

# EXPPO: EXecution Performance Profiling and Optimization for CPS Co-simulation-as-a-Service

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## Abstract

A co-simulation may comprise several heterogeneous federates with diverse spatial and temporal execution characteristics. In an iterative time-stepped simulation, a federation exhibits the Bulk Synchronous Parallel (BSP) computation paradigm in which all federates perform local operations and synchronize with their peers before proceeding to the next round of computation. In this context, the lowest performing (i.e., slowest) federate dictates the progression of the federation logical time. One challenge in co-simulation is performance profiling for individual federates and entire federations. The computational resource assignment to the federates can have a large impact on federation performance. Furthermore, a federation may comprise federates located on different physical machines as is the case for cloud and edge computing environments. As such, distributed profiling and resource assignment to the federation is a major challenge for operationalizing the co-simulation execution at scale. This paper presents the Execution Performance Profiling and Optimization (EXPPO) methodology, which addresses these challenges by using execution performance profiling at each simulation execution step and for every federate in a federation. EXPPO uses profiling to learn performance models for each federate, and uses these models in its federation resource recommendation tool to solve an optimization problem that improves the execution performance of the co-simulation. Using an experimental testbed, the efficacy of EXPPO is validated to show the benefits of performance profiling and resource assignment in improving the execution runtimes of co-simulations while also minimizing the execution cost.

## Keywords:

resource management, cyber-physical systems, distributed simulation, cloud computing, latency-sensitive, performance, gang scheduling

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## 1. Introduction

Cyber-physical systems (CPS) such as those commonly found in smart city, smart manufacturing, and transactive energy systems must make time-sensitive control decisions to ensure their safe operations. However, such CPS are an amalgamation of multiple dynamic systems such as transportation systems, vehicle dynamics, control systems, and power systems with interconnected networks of different scales and properties. The assurance that such complex systems are safe

and trustworthy requires simulation capabilities that can rapidly integrate tools from multiple domains in different configurations. Co-simulation is an attractive option for interlinking such multiple simulators to simulate higher-level, complex system behaviors.

The IEEE 1516-2010 High Level Architecture (HLA) defines the standardized set of services offered to a process in a distributed co-simulation [1]. A co-simulation in HLA comprises individual simulators called federates that are grouped into a logical entity called a federation. Each federate can have diverse computation and networking resource requirements, which must be considered when assigning resources to the simulators when the federation is deployed to cloud-fog-edge computation environments. The wall-clock execution time required for each federate's computation step varies based on this resource allocation. The variation in execution times can result in some federates waiting on oth-

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ers before they can proceed to the next stage of computation. Federates that require more wall-clock time for their computation steps (the low performing federates) increase the overall completion time (or *makespan*) of the entire simulation. This can potentially violate the simulation completion deadline. Thus, resource allocation and resource configuration selections are very important for the overall performance of distributed simulations.

Although co-simulations have traditionally been hosted in high-performance computing (HPC) clusters, there has been an increasing trend towards the adoption of cloud computing for simulation jobs. It is in this context that the Docker container run-time platform [2] provides a solution for running federates across different computation platforms by providing a unified packaging of simulation code with its software dependencies. But recent research [3] has shown that there are operational challenges, such as performance aware resource assignments, that need to be considered for running simulations in cloud environments. Furthermore, there are several infrastructure-related complexities that must be considered to run these distributed simulations in a cloud environment.

In this work, we focus on the problem of deployment of the CPS co-simulations in a cloud computing environment which optimizes resource allocation to the co-simulations such that they have the lowest makespan and execution cost. This work presents a performance profiling, simulation run-time optimization and resource configuration platform called EXPPO - (*EXecution Performance Profiling and Optimization*). EXPPO uses distributed tracing [4] to assess federate level performance for each computational step. These performance characteristics are used at runtime to determine whether changes to the resource allocation of the federation could enable shorter makespan for the simulation run. To enable distributed tracing, EXPPO utilizes the *opentracing* [5] specification for measuring the wall-clock time spent in each computational step of the simulation. To shield the developers from complexities when embedding tracing code in the simulation application logic, EXPPO leverages generative aspect of Model Driven Engineering (MDE) to auto-generate source-code snippets and configuration files for the simulation run. To provide resource recommendations for the individual federates, EXPPO uses the tracing information to build a resource-performance model and solves an optimization problem to find the resource allocation with the lowest makespan and cost for the co-simulation. For this paper we complement and build upon our previous work presented at [6]:

The main contributions of EXPPO are:

1. Demonstration that the default resource configuration for a federation has scenarios where a federate with a longer wall-clock execution time for its computational step increases the makespan of the co-simulation;
2. Development of an approach to generate tracing probes inside the source code of the simulation logic, which is required to perform distributed simulation profiling;
3. Development of a new resource recommendation engine which uses an optimization algorithm for finding the resource configuration for a federation that minimizes its overall makespan and cost; and
4. Validation of EXPPO showing its benefits on the performance of distributed simulation execution.

The rest of the paper is organized as follows. Section 2 provides a motivating use case for EXPPO and lists its key requirements. Section 3 provides an overview of the EXPPO framework. Section 4 describes the key EXPPO components and its resource configuration and optimization algorithms. EXPPO's cloud architecture and co-simulation framework are described in Section 5. The experiment evaluation results are described in Section 6. Related works are described in Section 7, discussion on running co-simulations in cloud computing environments is provided in Section 8 and Section 9 concludes the paper.

## 2. Motivation and Solution Requirements

The computation pattern of many time-stepped co-simulations follows the Bulk Synchronous Parallel (BSP) model [7]. In this model, every participating simulation completes its computation for a given time step, waits for the other participating simulations to complete their computations, exchanges new state information with its peers, and only then proceeds to the next time step. Thus, if one simulation takes more time to execute, the other simulations are forced to wait for it and remain idle. This is illustrated in Figure 1 which shows an HLA federation with three federates. Each federate executes one computation task repeated each time step. Federate 1 takes the longest to execute, and the other two federates are waiting for Federate 1 to complete before moving to the next computation step. This increases the makespan of the entire simulation, thereby decreasing the performance of the simulation execution.

Recently, co-simulations have been deployed using container management solutions [8] [9], such as Docker

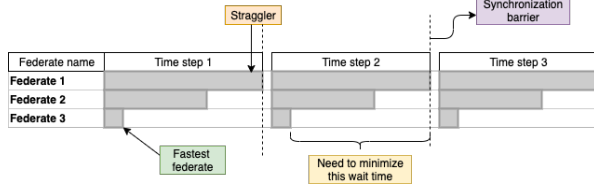


Figure 1: Performance visualization of an example federation.

Swarm [10], Kubernetes [11], Mesos [12]. However, these solutions use queue-based scheduling, wherein the containers are allocated to machines one at a time in sequence. Hence, there could be instances where the cluster runs out of resources to schedule all the federates of the federation. In these instances, a few federates are deployed while the remaining federates are stuck in the scheduler queue until more resources become available. This will cause the entire simulation to stall, since all the federates are required to participate in the simulation to progress. For instance, from Figure 1, if there are resources to deploy Federate 1 and Federate 3, and not sufficient resources to deploy Federate 2, the job scheduler should not deploy any federates and should wait until more resources are available. This requires a bag-of-tasks scheduling mechanism for deploying these federates on the cloud computing environments.

