bound parameter  $\alpha = 10$ , we then varied the number of cores, obtaining the results reported in Table 5.

Table 5: Advection Reaction Model - Adaptive  $\Delta t$  -  $\alpha = 10$ 

Cores	Init Time (ms)	Simulation Time (seconds)	Memory (MBytes)	Speedup	Error
1	174	2650	1058	1.00	0
2	672	2251	1626	1.18	3.63e-06
4	720	1218	2763	2.18	3.54e-06
8	733	716	5035	3.70	3.75e-06
16	871	381	9580	6.96	3.05e-06
32	1015	207	14573	12.80	3.70e-06
48	1270	178	23647	14.89	4.99e-06
62	1468	135	31614	19.63	4.50e-06

Notice that the error does not change significantly and the speed up follows a very similar evolution to that obtained using a fixed parameter  $\Delta t = 1e - 4$ .

It is worth mentioning that obtaining the optimal value  $\Delta t = 1e - 4$  required performing several experiments, while the value  $\alpha = 10$  is just the default choice for the adaptive algorithm.

In order to compare the performance of the parallel QSS Solver with that of a classic parallel ODE solver, we simulated the same model using the parallel extension of the CVODE package called PVODE [46] on the same server, under the same tolerance settings. The PVODE solver was configured to use the OpenMP library and the Adams numerical algorithm (BDF did not work due to the very large size). The results are reported in Table 6.

Table 6: Advection Reaction Model - CVODE Solver - Simulation Results

Cores	Simulation Time (seconds)	Speedup
1	12552	1
2	6814	1.84
4	3581	3.50
8	1854	6.77
16	1293	9.70
32	909	13.80
48	741	16.93
62	751	16.71

From the results, we also observe an almost linear speed-up until 48 cores, where it saturates. The QSS solver is about 4.73 times faster than CVODE running on a single core. Then, running on 62 cores, the QSS solver becomes about 6.3 times faster than PVODE.

These figures show that the QSS solver is faster in a sequential simulation and the

difference becomes bigger using 62 cores. It is worth mentioning that the QSS solver is not optimized to deal with 2D arrays and the simulation code produced is very inefficient.

### 6.3. Power Control of an Air Conditioner Population

This model, also taken from [11], extends the model of Section 6.1, by adding a central control of the total power consumption. Towards this end, the total AC population is divided into local sections  $S = \{s_1, \dots, s_n\}$  and the power consumed by the  $k$ -th section is computed as:

$$p_i(t_k) = \sum_{j \in s_k} P_j \quad (27)$$

and the total power consumption of the AC population is computed as:

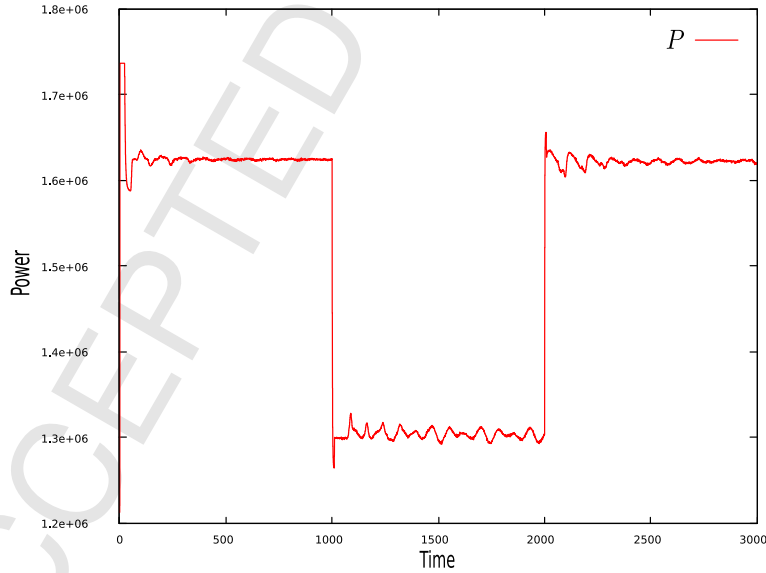
$$P(t_k) = \sum_{i \in S} p_i \quad (28)$$

The global control system regulates the total power  $P(t_k)$  so that it follows a desired power profile  $P_r(t)$ . To achieve this goal, a proportional integral (PI) control law is used to compute the common reference temperature as:

$$\theta_r(t_k) = K_P \cdot [P_r(t_k) - P(t_k)] + K_L \cdot \int_{\tau=0}^{t_k} [P_r(\tau) - P(\tau)] d\tau \quad (29)$$

Figure 6 depicts the output of the power control of a sequential simulation of the model.

