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Agents and artefacts for multiple models co-evolution

Building complex system simulation as a set of interacting models

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ABSTRACT

Complex systems simulations generally involve the interaction of different scientific fields. Human economies, ecosystems or dynamic computer networks such as P2P are good examples. Since models and simulators already exist in those fields, designing the simulation as a society of interacting and co-evolving models appears attractive. Beyond the technical issues to make different simulators cooperate, the challenges are to make the co-evolution design and implementation easier for the users that rarely know intricate modelling and simulation tools, and to facilitate the collaboration of different experts. Agents and artefacts (A&A) paradigm simplifies the design and the implementation of a society of interacting and co-evolving models. That is, the addition, the removal or the interchange of models require less effort. Contrary to classical approaches, we have built a decentralized co-evolution architecture based upon A&A and a data-driven coordination model. In this article, beyond the architecture presentation, we focus on the benefit provided by A&A used for multiple models co-evolution.

Categories and Subject Descriptors
I.6.5 [Simulation and Modelling]: Model Development

General Terms
Design

Keywords
Simulation techniques, tools and environments, Software engineering, Complex systems

1. INTRODUCTION

Modelling and Simulation (M&S) techniques and software engineering have always been in touch with each other. In [4] the authors declare that [...] the real limits on the future adoption of simulation may rest on our ability to represent complex systems and to do it easily, which can be constructed as a matter of modelling style. [...] Thus the distinction in programming style is not what can and can't be done but what can be done easily.

A complex system is composed of a set of interacting parts that as a whole exhibits properties that cannot be predicted from the simple sum of the individual parts properties. Human economies, social structures, climate or ecosystems are good examples of complex systems. Equation based modelling cannot represent interactions among components and their impact on the global system behaviour. Multiagent approach offers an interesting alternative [17].

Complex systems modelling also involves the interaction of different scientific domains or different abstraction levels. This is the case in biology, for example, where models from chemistry and biology (organ, tissues) may be needed [15] or in dynamic networks with models from sociology and computer networks [12]. This way, different specialists work on the same simulation. Each one brings its own models and simulators.

The challenge is then to allow those scientists to build a complex simulation from their own building blocks. Moreover, we should keep in mind that they are probably not familiar with the intricate modelling and simulation tools and theories. One way to facilitate the interaction of heterogeneous models is to build the simulation as a society of interacting models. Models should be seen as component we can weave together (as in component-based software engineering).

Some work has already been done in this way. The high level architecture (HLA) [5] is a standard that proposes to use a central infrastructure in order to make models interact. VLE [6] and JAMES II [3] are both modelling and simulation frameworks that allow to couple different models (issued from different formalisms). Agents and artefacts (A&A) paradigm [8] has been used in order to build a complex systems simulation as a society of interacting and co-evolving models [1].

In this paper we propose modelling engineering concepts to simplify the design of distributed and decentralized complex systems simulations. To do this, we use the perspective of A&A. However, contrary to approaches such as HLA or A&A used in [1], we do not rely on a global scheduler in order to synchronize the different interacting models. Instead, we propose a data-driven coordination model that allows a decentralized synchronization [13]. On the other hand, contrary to approaches such as VLE [6] and JAMES II [3] that imposes to build models directly in the modelling framework, we concentrate on reusing existing models and simulators and making them interact. Since our work fo-
cuses on model coupling level, innovations from the parallel and distributed computation domain may of course be used to implement our approach.

The outline is the following: in section 2, we present the challenges related to the coupling and the coordination of different simulators. In section 3, we propose a coordination model and an architecture based upon A&K. In section 4, we developed a proof of concepts. We use our concepts to make some existing Netlogo models interact. Finally, discuss the advantages and the limits of such an approach.

2. CHALLENGES AND RELATED WORKS

2.1 A modular approach

When we talk about complex systems, we usually think of decentralized, open and large scale systems. It is generally accepted that there is no central point of control. Actually, global properties emerge from the interaction of subsystems (or entities). Moreover, entities can enter and leave the system and their number is admitted to be large. A good example is P2P networks, in which there is no central point of control, people can connect and disconnect themselves on the fly and the size of these networks have reached several million users.