The resource configuration also plays an important role in the execution time of the federates. Figure 2(a) depicts the performance of a federate when assigned different numbers of cores for executing a computation step. This federate is running a *Freqmine* application from the Princeton Application Repository for Shared-Memory Computers (PARSEC) benchmark [13] as its computation task. This figure shows that the execution time generally decreases with the increasing amount of resources assigned to the federate. Thus, there is a potential for minimizing the wait time by appropriately configuring the resources assigned to the federates. Minimizing the wait time can be done either by providing the highest resource configuration for all the federates, or finding a resource configuration that considers the cost of assigning the resources to the federates. However, deciding what resource configuration to select for a given computation is a non-trivial task for a simulation developer who may not have the domain expertise of configuring and running applications in cloud computing environments. Performance profiling of the federates can help in understanding the relation between the resource assignment and the execution performance. However, these simulations might be deployed across different physical and/or virtual host envi-

Property	Type I	Type II	Type III
Floor Area ( $m^2$ )	55.7	97.5	143.9
Fixed Plug Load (W)	800	1250	1750
Solar Panel Area ( $m^2$ )	13.9	24.4	35.9
Indoor Set Point ( $^{\circ}C$ )	20	22	24
Heat Pump (KW)	1.57	1.57	1.57
Solar Cell Efficiency (%)	19	19	19

Table 1. Configuration used for different type of houses

Symbols	Definition
$\gamma$	Solar Insolation ( $W/m^2$ )
$\mu$	Solar Heat Gain Coefficient (scalar)
$HA$	House effective area exposed to Sun ( $m^2$ )
$\omega$	Conversion constant (scalar)
$\kappa$	Lumped Thermal Conductance ( $W/^{\circ}C$ )
$p$	Solar Panel Area ( $m^2$ )
$\tau$	Time Constant of heating/cooling (sec)
$T_{in}$	House Indoor Temperature ( $^{\circ}C$ )
$T_{out}$	Outdoor Temperature ( $^{\circ}C$ )
$t$	Simulation time (sec)
$P$	Electrical energy transfer (W)
$Q$	Thermal energy transfer (W)
$hp$	Power Consumed by Heat pump (W)

Table 2. Symbols used in the equations of case study and their definitions

ronments when running in cloud computing platforms; performance profiling for such distributed simulations can be very challenging.

### 2.1. Co-simulation Use Case

This section presents a use case of a complex CPS scenario executed as a co-simulation to analyse and answer questions about the system and its components. This experiment highlights the challenges in co-simulation that the EXPPO framework can help address. The goal for this simulation is to understand the power generated by installation of solar panels in a community and answer to questions "How much power we can generate by installing solar panels in this community? After installing the solar panels, can the community become a net-zero-energy community?".

The community consists of 30 houses of three different types, where each type consists of 10 houses. Type classifications are based on the square feet area and the corresponding plug load of the house. The details of configuration used for different type of houses is shown in Table 1. Before the simulation starts, the indoor temperature of each house is a random variable, which is drawn from a uniform distribution between 17 to 32  $^{\circ}C$  i.e.,  $T_{indoor,initial} \sim Uniform(17, 32)$ .

The operation of the cooling system depends upon the indoor temperature. The cooling system turns on when

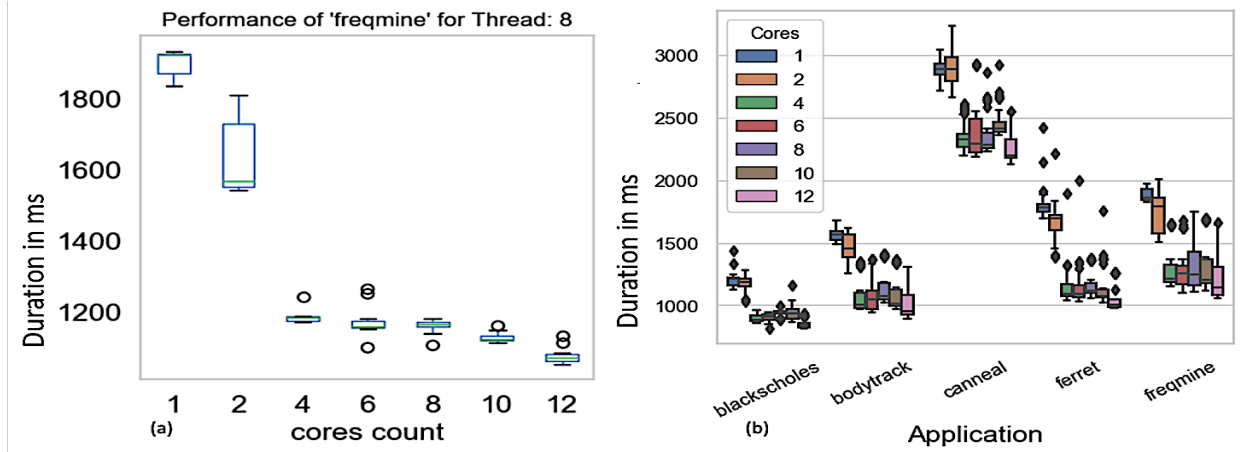


Figure 2: (a) Performance of the federate running the PARSEC Freqmine application using 8 threads for different resource configuration selections. (b) Performance of five PARSEC benchmark applications for different resource configuration selections.

the thermostat measures an indoor temperature greater than sum of the indoor set point and the thermostat hysteresis value. The model used for the simulation follows thermodynamics and electrical laws with some approximations.

Thermodynamics of a house depend upon two factors, solar insolation and the cooling system. The solar insolation has positive heat flux density and increases the indoor temperature while the cooling system has negative heat flux density and decreases the indoor temperature. Thermodynamics effect of solar insolation on the house depends on the amount of solar insolation, house effective area exposed to sunlight and solar heat gain coefficient. Solar insolation has a dual role in this simulation, both increasing the temperature of the house and generating electrical power using solar panels. Refer table 2, for definition of all symbols used in this case study. Thermodynamics and electrical laws related to the solar panels are:

$$Q_{solar} = \gamma * HA * \mu \quad (1)$$

$$P_{solar} = -\gamma * \omega * p \quad (2)$$

The cooling system is based on heat pumps that move thermal energy in the opposite direction by absorbing heat from a house and releasing it to the outside atmosphere. This process is opposite to the second law of thermodynamics so external power is needed to accomplish the work of transferring energy from the heat source to the heat sink. This results in the consumption of electrical power when the air is cooled. When cooling indoor air, the cooling system needs to cool both air and the water present in the atmosphere. Both sensible heating (cooling of air) and latent heating (cooling

of water) are approximated assuming 50 % relative humidity, air density of  $0.624 \text{ m}^3/\text{kg}$  with an air flow rate at  $0.378 \text{ m}^3/\text{sec}$ .