Figure 6: Power Control of an Air Conditioner Population Model - Output Trajectory



We simulated a population of 248000 ACs with 248 local sections, so in this case the model contains 248000 state variables and 496498 zero-crossing functions. We used the

QSS3 method with a quantization of  $\Delta Q_{rel} = 1e - 4 = \Delta Q_{min} = 1e - 4$  and the model parameters set as in Section 6.1.

The error introduced by the parallel implementation was measured on the total power consumption of the ACs ( $P(t_k)$ ) according to Eq. (18).

Table 7 reports the simulation results obtained for different  $\Delta t$  values using 62 cores.

Table 7: Power Control of an Air Conditioner Population Model - Fixed  $\Delta t$  - Using 62 Cores

$\Delta t$	Simulation Time (seconds)	Speedup	Error
1e-05	2009	1.56	1.42e-04
1e-04	761	4.13	1.82e-04
1e-03	432	7.27	1.81e-04
1e-02	250	12.56	1.85e-04
1e-01	200	15.70	2.95e-04
5e-01	94	33.39	4.99e-04
1	90	34.88	5.50e-04
2	124	25.32	5.84e-04

As expected, the speedup and error grow with the value of  $\Delta t$ . Starting from  $\Delta t = 1$ , there is no appreciable gain in the speedup, so we used this value for the next experiments. This time, the maximum speedup is in the order of 35 for 62 cores (it was 52 in the first example, that did not need synchronization). In this example, the different LPs communicate discrete changes (changes in the power consumption of each section or changes in the global reference temperature). As explained in Section 3.7 the communication of discrete changes enforces a logical process to wait until the receivers process the corresponding change, what slows down the simulation.

Then, using  $\Delta t = 1$ , we simulated the model for different number of cores obtaining the results reported in Table 8.

Table 8: Power Control of an Air Conditioner Population Model - Fixed  $\Delta t = 1$

Cores	Init Time (ms)	Simulation Time (seconds)	Memory (MBytes)	Speedup	Error
1	106	3139	458	1.00	0
2	240	1830	655	1.72	9.18e-04
4	267	1019	1048	3.08	1.02e-03
8	294	648	1835	4.84	2.02e-04
16	382	426	3440	7.37	1.25e-03
32	526	176	6651	17.84	5.46e-04
48	673	117	9830	26.83	5.72e-04
62	796	90	12648	34.88	5.50e-04



The results are very similar to those of the previous example in terms of initialization times, memory consumption and errors. The speedup, again, grows almost linearly.

Then, we simulated the system with the adaptive algorithm for different error bound parameter  $\alpha$ , obtaining the results reported in Table 9.

Table 9: Power Control of an Air Conditioner Population Model - Adaptive  $\Delta t$  - Using 62 Cores

$\alpha$	Simulation Time (seconds)	Speedup	Error
2	128	24.53	3.88e-04
10	91	34.50	1.47e-03
20	92	34.12	1.46e-03
50	89	35.27	1.39e-03
100	90	34.88	1.59e-03

Again, the value  $\alpha = 10$  has an almost optimal performance. Using this parameter, we simulated the model for different number of cores, obtaining the results reported in Table 10.

Table 10: Power Control of an Air Conditioner Population Model - Adaptive  $\Delta t$  -  $\alpha = 10$

Cores	Init Time (ms)	Simulation Time (seconds)	Memory (MBytes)	Speedup	Error
1	106	3139	458	1.00	0
2	195	1799	655	1.75	1.36e-03
4	209	1032	1048	3.04	1.43e-03
8	210	651	1835	4.82	4.77e-04
16	251	426	3440	7.37	1.33e-03
32	313	172	6651	18.25	9.01e-04
48	381	117	9830	26.83	1.09e-03
62	425	91	12648	34.50	1.47e-03

Again, the results are very similar to those obtained with the optimal value  $\Delta t = 1$  in Table 7.