However, complexity occurs through the interaction of different abstraction levels. For example, prediction of P2P networks performances are difficult because it involves the interaction of socio-economic parameters (users willingness to share and to download data) and network parameters (bandwidth, download time). Models exist in both humanities and computer network sciences. Moreover, modelling a system as a set of interacting subsystems is simpler and requires a collaborating team of experts. Thus, it appears that, in order to predict the behaviour of a complex system such as P2P networks, it is necessary to couple heterogeneous models [12]. It may also be necessary to interchange a model by another one in order to compare simulation results. In the same way, a new instance of a model may be added in order to scale up the simulation.

We think that, in order to build a complex system simulation from heterogeneous models, specialists from the different concerned fields should be able to, firstly, reuse the existing models and simulators they develop with as little modifications as possible and, secondly, easily add, remove or intercange the models.

Existing modelling and simulation frameworks (such as JAMES II [3] or VLE [6]) imposes that every specialist involved in the simulation should know the framework and modify its own models in order to include them. Conversely, we think that models should be seen as component we can weave together (as in component-based software engineering). The challenge is then to modify as little as possible the existing models, to make them interact in such a way that it is transparent for the people involved in the simulation design.

In order to build concepts generic enough, we make several assumptions. Models are black boxes with input and output ports (as in [19], see figure 1) and there exist simulators functions that give us the current and the next simulation time (see section 2.3).

2.2 Exchanging data between simulators

When we couple different models the first thing to do is to represent the data flow between each model (as in figure 1). Since these models represent several dynamics at different scales and have been implemented independently, each model has its own representation of time and data. In the same way, each simulator proceeds its own execution. Coherence and compatibility issues appear when coupling different models and simulators. The next sections present the related challenges and issues.

2.2.1 Coherence

Scales or dimensions in which a piece of data is represented could be different from a model to another. Moreover the representation can be discrete or continuous. For example, a position pos1 = < x, y, z > (with x, y and z expressed in meters) in a first model can be represented only in two dimensions in a second one: pos2 = < x, y > (with x and y expressed in kilometers). A solution proposed in [1] is to define operations (projection, discretization, reduction) in order to achieve this coherence.

The same way, each model could have its own time representation. For synchronization purpose, we need to assure, for example, that a time value t1 ∈ R+ in a first model correspond to a time value t2 ∈ N in a second one. We propose to express an operation that makes the correspondence between both time values.

2.2.2 Compatibility

Each simulator could implement a single piece of data in its own way (integer, float, etc.) or some simulators may not implement all aspects of a given model. These challenges are discussed in [9]. A solution is to add an entity (a program) between the simulators. Its role is to translate the data in order to respect the compatibility between simulation tools.

2.2.3 Synthesis

Coupling different models and simulators implies to deal with coherence and compatibilities issues that are neither models, nor simulators purpose. Since we try to modify as least as possible the existing models, a solution that appears is to design of a new entity in charge of these issues (see figure 2). This is what is proposed in [1], where an artefact [8] is define in order to deal with these issues.

2.3 Coordination in co-evolution

The goal of time management in co-evolution is to ensure that simulation events (or steps) are executed in the correct
order (local causality constraint [2]). Two main approaches have been proposed to coordinate interacting simulators: optimistic and conservative [2]. We assume that the existing simulators we reuse have been developed independently and where not thought for distributed simulations. So they do not have a roll-back capability (see optimistic approach): they cannot go back into the simulation process in order to take new input events into consideration. As a consequence, we focus on the conservative mode. In the latter, the coordination model has to determine when a simulation event (or step) is safe to process.

**Definition 2.3.1.** For a model $M_i$, a simulation event (or step) associated with the current simulation time $ct_i$ is said to be safe to process if all the input events received after this event execution are timestamped with a time value $> ct_i$.

For example, in the figure 2, suppose the model $M_1$ is at simulation time $ct_1 = 8$. Its next event to process $ne_1$ is at time $nt_1 = 12$. This event $ne_1$ is said to be safe if no other event will be send to $M_1$ with a timestamp $< 12$. In our example, $M_1$ has to know if $M_3$ will send events with timestamps $\in [8, 12]$. If not, $ne_1$ is safe and $M_1$ can process it. This way, simulators stay synchronized and the causality constraint is satisfied.