The thermodynamics and electrical equations related to cooling system are approximated to:

$$Q_{pump} = -(100.3 * T_{out}) + 3097.8; \quad (3)$$

$$P_{pump} = hp \quad (4)$$

The overall effect of solar insolation, solar panels, the cooling system and plug load can be summarised as:

$$Q = Q_{pump} + Q_{solar} \quad (5)$$

$$P = P_{pump} + P_{solar} + P_{plugload} \quad (6)$$

Here, Q is the net exchange of thermal energy (heat) between house and the outdoor environment and P is the net power consumption/production by the solar panel and utilities cumulatively (-ve value indicates production and +ve value indicates consumption). Overall change in the indoor temperature depends on net heat transfer due to difference between indoor and outdoor temperature, net heat transfer by solar insolation and the cooling system. For calculating the indoor temperature at any point in simulation, the rate of change of temperature is assumed to have a linear relationship with the heat transfer. The following equation guides the indoor temperature at any point in simulation:

$$T_{in} = T_{out} + Q/\kappa + (T_{in} - T_{out} - Q/\kappa) * e^{-t/\tau}$$

where  $T_{in}$  and  $T_{out}$  are indoor and outdoor temperature respectively in  $^{\circ}\text{Celsius}$ .

This scenario is derived from [14]. The interaction model among the federates is shown in Figure 3. Total

number of federates in the simulation is 34 (30 houses, 1 metronome, 1 weather source, 1 utility, and 1 federation manager). The simulation runs for a total of 150 hours of simulation time, with a logical time step of 5 minutes.

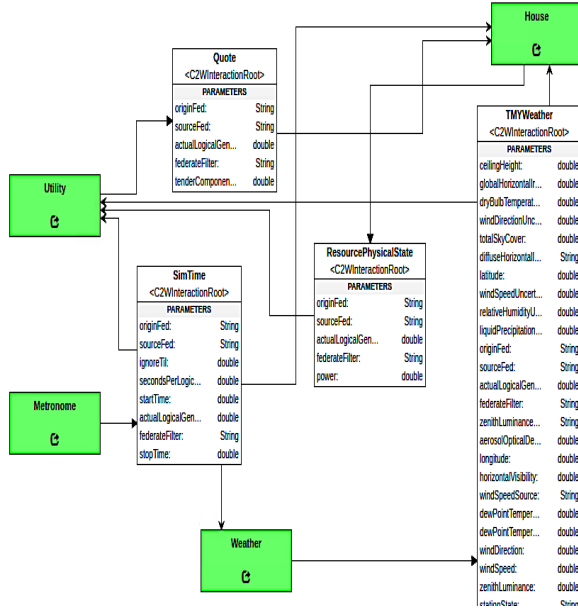


Figure 3: WebGME interaction model with one federate of each type. In simulation, there are 30 instances of the house federate with different configurations.

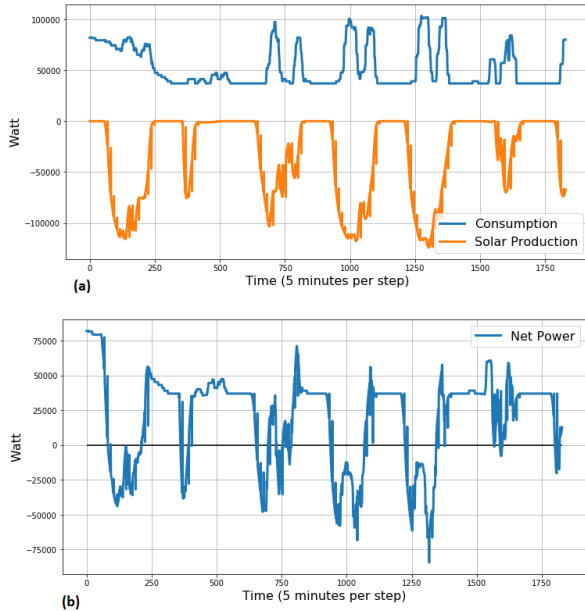


Figure 4: Consumption and solar production (a) and net power (b) over time for the simulated use case

The results of the simulation outcome are shown in Figure 4. Figure 4(a) shows the power consumed by all 30 houses and the solar power generated by the solar panel installed in the community. Figure 4(b) represents the net power (consumption - production) for the community. Simulation results show that total electrical power consumption by the community is 632 KWh and total electrical energy production by solar panel installed at community is 401 KWh for total duration of simulation. The net power for this configuration is negative as there is more consumption than production. By installing solar panels in 1/4th of floor area at each house, we can reduce the net power consumption but we are still in deficiency of power production. With this configuration, it is not possible to become a net zero community. This deficiency can be fulfilled by either further adding some more solar panels or getting extra power from utilities.

**Lessons Learned:** However, in this study we realized that if we had to scale the community to large number of houses greater than the current selection of 34 simulation federates, our compute system (13 GB RAM, Intel i7-8 core processor, 3.6GHz) on which the co-simulation experiment was running was insufficient to meet the demands of the computation load of the federates leading to drastic performance and the makespan degradation of the simulations. Also, due to the insufficient system memory resources, our simulation experiment would encounter memory buffer-overflow errors leading to failure of the entire simulation execution. Furthermore, in such kind of simulation study, there is a need to perform different actions to the inputs of the simulations, generating adhoc events to the running simulations or performing changes to the simulation world environment based on observation of certain events happening inside the simulation. Conducting all such variants of experiments requires running many counts of experiments which will be time consuming if ran in a sequential manner in a computing platform. Thus there is a need for running such experiments in parallel thereby reducing the overall makespan of the completion of the experiments. Cloud computing offers the ability to run such experiments at scale, thus there is a need to leverage cloud computing resources to run these CPS co-simulations.

## 2.2. Requirements for EXPPO

Building on the above use case, below are the four key requirements EXPPO aims to satisfy:

- **Requirement R1.** *Conduct distributed performance tracing of the federates:* To understand the

bottlenecks in federation performance, there is a need for logging and gathering execution performance traces of the federates. The tracing infrastructure needs to handle federates which are distributed across multiple physical hosts. Hence, the tracing infrastructure should be able to correlate traces from different federates of a federation.

- **Requirement R2.** *Reduce complexity in provisioning software probes:* Requiring the developer to manually write source code for performance tracing for the federation can result in complexities and errors which need to be minimized. The developer needs to understand the tracing software and write code which adheres to the tracing software requirements. This is tedious for the developer, who now apart from writing the simulation logic must also setup and configure the tracing infrastructure. EXPPO should reduce the manual configuration of the tracing information, thereby reducing repeated effort by the developer.
- **Requirement R3.** *Recommend resource configuration to minimize the co-simulation makespan and cost:* It can be challenging for an end user to determine the resource requirements for a federation because each federate can be configured differently. A bad resource configuration can have inadvertent effect on the completion time of the simulation. However, the choice of resource configuration also has an associated cost. Hence, the configuration must be chosen such that it satisfies both quality of service (QoS) and cost factors.
- **Requirement R4.** *Provide a gang-scheduling algorithm for executing simulations:* The runtime platform should support deployment of multiple federate using bag-of-tasks scheduling (or gang scheduling) algorithm.