#### 6.4. Spiking Neural Network

The last model, adapted from [47], represents a network of spiking neurons. The  $i$ -th neuron is modeled by three differential equations:

$$\begin{aligned} \tau \cdot \dot{v}_i(t) &= v_{rest} - v_i(t) + g_i^{ex}(t) \cdot (E^{ex} - v_i(t)) + g_i^{inh}(t) \cdot (E^{inh} - v_i(t)) \\ \tau^{ex} \cdot \dot{g}_i^{ex}(t) &= -g_i^{ex}(t) \\ \tau^{inh} \cdot \dot{g}_i^{inh}(t) &= -g_i^{inh}(t) \end{aligned}$$

where  $v_i(t)$  represents the membrane potential,  $g_i^{ex}(t)$  is the excitatory conductance, and  $g_i^{inh}(t)$  is the inhibitory conductance. The definition and values for the remaining parameters can be found in [47].

Whenever the membrane potential  $v_i(t)$  reaches the threshold value  $-50$ , the neuron performs a spike, resetting its potential to the value  $v_i(t) = v_{rest} = -60$ . The spike is transmitted to the post synaptic connected neurons, that change their excitatory or inhibitory conductance depending on the type of neuron that performed the spike. If the  $i$ -th neuron is of excitatory type, the excitatory conductance of their post-synaptic neurons is changed according to

$$g_j^{ex}(t^+) = g_h^{ex}(t) + \Delta g^{ex}$$

for all  $j \in Post_i$  (the set of post synaptic neurons of neuron  $i$ ). If the  $i$ -th neuron is of inhibitory type, the inhibitory conductance of their post-synaptic neurons is changed as

$$g_j^{inh}(t^+) = g_h^{inh}(t) + \Delta g^{inh}$$

for all  $j \in Post_i$ . For these synaptic connections, we used parameters  $\Delta g^{ex} = 0.4$  and  $\Delta g^{inh} = 1.6$ .

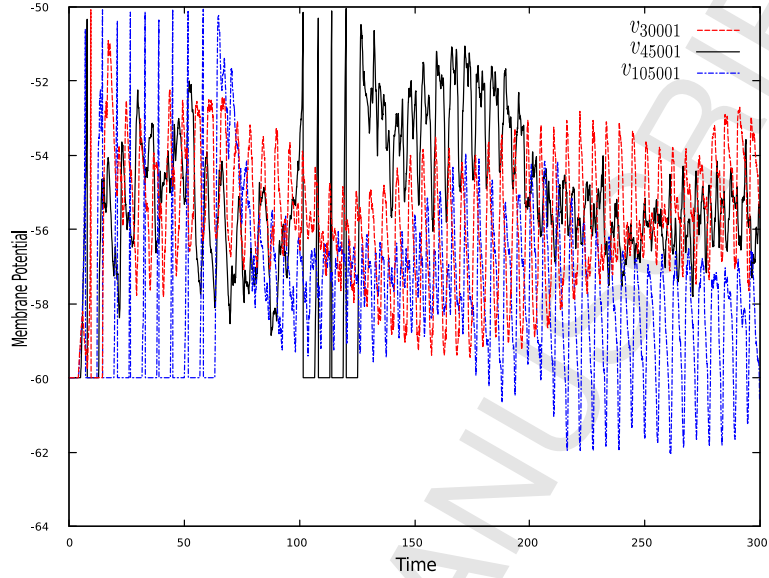
After a neuron spikes, it enters a refractory period during it does not change its membrane potential.

We considered a network of 300,000 neurons, with 80% of excitatory type and 20% of inhibitory type randomly distributed. Each neuron has 200 random post synaptic connections, limited to the 800 closest neurons.

Additionally, there is a set of 60,000 neurons that receive input spikes from an external source during the first 100 mili seconds.

The system has a total of 900,000 state equations, and 660,000 zero crossing functions. Example output trajectories for three neurons are depicted in Fig.7.

Figure 7: Spiking Neurons Model - Output Trajectories



We simulated this system using QSS2 algorithm with a tolerance  $\Delta Q_{rel} = \Delta Q_{abs} = 1e - 3$  until a final time  $t_f = 300$ .

This model has a chaotic behavior and a small change in the tolerance settings produces a totally different result if we look at the evolution of an individual neuron. Thus, rather than measuring errors, we measured the activity of the neuron network, counting the number of spikes performed during the simulation.

Due to this chaotic behavior, not only the errors but also the speedups change between different simulation runs under the same parameter settings. Thus, the results reported here are computed as the average over 10 experiments.

As in the previous examples, we started by varying the parameter  $\Delta t$  using 62 cores. The results are reported in Table 11

Table 11: Spiking Neurons Model - Fixed  $\Delta t$  - Using 62 Cores

$\Delta t$	Simulation Time (seconds)	Speedup	Average Spikes
1e-04	2201	2.45	15.80
1e-03	726	7.44	15.92
1e-02	410	13.17	15.92
1e-01	304	17.76	15.93
5e-01	227	23.79	15.87
1e+00	214	25.24	15.84
2e+00	216	25.00	15.62

As expected, the speedup increased with  $\Delta t$  reaching a maximum of about 25 times. Regarding the number of spikes, it did not experience noticeable changes, so the activity was similar.