Conservative coordination can be done by using a central and global scheduler that synchronizes all the simulators as in [1, 9, 5]. In these solutions, models and simulators need substantial modifications in order to be controlled by the scheduler. This prevents to easily reuse the existing M&S tools. In section 3, we remove this central scheduler and we propose a novel decentralized coordination.

## 3. PROPOSITION

We present the concepts used in order to build complex simulation as a society of interacting models. We do not target on-line nor real time simulations, which directly interact with the reality.

### 3.1 Goals

We use the A&A paradigm [8] to facilitate the design and the implementation of a society of co-evolving models. Our approach is intended to make the design and the implementation of such simulations transparent. This way the specialists involved in the simulation do not care about coordination issues and only focus on the modelling aspects. The architecture we propose is modular and decentralized. It requires as little modifications as possible to integrate existing models and simulators and, more important, these modifications are done once for all. Thus, to design a simulation of heterogeneous models becomes a building block game.

We propose to use objective coordination. That is, coordination does not rely on a single entity but is provided by the surrounding environment. This method is well known in the field of situated multiagent systems [16, 10] (stigmergy) or in parallel systems [11] (shared memory). This provides a way to loosely couple and to coordinate the interacting processes. In our case, the simulators interact through the set of data they exchange. Thus, their surrounding environment is composed of the coupling-artefacts between them and the set of exchanged data.

### 3.2 Validity interval and coordination

Our coordination model principle is the following, each model $M_i$ holds a current simulation time value $ct_i$ and knows the simulation time value for the next event (or next step) to be processed $nt_i$. That is, when a model $M_i$ is executed, it produces data $\delta_i$ at time $ct_i$. These data $\delta_i$ will not change until the next time $nt_i$ when the model $M_i$ is executed. As a result, we can say that $\delta_i$ are valid for the simulation time interval $\Gamma_i = [ct_i, nt_i[$.

A simulator can execute a model if the simulation event to process is safe (see definition 2.3.1). Then, the issue for the simulator is to know when an event is safe. It can be solve if the simulators exchange both the data and the corresponding validity interval: $< \delta_i, \Gamma_i >$. Indeed, a simulation event is said to be safe to process if $ct_i \in \Gamma_i$ for all the input data $(i \neq j)$.

Coordination is neither the simulator, nor the model purpose. It is the environment role. In our case, the coupling-artefacts besides being in charge of coupling issues (see section 2), are also in charge of coordination. That is, when data are transmitted through the coupling-artefact, the latter saves them (as a shared memory), then, when a model $M_i$ needs input data for the time $ct_i$, it asks the coupling-artefact for safe data (i.e. data $< \delta_i, \Gamma_i >$ that fulfills the condition $ct_i \in \Gamma_i$ (with $i \neq j$)). This way, models can interact and coordinate themselves in a loosely and decentralized way.

We have developed a formal version [14] of this coordination model and we have proved that coordination occurs between models and that the system is alive and deadlock free with $k$ models $(k \in \mathbb{N})$.

### 3.3 Architecture overview

Our architecture is intended to make the design and the implementation of a simulation of heterogeneous models transparent, modular and decentralized. In this section, we presents the concepts from A&A we use.

The coupling-artefacts are in charge of the coordination and the coupling aspects. The main contribution of our work is that data exchanges and coordination happen through the simulators environment (the coupling-artefacts) so the models execution can be decentralized. We previously define one coupling-artefact for each link between models (see figure 2). Data exchanged between two models flow through a coupling artefact (see figure 3).