### 3. Overview of EXPPO

Figure 5(a) shows the workflow and components of EXPPO.

The *design phase* requires the developer to model the federation using the federation development toolkit. This toolkit is based on the Web-based Generic Modeling Environment (WebGME) [15]. To measure the execution time of a federate in a time step, the simulator needs to embed a tracing code initializer and logger to record the execution time completion of the computation step. The tracing initialization code and the

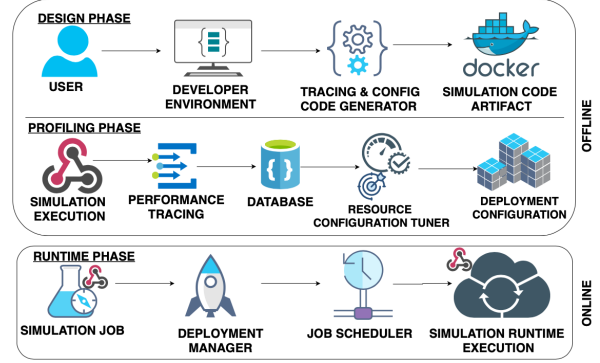


Figure 5: Workflow of EXPPO illustrating the connections between different components of the system.

tracer configuration files are auto-generated by the custom WebGME model interpreters. Once the required code is generated and the user has implemented the necessary simulation logic, the federate is compiled into an executable image, which is then packaged inside a Docker container.

During the *profiling phase*, each federate is executed and profiled under different resource configurations. The execution time for each federate is logged using the tracing information, and this tracing information is stored in a centralized database. The resource configuration tuner uses the recorded logs together with user provided objectives to optimize the resource configuration for each federate. Finally, the optimized resource configuration is used to configure the co-simulation deployment accordingly.

During the *runtime phase*, the simulation job information for the federates in the co-simulation is submitted to the deployment manager. The job information includes the name of the federate Docker image, resource configuration requirements, etc. The deployment manager submits the scheduling information to the job scheduler, which handles the execution of the jobs on the co-simulation runtime execution platform.

### 4. Design Elements of EXPPO

EXPPO allows users to design and deploy co-simulations on distributed compute infrastructures supported by Docker-based virtualization. Its generative capabilities simplify the auto-generation of performance monitoring instrumentation, configuration and probing for the different federates. Its resource configuration tuner optimizes resource allocations to federates to lower the makespan and execution cost for the federation execution. Its runtime platform supports parallel



execution of different federations on its shared compute infrastructure. This section details each of the EXPPO components.

#### 4.1. Performance Profiling of Federates

Understanding the performance characteristics of the co-simulation execution is of paramount importance when deciding how to run federates in runtime execution environments. Individual federates have different resource needs, and their execution performance will vary depending on how the resources are assigned. Thus, understanding the performance profiles of each federate is critical to the problem of optimizing the resource allocation and thereby lowering the makespan and execution cost of the federation execution. EXPPO leverages distributed tracing to assist in the logging of time stamps of distributed events generated in the federation (Requirement **R1**). It leverages *Opentracing* instrumentation [5] to track execution time spent during each computation time step of the federate. The execution time is then logged into a time-series database for conducting performance analysis. It leverages MDE technologies [16] such as Domain Specific Modeling Language (DSML), code generators and model interpreters to synthesize performance profiling software artifacts which can be used for performance profiling of federates (Requirement **R2**). An example code snippet is shown in Listing 1.

Here in Listing 1, we see that at line 1, we first initialize and activate the *Opentracing*'s profiling software module for monitoring the federate execution time performance. The profiling consists of monitoring the execution time at computation phase(*ExecuteState*) as well as waiting phase(*WaitState*) at the barrier synchronization. Each phase(also called as *Span*) has a start time and end time associated with it which is used for performance monitoring for individual phases. Lines 4-10 monitors the execution time for *WaitState*. Lines 11-17 monitors the execution time during the *ExecuteState* of the federate operation.

```

1  try (Scope scope1 =
      fedtracer.startFederateSpan("TimeStep"))
2  {
3    scope1.span().setTag("timestep",currentTime);
4    try (Scope scope3 =
          fedtracer.startFederateSpan("WaitState"))
5    {
6      scope3.span().setTag("timestep", currentTime);
7      atr.requestSyncStart();
8      enteredTimeGrantedState();
9      scope3.close();

```

```

10 }
11 try (Scope scope2 =
      fedtracer.startFederateSpan("ExecuteState"))
12 {
13   scope2.span().setTag("timestep",currentTime);
14   scope2.span().setTag("customTag",this.cmd);
15   executeComputation();
16   scope2.close();
17 } catch (InterruptedException e){
18   e.printStackTrace();
19 }
20 }

```

Listing 1: Profiling code snippet in Java language generated leveraging the MDE techniques.

#### 4.2. Federation Resource Configuration Optimization

When running a federate in a Docker container, cloud providers usually have multiple resource configurations from which the user can select for their application. However, without analyzing the resource dependency of the application, it may be challenging for the user to select the resource configuration that meets the application's quality of service (QoS) requirement while at the same time minimizing the execution cost in terms of the cost of resources utilized. Figure 2(b) shows how the different resource configurations impact the execution time of the federate which is running applications from the PARSEC benchmark. Furthermore, in a co-simulation, the resource selection becomes critical as every federate's execution time will contribute to the co-simulation's performance. To address this issue (Requirement **R3**), EXPPO provides a resource configuration recommendation system that selects the resource assignment which optimizes the application's QoS performance and user's budget requirements.

Consider the following optimization problem: Suppose the system has a set of homogeneous machines, each with  $k$  processing cores. Let  $F = \{f_1, f_2, \dots, f_n\}$  denote a federation (co-simulation) that consists of a set of  $n$  federates. Each federate is assumed to consist of many steps, and the execution time of each step is a function of the number of allocated cores for the federate (i.e., the federate is a parallel application). Formally, a resource assignment  $R$  is defined as a set of resources (CPU cores) assigned to the federates. Given a resource assignment  $R = [r_1, r_2, \dots, r_n]$  to each federate, where  $r_j \in R$  denotes amount of resources (CPU cores) assigned to federate  $f_j \in F$ , the execution time of federate  $f_j$  can be expressed as  $t_j(r_j)$  when assigned  $r_j$  cores for executing a single time step of the simulation. For performance reasons, assume a federate can-

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**Algorithm 1: Resource Configuration Tuner**