Table 12 shows the speedup varying the number of cores using  $\Delta t = 0.5$ .

Table 12: Spiking Neurons Model - Fixed  $\Delta t = 5e - 1$

Cores	Init Time (ms)	Simulation Time (seconds)	Memory (MBytes)	Speedup	Average Spikes
1	1244	5402	2555	1.00	15.84
2	1806	3331	2916	1.62	15.86
4	1809	2030	3670	2.66	15.94
8	2071	1349	5177	4.00	15.85
16	2176	1012	8257	5.33	15.93
32	2594	439	14352	12.30	16.16
48	3104	288	20480	18.75	15.81
62	3536	231	25851	23.38	15.80

The initialization times, memory usage, and speedup show similar evolutions to those of the previous examples, and again, the speedup is almost linear.

The use of the adaptive algorithm varying the error bound parameter  $\alpha$  produces the results reported in Table 13

Table 13: Spiking Neurons Model - Adaptive  $\Delta t$  - Using 62 Cores

$\alpha$	Simulation Time (seconds)	Speedup	Average Spikes
2	388	13.92	15.94
10	344	15.70	15.90
20	315	17.14	16.02
50	252	21.43	15.75
100	236	22.88	15.84

This time, the best results are obtained for  $\alpha = 100$ . However, due to the chaotic behavior of the model, we cannot estimate the effects of the errors introduced by the lack of synchronization.

Using  $\alpha = 10$  and varying the number of cores, we obtained the results of Table 14.

Table 14: Spiking Neurons Model - Adaptive  $\Delta t$  -  $\alpha = 10$ 

Cores	Init Time (ms)	Simulation Time (seconds)	Memory (MBytes)	Speedup	Average Spikes
1	1244	5402	2555	1.00	15.84
2	1805	3367	2916	1.60	16.41
4	1806	2040	3670	2.64	15.50
8	1935	1358	5177	3.97	16.04
16	2196	1049	8257	5.14	15.79
32	2654	486	14352	11.11	15.77
48	3123	354	20480	15.25	15.93
62	3512	344	25851	15.70	15.84

## 7. Conclusions

In this article we presented novel methodologies for the parallel simulation of continuous time and hybrid systems using QSS algorithms. Also, we described their implementation in a software simulation tool.

The methodologies are based on the use of non strict synchronization between logical processes and introduce an additional numerical error to that of the QSS approximation. We showed that this additional error is bounded depending on a parameter  $\Delta t$ , and we designed an adaptive algorithm for dynamically computing that parameter in order to keep the error bounded according to the desired tolerance.

The results obtained on four large systems (one of them being purely continuous and the remaining having discontinuities) showed linear speedups with the number of cores. For a maximum of 62 cores used, the speedup reached values between 22 and 45 in the different cases. Taking into account that the sequential QSS simulations of those examples are faster than those of classic discrete time numerical algorithms, the parallel QSS results can increase the advantages.

This is the first work reporting parallel simulations with QSS methods in models with a size of the order of one million variables (accounting states and discontinuity handlers).

Regarding future work, we consider the following items:

- The results should be extended to a wider set of models, including in particular other types of PDEs.
- Although the tool includes the automatic partitioning algorithms of [45], they must be adapted and optimized according to the features of the this particular parallel implementation.
- In many problems, like in that of the Advection–Reaction equation, the results can be improved using dynamic load balancing. To this end, we plan to extend the tool implementing dynamic partitioning algorithms.
- In order to simulate using more LPs, we need to extend the techniques and the implementation to consider distributed memory architectures, where the synchronization strategy must be modified.

- We are currently working on the automatic generation of partitioned sub-models in order to facilitate the use of the model partitioned implementation, which is far more efficient in terms of memory consumption.

The Parallel Stand-Alone QSS solver is an open source project, available at the site <http://sourceforge.net/projects/qssengine>. The models used in this article are part of the distribution.

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- Novel parallelization techniques for Quantized State System (QSS) ODE simulation are proposed.
- A theoretical analysis of the parallelization error introduced is presented.
- The novel techniques are implemented on multi-core architecture.
- The presented implementation is deeply evaluated on four large scale models.