We use two other kinds of entities to deal with separation of concerns. To format data (i.e to associate $\delta_i$ and $\Gamma_i$), to send data through the coupling-artefact and to ask for
Table 1: Artefacts functions summary

<table>
<thead>
<tr>
<th>Artefact</th>
<th>Functions</th>
<th>Role</th>
</tr>
</thead>
<tbody>
<tr>
<td>Model-Artefact</td>
<td>init()</td>
<td>Initialize $M_i$</td>
</tr>
<tr>
<td></td>
<td>run()</td>
<td>Run one simulation event, step or time interval</td>
</tr>
<tr>
<td></td>
<td>getOutputData()</td>
<td>Return output $\delta_i$ from $M_i$</td>
</tr>
<tr>
<td></td>
<td>setInputData()</td>
<td>Send input $\delta_j$ data to $M_i$</td>
</tr>
<tr>
<td></td>
<td>getCurrentTime()</td>
<td>Return $ct_i$</td>
</tr>
<tr>
<td></td>
<td>getNextTime()</td>
<td>Return $nt_i$</td>
</tr>
<tr>
<td>Coupling-Artefact</td>
<td>post()</td>
<td>Manage the end of the simulation (storing data, etc.)</td>
</tr>
<tr>
<td></td>
<td>read()</td>
<td>Return a safe data $&lt;\delta_i, \Gamma&gt;$</td>
</tr>
</tbody>
</table>

safe data is neither the simulator, nor the model, nor the coupling-artefact purpose. So we define an entity which behaviour will be to manage the data flows for a given model: the model-agent. Its role is to get the data $\delta_i$, the $ct_i$ and $nt_i$ values from the model $M_i$, to send the $<\delta_i, \Gamma_i>$ tuples to the output coupling-artefacts, to ask the safe data $\delta_j$ from the input coupling-artefacts and to send them to $M_i$. The model-agent behaviour is described in figure 4.

In order for the model-agent to interact with the models, and in order to modify as little as possible the existing models $M_i$, we define an interface that allows the model-agent to achieve its purposes: the model-artefact. It allows the model-agent to initialize, execute the model $M_i$ and to manage its input and output ports. The table 1 lists both the coupling and model-artefacts functions.

Figure 3: Architecture overview: from concepts to implementation

The whole architecture is described in the figure 3. It is modular, transparent and decentralized so the simulation can be seen as a set of distributed and reusable components. Models are in charge of modelling. The model-agents are in charge of executing the models. Finally, the coupling-artefacts are in charge of the coordination process. This way, we can easily add, remove or interchange models without being concerned with coupling and coordination issues (see section 4).

4. PROOF OF CONCEPT

The objective of this section is to present how to build a complex simulation as a society of interacting models. The simulation results validation is out of the scope of this article. We present a proof of concept illustrated by NetLogo [18] models co-evolution. We choose those simple models for the sake of illustration: they are straightforward to understand and we think they highlight some of the coupling and synchronization problems we raised in section 2. We discussed more evolved examples in section 5.

We implement our concepts in a framework called AA4MM\(^1\). It is coded in Java and rely upon the Java Messaging Service in order to deal with shared memory purposes. Since this framework, at first, has been developed in order to build a proof of concepts and since the discussion about implementation choices and performances is out of the scope of this article, we do not give a lot of details about this framework. Interested readers can find more details on the AA4MM web page.

4.1 A sheepfold study made of sheep, grass and wolves models co-evolution

Imagine we want to predict the impact of a sheepfold. For example: what are the best herding strategies? What is the risk for a sheep to be eaten by a wolf? etc. We want to model an ecosystem made of wolves, sheep, shepherds and grasslands. The first model $M_1$ represents sheep moving and shepherds trying to herd them. The second one $M_2$ represents sheep\(^2\) eating grass and gaining energy. The last one $M_3$, is a prey predator model that represents the wolf predation.

Our goal is to design the sheepfold simulation as a set of interacting and co-evolving models. We want to couple the three above-mentioned models using the concepts we present in section 3.

4.2 Integration basic steps

At first we only consider $M_1$ and $M_2$. Assume we are more

\(^1\)http://www.loria.fr/~siebertj/aa4mm/

\(^2\)Originally rabbits. We changed species.
preoccupied by the impact of sheep on the grasslands. We target the addition of $M_3$ later (section 4.4). Both specialists of sheep model and grassland model should be able to make their own models and simulators interact.