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**Input :** Execution time  $t_j(r)$  for each federate  $f_j$  in federation  $F$  when allocated different amounts of resources  $r$ , where  $1 \leq r \leq k$ .  
**Output:** A resource assignment  $R^* = [r_1, r_2, \dots, r_n]$  for each federate in the federation that minimizes a linear combination of makespan and cost.

```
1 Initialize  $r_j \leftarrow 1$  for all  $1 \leq j \leq n$ ;  
2 Compute  $G \leftarrow (\max_j t_j(r_j)) \cdot (\alpha + \beta \sum_j r_j)$ ;  
3  $R^* \leftarrow [r_1, r_2, \dots, r_n]$ ;  
4  $G^* \leftarrow G$ ;  
5 while  $\sum_j r_j < nk$  do  
6    $j \leftarrow$   
   Index of a federate with longest exec. time;  
7   if  $r_j = k$  then  
8     break;  
9   else  
10    Increment  $r_j$  to the next higher profiled  
    resource amount;  
11    Update  $G \leftarrow (\max_j t_j(r_j)) \cdot (\alpha + \beta \sum_j r_j)$ ;  
12    if  $G < G^*$  then  
13       $R^* \leftarrow [r_1, r_2, \dots, r_n]$ ;  
14       $G^* \leftarrow G$ ;  
15    end  
16  end  
17 end
```

---

not be split among two or more machines, so we have  $1 \leq r_j \leq k$ . The makespan  $M$  for every computation step for the entire federation  $F$  is dictated by the slowest running federate (i.e., the straggler), and is defined as  $M = \max_j t_j(r_j)$ . The execution cost  $C$  is given by the total resource used by all the federates over the makespan duration. Since the cores will be reserved for the federation until the slowest federate is done executing, the cost is defined as  $C = M \cdot \sum_j r_j$ . The resource configuration recommender needs to find a resource assignment  $R^*$  that minimizes  $G = \alpha M + \beta C = M(\alpha + \beta \sum_j r_j)$ , where  $\alpha$  and  $\beta$  denote the user-defined weights to the application's QoS and the execution cost, respectively.

Two additional assumptions are made to solve this optimization problem: (1) federate execution time does not increase with the amount of resources (number of cores) assigned, i.e.,  $r_j \leq r'_j$  implies  $t_j(r_j) \geq t_j(r'_j)$ ; (2) federate execution cost does not decrease with the amount of resources assigned, i.e.,  $r_j \leq r'_j$  implies  $c_j(r_j) \leq c_j(r'_j)$ , where  $c_j(r) = r \cdot t_j(r)$ . These are realistic assumptions as many practical applications have *monotonically increasing and sub-linear* speedup functions [17, 18], such as those that follow Amdahl's law

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**Algorithm 2: First Fit Decreasing (FFD)**

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**Input :** Resource assignment  $R = [r_1, r_2, \dots, r_n]$  for all federates in a federation. Current available resource  $A = [a_1, a_2, \dots, a_m]$  of all  $m$  machines in the system.  
**Output:** Machine allocation  $L = [\ell_1, \ell_2, \dots, \ell_n]$  of all federates in the system.

```
1 Sort the resource assignments of all federates in  
  non-increasing order, i.e.,  $r_1 \geq r_2 \geq \dots \geq r_n$ ;  
2 Initialize  $\ell_j \leftarrow 0$ ,  $\forall 1 \leq j \leq n$ ;  
3 for  $j = 1, 2, \dots, n$  do  
4    $fit \leftarrow false$ ;  
5   for  $i = 1, 2, \dots, m$  do  
6     if  $r_j \leq a_i$  then  
7       Update  $a_i \leftarrow a_i - r_j$ ;  
8       Set  $\ell_j \leftarrow i$ ;  
9        $fit \leftarrow true$ ;  
10      break;  
11    end  
12  end  
13  if  $fit = false$  then  
14    // revert allocations done so far  
15    for  $k = 1, 2, \dots, j-1$  do  
16      Find  $i \leftarrow \ell_k$ ;  
17      Update  $a_i \leftarrow a_i + r_k$ ;  
18       $\ell_k \leftarrow 0$ ;  
19    end  
20    break;  
21 end
```

---

[19]. As such, the optimization problem can be solved by examining all possible makespan values while guaranteeing the minimum cost. Algorithm 1 presents the pseudo-code of this solution with a time complexity of  $O(n \log n)$  by maintaining a priority queue for all jobs. Similar approaches can be applied to find the minimum makespan subject to a cost budget or the minimum cost for a target makespan.

As seen in Algorithm 1, we initially assign 1 to  $r_j$  for all the federates (Line 1). In lines 2-4, we find resource assignment  $R^*$  that minimizes  $G$  for the current value of  $r_j$ . In Lines 5-17, we determine the  $R^*$  for all the federates in the federation that minimizes the linear combination of makespan and cost (Line 11).

#### 4.3. Federation Machine Scheduling Heuristics

A custom scheduler is required to deploy all the federates in the federation to their respective distributed computing environments. Since the federates cannot run independently of the federation, the scheduling scheme must simultaneously run all of the federates of



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**Algorithm 3: Best Fit Decreasing (BFD)**

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**Input :** Resource assignment  $R = [r_1, r_2, \dots, r_n]$  for all federates in a federation. Current available resource  $A = [a_1, a_2, \dots, a_m]$  of all  $m$  machines in the system.

**Output:** Machine allocation  $L = [\ell_1, \ell_2, \dots, \ell_n]$  of all federates in the system.

```
1 Sort the resource assignments of all federates in
  non-increasing order, i.e.,  $r_1 \geq r_2 \geq \dots \geq r_n$ ;
2 Initialize  $\ell_j \leftarrow 0, \forall 1 \leq j \leq n$ ;
3 for  $j = 1, 2, \dots, n$  do
4    $best\_i \leftarrow 0$ ;
5    $best\_a \leftarrow \infty$ ;
6   for  $i = 1, 2, \dots, m$  do
7     if  $r_j > a_i$  or  $a_i - r_j \geq best\_a$  then
8       break;
9     else
10       $best\_i \leftarrow i$ ;
11       $best\_a \leftarrow a_i - r_j$ ;
12    end
13  end
14  if  $best\_i \neq 0$  then
15    Update  $a_{best\_i} \leftarrow a_{best\_i} - r_j$ ;
16    Set  $\ell_j \leftarrow best\_i$ ;
17  else
18    // revert allocations done so far
19    for  $k = 1, 2, \dots, j-1$  do
20      Find  $i \leftarrow \ell_k$ ;
21      Update  $a_i \leftarrow a_i + r_k$ ;
22       $\ell_k \leftarrow 0$ ;
23    end
24    break;
25 end
```

---

the federation (Requirement **R4**). This is referred to as *Gang scheduling* or *Bag-of-tasks scheduling* in the literature. To achieve this, EXPPO supports two heuristics to simultaneously schedule the federates on a fixed number  $m$  of available machines while utilizing the resource configuration results obtained from Section 4.2. These approaches handle the case where some of the machines are loaded with other compute tasks unrelated to the federation.

The first heuristic is inspired by the First-Fit Decreasing (FFD) algorithm for bin packing and is described in Algorithm 2. The heuristic first sorts all the federates in decreasing order of resource assignment and then tentatively allocates each one of them in order onto the first available machine (Lines 1-2). If all federates in the federation can be successfully allocated, then the schedule is finalized (Lines 5-12); otherwise, the entire federation

will be temporarily put in a waiting queue to be scheduled later (Lines 13-20). The time complexity of the heuristic is  $O(n(\log n + m))$ . The other heuristic is based on the Best-Fit Decreasing (BFD) algorithm as shown in Algorithm 3 that works similarly to FFD, except that it finds, for each federate, a best-fitting machine (i.e., with the least remaining resource after hosting the federate) (Lines 6-16). Similarly, if all the federates in the federation can be successfully allocated, then the schedule is finalized (Lines 15); otherwise, the entire federation will be temporarily put in a waiting queue to be scheduled later (Lines 17-24). Note that since finding the optimal schedule (or bin packing) is an NP-complete problem, these heuristics may not always find a feasible allocation for a federation even if one exists. However, if an allocation has been found, it is guaranteed to produce the optimal combination of makespan and cost for the federation by using the resource configuration from Section 4.2.

## 5. Co-simulation as a Service Middleware

Figure 6 shows the different components and workflow of the EXPPO co-simulation framework. It is called the Co-simulation-as-a-Service (CaaS) middleware because it enables users to automatically deploy groups of service-based applications to a cloud environment without any concern of resource allocation, application lifecycle monitoring, and cluster management. The functionality of each component is described below.

In ①, the *FrontEnd* component allows a user to submit a simulation job descriptor in JavaScript Object Notation (JSON) using a Representational State Transfer (REST) Application Programming Interface (API). The *FrontEnd* component is implemented using Flask [20] which is a web framework written in Python language. Each simulation job contains a list of federates, and each federate is run on an individual Docker container. The simulation job descriptor also includes meta-information for each federate, for example, resources required, running status, container image details, etc. The *FrontEnd* creates a record in a database ④ for each incoming job, then relays the job identifier to *JobManager* ② which handles resource management. Instead of deploying jobs immediately, the *JobManager* stores received jobs in a local queue and consults the database about the latest status of cluster resources. It then periodically transfers the information of pending jobs and available resources to *JobScheduler* ③, which is a pluggable component that implements

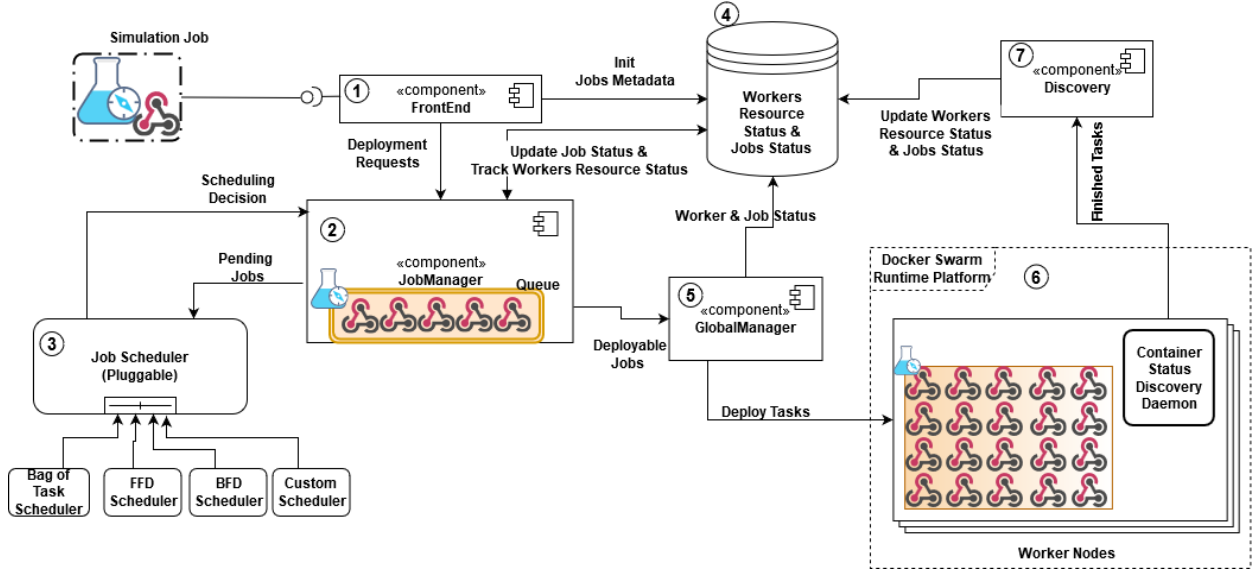


Figure 6: Co-simulation-as-a-Service showing different components and interaction in EXPPO.

multiple scheduling algorithms (such as Algorithm 2 and 3). The JobScheduler either replies with a scheduling decision if the submitted jobs are deployable, or returns a *KeepWaiting* signal to notify the JobManager that resources are insufficient. The JobManager then forwards deployable jobs to *GlobalManager* (5), synchronizes job status with the database, and deletes the deployed jobs from its queue. The GlobalManager is responsible for managing participants of the Docker Swarm Runtime Platform (6). It launches the master node of the Docker Swarm cluster and accepts registration requests sent from worker nodes. Every joined Worker Node runs a CaaS-Worker daemon that is used to receive and perform commands sent from the GlobalManager. The GlobalManager parses received deployment requests and spawns containers in specific Worker Nodes. Additionally, the Worker Nodes employ a CaaS-Discovery daemon to track the status of containers, and reports a *StatusChanged* signal to the Discovery component (7) when a task is completed.

## 6. Experimental Evaluation

### 6.1. Experimental Setup

EXPPO was validated using seven homogeneous compute servers with a configuration of 12-core 2.1 GHz AMD Opteron central processing units, 32 GB memory, 500 GB disk space, and the Ubuntu 16.04 operating system. The runtime platform was based on

Docker engine version 19.0.5 with swarm mode enabled. There was one client machine that submitted simulation job requests. The front end, job manager, job scheduler and the global manager components were running on a single shared compute server, and five compute worker servers were deployed for running the simulation jobs.

The simulation job consisted of three federates each running a unique application from the PARSEC benchmark: *frequmine*, *blackscholes* and *ferret*. These applications were chosen as they are realistic representations of real-world simulation tasks [13]. During the federation execution, each federate executed its application at every logical time step, for a total number of 100 logical time steps. The implementation of the co-simulation federation was done using Portico HLA [21]. The BFD scheduler was used as the default scheduler in the experiments. The default weights for the QoS and the cost were set to  $\alpha = 1$  and  $\beta = 0.5$ . These weights are chosen so that they correspond to a specific brand of QoS that prioritizes low makespan over resource usage, which is representative of CPS applications.

### 6.2. Experimental Results

EXPPO recommended a resource configuration of 4 cores for *frequmine* federate, 4 cores for *ferret* federate and 1 core for *blackscholes* federate. Two baseline approaches were used to compare the performance of resource configuration selection of EXPPO. In the first approach (*least configuration*), all the federates were assigned the lowest possible configuration of 1 core each.