The first thing to do to couple different models is to define the input and output ports of each model and the connections between them. The sheep movements are described by the sheep model $M_1$. That is $M_1$ provides the set of sheep positions $SP$. In order to be executed, $M_2$ needs to update its own local sheep positions from $M_1$. On its part, the sheep movement is influenced by the sheep energy $SE$. Energy is provided by $M_2$. As a consequence, $M_1$ needs to update its own local sheep energy values from $M_2$. Figure 5 depicts both conceptual links between $M_1$ and $M_2$ and how it is implemented.

### 4.2.1 Existing models integration

The second thing to do to couple different models is to build the *model-artefacts* for each involved model. Each specialist have to create a *model-artefact* for its model. Consider $M_1$, its behaviour is given by the algorithm 1. The *model-artefact* in charge of $M_1$ must implement the functions listed in table 1.

The $init()$ function sets the number of sheep, shepherds and their initial positions in $M_1$. Since Netlogo processes simulation step by step, the $run()$ function executes one simulation step of $M_1$, the $getCurtime()$ and $getNextTime()$ returns $ct_1$ (the number of clock ticks) and $nt_1 = ct_1+1$. $getOutputData()$ function returns all sheep positions $SP$, $setInputData()$ sets into $M_1$ the sheep energy $SE$.

The same work is done for $M_2$ and $M_3$. All those functions are implemented by simply calling procedures through the provided API. As a consequence, we do not modify the original model. However, the simulators has to provide the mentioned functions ($run()$, $init()$ etc.). Thus, we modify the execution process.

### Algorithm 1: Model $M_1$ (one step)

| Input: Sheep energy levels $SE$ |
| Output: Sheep Positions $SP$ |
| Set the sheep energy levels from $SE$. |
| foreach Sheep Positions $SP$:
  | Shepherd $B$ moves randomly. |
  | if $B$ is not carrying a sheep. |
  | then Search for a sheep. |
  | else Find a herd to drop the sheep. |
| foreach Sheep $S$ not carried:
  | $S$ moves randomly. |
| return $SP$ |

#### 4.2.2 Coupling models

In our example, one coupling-artefact $cA_1$ is in charge of the sheep positions. The other one $cA_2$ deals with sheep energy (figure 5).

Consider $cA_1$, its $post()$ function allows $M_1$ for sending the sheep positions with their validity interval: $<SP, \Gamma_1>$. The $read()$ function provides safe sheep positions to $M_2$. We assume that both models $M_1$ and $M_2$ have the same representation of the sheep positions. We will see in section 4.3 how to deal with different scales and representation. It is worth noticing that the specialists involved in the simulation do not have to deal with coordination issues since the coordination model is already implemented into the coupling-artefact.

### 4.2.3 Model-agent behaviour

Once the artefacts are built, the last thing to do is to build the model-agents that will use them. One model agent is created for each model. The generic model-agent behaviour is presented in figure 4. In our example, agents have to create and to initialize their own model via the proper model-artefact functions $init()$. Then, they post the initial output data ($SE$ and $SP$ at time 0).

The agents manage the simulation process as follows.

1. Read the input data $SE$ (resp. $SP$) from the coupling-artefact $cA_2$ (resp. $cA_1$). Set them to the model one.
2. Execute the model $M_1$ (resp. $M_2$) (i.e call the model-artefact $run()$ function).
3. Get the current and next time values $ct_1$ and $nt_1$ (resp. $ct_2$ and $nt_2$).
4. Get output data $SP$ (resp. $SE$) from the model-artefact. Post them to the coupling one $cA_1$ (resp. $cA_2$).

The agents do this loop until the simulation ends.

#### 4.2.4 Synthesis

In this section, we presented a case study in which a simulation is built from three models. Notice that those models come from different domains, have been designed independently and can be ran separately. We assume that the specialists of each domain wish to couple those models in order to build a more complex simulation. In order to make those models interact they need to deal with issues such as coupling, coordination and model execution. However, they are not familiar with distributed simulations and coordination process.

The aim of our framework is to facilitate the creation of this simulation by making the coupling and coordination issues transparent. When the specialists need to couple models, they need to define the data flows between them and to implement a coupling-artefact for each link. They do not have to care about the synchronization process since the coordination model is already implemented into the coupling-artefacts and the exchanged data. A model-artefact has to be created for each model to interface. There are six functions to implement (see table 1). We rely upon the work in [19] in order to define and make these functions as generic as possible, anyway existing models may be adapted in order to match them. In the next sections, we show that, having done these simple modifications, it is straightforward to add or interchange models and also to deal with the interaction of models designed in different scales.