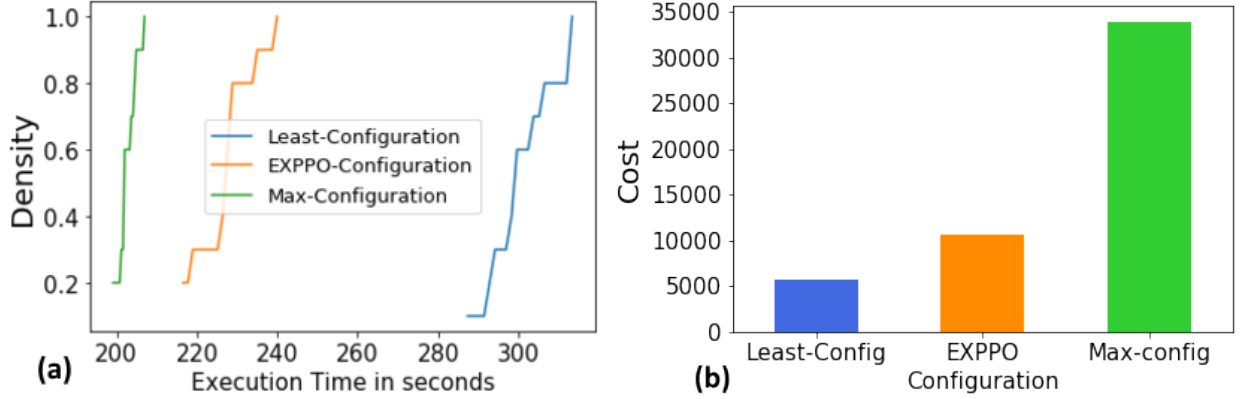


Figure 7: (a) Execution time for the EXPPO resource configuration compared to other strategies. (b) Cost for the EXPPO resource configuration compared to other strategies.

In the second approach (*max configuration*), all the federates were assigned the highest possible configuration of 10 cores each. The performance data was collected over 10 simulation jobs.

Figure 7(a) shows the cumulative distribution function (CDF) of the execution time of EXPPO compared to the other two approaches. The resource configurations selected by EXPPO for the federation had a 90th percentile execution time of around 230 seconds, which was significantly better than the 320 seconds for the *least configuration*. Its performance was close to that of the *max configuration* (90th percentile execution time of around 200 seconds), with the difference due to its lower resource allocation to conserve the cost.

Figure 7(b) shows the cost analysis of the three strategies using the cost function defined in Section 4-B. As can be seen, resource configuration selected by EXPPO incurred a larger cost than the *least configuration* due to the higher resource allocation to reduce execution time, but it had a substantially lower cost compared to the *max configuration*.

Next, we show the comparison of the best-fit decreasing (BFD) and the first-fit decreasing (FFD) scheduling strategies on the simulation jobs. We can see from Figure 8 that the 90th percentile execution time of the jobs for BFD is better compared to the FFD. This shows that BFD has better performance in terms of 90th percentile execution time, and is therefore used as the default scheduler in EXPPO.

Overall, the results show that EXPPO is able to select resource configurations and schedule the federates in such a way that minimizes the combination of execution time and cost of the simulation successfully.

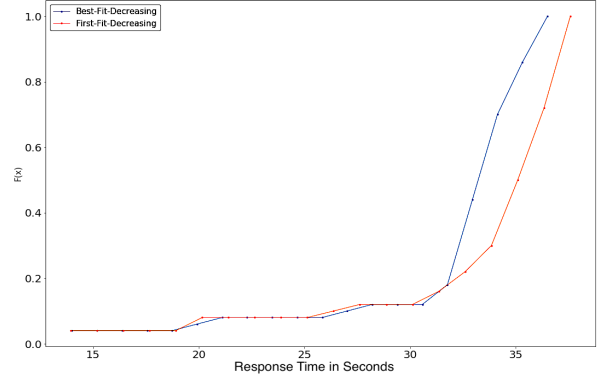


Figure 8: Execution time comparison when running the jobs using the Best-Fit Decreasing (BFD) and First-Fit Decreasing (FFD) scheduling strategy. We can see that the 90th percentile execution time for all the jobs is better when using BFD compared to FFD in EXPPO.

### 6.3. Validating the Resource Configuration Tuner

We also study the effectiveness of the resource configuration tuner for different values of  $\alpha$  and  $\beta$ , and show that EXPPO provides resource configurations for the simulation jobs with the lowest  $G$  value as defined in Algorithm 1 compared to the baseline strategies. The  $G$  value is the combined objective involving both makespan and cost of the simulation jobs. We set different values for the  $(\alpha, \beta)$  pairs with  $\alpha$  varying between  $[10, 100]$  and  $\beta$  varying between  $[0.1, 1.0]$ . We show that no matter what values, our algorithm is always producing a better  $G$ . We tested our scheme for values of  $(\alpha, \beta)$  pairs as shown in Table 3.

Figure 9 shows the performance of the EXPPO compared to the *least configuration* and the *max configuration* baseline strategies in terms of combined objective  $G$ , the cost and the makespan. We can see from Figure

Set	$\alpha$	$\beta$
1	10	0.35
2	10	0.60
3	10	0.85
4	35	0.10
5	35	0.35
6	35	0.60
7	35	0.85
8	60	0.10
9	60	0.35
10	60	0.60
11	60	0.85
12	85	0.10
13	85	0.35
14	85	0.60
15	85	0.85

Table 3: Pairs of  $(\alpha, \beta)$  used for the resource configuration tuner validation.

9(a) that EXPPO always find configurations that have the best values of  $G$  compared to *least configuration* and *max configuration*. This is because the other two baselines only focus on one objective, i.e., either best performance (*max configuration*) or lowest cost (*least configuration*). However, EXPPO tries to minimize the combination of cost and makespan for running the simulation jobs. Similarly, we can see in Figure 9(b) that EXPPO is able to find resource configurations that have lower cost compared to *max configuration*, while from Figure 9(c), we can see that EXPPO is able to find resource configurations that have substantially lower makespan compared to *least configuration*.

## 7. Related Work

In [8], the authors presented a Kubernetes co-simulation execution platform for cloud computing environments. Similarly, [9] presented a Docker swarm co-simulation platform for running mixed electrical energy systems simulations. However, these platforms do not use a gang-scheduling based simulation deployment strategy.

For resource recommendation, [22] presented a data-driven approach for selecting the best resource configuration for a virtual machine from a set of different configuration options. [23] explored the cost-sensitive allocation of independent tasks to cloud computing environments. However, these approaches are different from EXPPO as they focus on a single task rather than a pool of BSP tasks.

In [24, 25, 26] scheduling of parallel and distributed real-time tasks is presented. The computation model supported in these tasks is fork-join computation model [27]. However, these methods do not consider the bulk-synchronous parallel computation model which is exhibited by the co-simulation in this paper.