### 4.3 Dealing with scales differences

Until now, we have assumed that time and space scales in both models $M_1$ and $M_2$ were the same. In the reality, specialists design their models independently. Most of the time, the models scales are different. For example, the grassland specialists are interested into how fields evolve each season although sheep specialists are more concerned with how the sheep population evolves each week.
4.3.1 Different space scales

Consider that one patch of grass correspond to a square of 2 × 2 patches in the sheep model \( M_1 \) (see figure 6). Since it is not possible to change the patches size in the grass model \( M_2 \), sheep positions produced by \( M_1 \) does not fit anymore with space in \( M_2 \). As a consequence, we need to add an operation in the coupling-artefact \( cA_1 \) when the grass model \( M \) agent reads the sheep positions. This operation consists in dividing each sheep position coordinates by a factor 2.

4.3.2 Different time scales

The same way, consider that the grass model \( M_2 \) may no longer be executed step by step but 2 steps by 2 steps; while the sheep model execution remains step by step (see figure 7). This is related to the model \( M_2 \) execution process. So we modify the related model-artefacts functions \( run() \) and \( getNextTime() \). That is, instead of executing only one simulation step, the \( run() \) function executes two simulation steps. Then the \( getNextTime() \) function returns \( n_{t_2} = c_{t_2} + 2 \). Since the coordination occurs only thanks to time values given by \( getCurrentTime() \) and \( getNextTime() \) functions, these are the only modifications to make.

4.4 Adding, or exchanging models

In this section, we present the advantages to consider model as component and the simulation as a distributed set of interacting models.

4.4.1 Simple addition

The sheep and grass specialists involved in the sheepfold simulation wish to predict the impact of wolves living in the neighbourhood. As a consequence, a new model \( M_3 \) should be added in the simulation. The challenge is to couple a new model to the previous ones without rebuilding the whole simulation (see figure 8).

The first thing to do is to define the new links between three the models. The specialists have to know which data are exchanged between \( M_1 \), \( M_2 \) and \( M_3 \). Two cases are possible. Either the exchanged data are the same as those existing or they bring into play new ones. On the one hand, we can reuse the existing coupling-artefacts. On the other hand, we need to create new coupling-artefacts (as described in section 4.2).

Consider \( M_3 \), this models defines the wolf predation dynamics: wolves move, catch a sheep and took energy from the sheep (as they eat them). This way, \( M_3 \) needs to update its own local sheep position values from \( M_1 \) and \( M_2 \) provides negative sheep energy values \( SE_3 \) when they eat the sheep. The links between the models are described in figure 8. Since they do not require some new data to exchange, we can reuse the existing coupling-artefacts \( cA_1 \) and \( cA_2 \).

Since it is already present, \( M_3 \) can ask the coupling-artefact \( cA_1 \) to give it the sheep positions \( SP \) from \( M_1 \). The only thing to do is to add an operation to \( cA_1 \) in order for \( M_3 \) to read that value. However, when \( M_1 \) ask \( cA_2 \) to give it the sheep energy values, it has to take into account both the positive values \( SE_2 \) given by the grass model \( M_2 \) and the negative values \( SE_3 \) given by the wolf model \( M_3 \). That is, we need to define a new operation in \( cA_2 \) in order to combine both energy levels. In our case, for a given sheep energy level, \( cA_2 \) needs to add the energy gain \( SE_2 \) and the energy loss \( SE_3 \). This way \( M_1 \) updates a single sheep energy value for each sheep.

4.4.2 A (little bit) more complicated addition

As we saw in section 4.3, \( M_1 \), \( M_2 \) and \( M_3 \) may have been designed independently and, as a consequence, scales in each model may be different. For example, the grassland model...
represents fields evolution for each season, the sheep model represents the sheep population evolution for each week and the wolves model represents the wolf predation techniques for each day. Assume that we couple the three model as in figure 8.

There is no need to care about coordination since the coordination model is already implemented into the coupling-artefacts and the exchanged data. The only modifications to make concern the run() and getNextTime() functions as in section 4.3.2.

In our example, the coupling-artefact $cA_1$ needs to provide sheep positions from $M_1$ to both models $M_2$ and $M_3$. Since the sheep positions in these models are different, $cA_1$ need an operation to translate sheep positions from $M_1$ to $M_2$ (as in section 4.3.1) and, in the same way, another one that translate sheep positions from $M_1$ to $M_3$.

Note that, since the coupling-artefacts store the data sent by the model-agents it is possible to define as many operations as needed in order to deal with the coupling issues (see section 2). As a conclusion, if works from sections 4.3 and 4.4.1 have already been done, no more modifications are needed in order to build the new coupling scheme.

### 4.4.3 Interchange

After some measurements, one may find that a model is not accurate enough and that another one should be better. In order to interchange models, the simulation designers simply need to check that they will have the same input and output ports. For example, in $M_1$, instead of moving randomly, the sheep can move accordingly to a flocking model (as [7]). Instead of rewriting a new model, we can simply replace $M_1$ by an existing flocking model $M'_1$ that have the same input and output ports. This way, we do not change the model-agents nor the coupling-artefacts. We only need to build a model-artefact for the new model $M'_1$ (see figure 9).

![Figure 8: Addition of a third model](image8)

![Figure 9: Interchanging models](image9)

### 4.4.4 Building new simulations

Imagine now that we are more interested in reintroducing wolves into the wild than in the sheep industry. However, we want to reuse the existing models and results we previously obtained. That is, we want to build new simulations from the same building blocks. In addition, new models may be needed. In order to build this simulation, we do not have to rewrite neither all the models, nor the coordination process. We only have to take the building blocks and to link them properly. This is one of the advantages to see simulation as a set of interacting and co-evolving models, and to have the concepts that allows to easily implement it.

Indeed, it is worth noticing that none of the model-agent needs to know each other, neither they know which reads the data they post. This simplifies the addition, the removal or the interchange of the models. The important thing is to check the dependencies between the models. That is, the interacting models should have the right input and output ports. Then, it is straightforward to implement new coupling-artefacts, to reuse the existing ones and to link them with the proper model-agents.

5. CONCLUSION

#### 5.1 Summary

In this article, we present an architecture to build simulations as a society of interacting and co-evolving models. It is based upon the A&A paradigm. We also propose an objective coordination model with decentralized models execution. Our concepts facilitate the building of such simulations since it makes the coupling and coordination issues transparent for the users. This way, people involved into the simulation design only have to care about modelling and about the data flows between models.

The architecture we propose is modular and decentralized so that any model could be added, removed or interchanged with as little modifications as possible. We have developed a formal coordination model and proved that synchronization happens without deadlocks for a number of simulators $k \in \mathbb{N}$. We implemented our concepts in order to build a proof of concept illustrated by several examples.

#### 5.2 Discussion and future directions
In this article, we have made the assumption that the complex system studied is divided into several abstraction levels. One may want a geographical division where the subsystems represent significantly different geographical subparts. As a consequence, entities can enter and leave the models or go from a model to another. This point of view raises new modelling and coupling challenges. We think the concepts presented in this article can help to solve these new issues.

We use this framework on an ongoing work on co-evolution of mobile ad hoc networks and social models. We then couple an event-based simulator (network model) with a step-by-step simulator (agent based model). It is part of a french ANR research project. We plan to use some of the experiments to improve and study the implementation performances.

Although the examples presented are quite simple and broadly compatible (both discrete event), we though that with the concept presented here we can be used to couple equation based models (EBM) with agent based ones. Technically, we need to resolve (or \( \text{run()} \)) EBM for time intervals instead of events or steps. It is possible to go from the micro scale to the macro one by using operations (as defined in section 2) such as sums, means or distributions, etc. ; the other way round by, for example, introducing randomness. Discrete event formalism and the work in [19] should be a good line of thinking in order to look further.

Another open issue would be in the validation process of the virtual laboratory environment: A multimodelling case study and users' guide. In MABS, pages 10–25, 1998. ACM.


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6. REFERENCES


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