In [28], the authors proposed a scientific workflows scheduling algorithm to minimize the execution time under budget constraints for deploying to cloud computing environments. [29] proposed an advance-reservation scheduling strategy for message passing interface (MPI) applications. Similarly, [30] presented an approach for gang-scheduling of jobs with different resource needs, such as a compute intensive task paired with a network or I/O intensive task. In [31], the authors present scheduling of multiple container workloads on shared cluster as a minimum cost flow problem (MCFP) constraint satisfaction problem. In [32], the authors proposed a locality-based process placements for parallel and distributed simulation. The evaluation of the proposed framework was carried out using the OMNET++ network simulator.

[33] proposed a hierarchical virtual machine (VM) based workload aware resource allocation for the federates of a simulation in cloud data center. However, the workload characteristics of the federates were restricted to single threaded applications. In [34], priority based scheduling of parallel job on high priority and low priority VMs to allow avoid under utilization of cloud resources and improve parallel job performance. Thus only two categories of resource assignment is possible in this scheme. In [35] authors presented different vector bin packing algorithms and first fit decreasing (FFD) heuristic to allocate resources in a shared cluster to the static workloads. The evaluation of which was shown through simulation. Similarly, [36] presented various FFD based algorithms for VM placement problems.

Compared to related work, EXPPO provides a resource recommendation engine which tries to minimize the makespan and cost for the entire federation (BSP tasks) rather than a single task. EXPPO uses a gang scheduling scheme based on heuristic bin packing techniques to deploy the entire federation on Docker container platform as one batch job. Furthermore, EXPPO provides automatic code generation of distributed tracing probes for performance profiling of the federates which maybe deployed on a distributed infrastructure, thereby relieving the developers from incurring complexities in writing code for the profiling of federates [37].

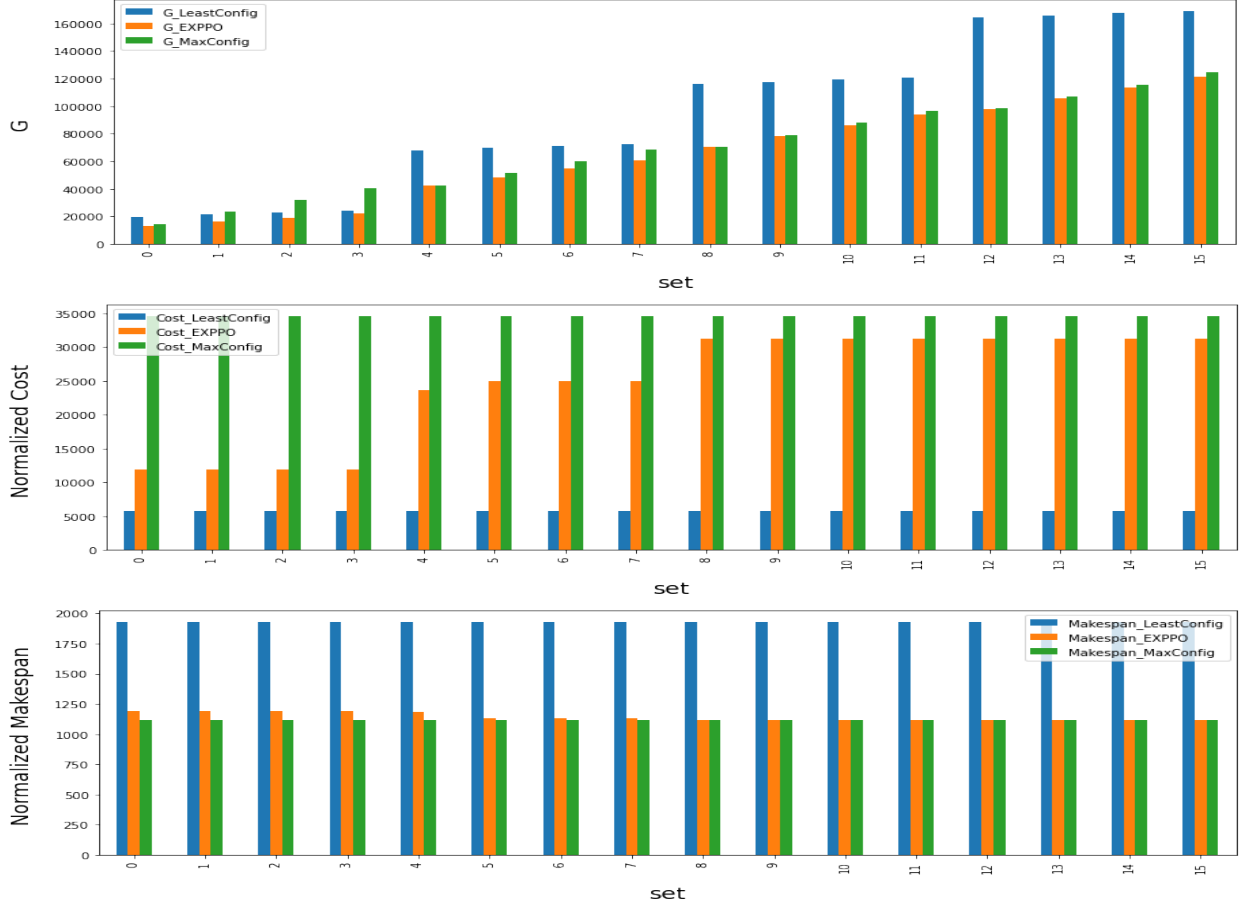


Figure 9: We illustrate the performance of resource configuration tuner for different values of  $(\alpha, \beta)$ . (a) shows that EXPPO is able to find resource configurations that have the lowest  $G$  value, which is the combination of cost and makespan of the simulation jobs. (b) shows that EXPPO is able to find consistently better resource configurations in terms of cost compared to *max configuration*. (c) shows that EXPPO is able to find resource configurations that have substantially lower makespan compared to *least configuration*.

## 8. Discussion

With the advancement of technologies such as 5G and edge computing, CPS are getting increasingly deployed in various safety critical applications. Co-simulations of such dispersed and diverse deployed CPS offers researchers a thorough understanding of the complex dynamics of the systems. However, much of the co-simulation tools used by the researchers in the CPS domains focus largely on conducting simulations on a single compute node and seldom use distributed computational capabilities to run such simulations. There is a large barrier to entry to make these tools leverage distributed cloud computing resources. For example, network emulation tools like mininet or ns-3 do not natively support execution on multiple hosts and are restricted to running in a single host machine. Similarly,

there is a high barrier to entry to the underlying co-simulation HLA middleware to adapt these technologies to run in the cloud computing environments. In our experiments, we used the Portico HLA middleware, where we faced several issues running this middleware directly on the Docker containers. One of the main issues was the networking between the federates hosted across the physical hosts. The middleware leverages the multicast approach for communication between the federates. However, the default networking support in the Docker swarm did not support multicast. Thus, considerable operational issues were faced to find out the right set of software drivers to support multicast networking for the Docker swarm cluster. We finally settled with the *weave* network plugin to allow us the support for federate communication across the physical host systems

after facing many accidental complexities involved in provisioning such systems. Also, each of these federations demanded running on a separate subnet address to allow for isolation of these multicast traffic within these subnets' address space. Weave networking allowed us to create these virtual subnets and alleviate the multicast traffic isolation issue.

Next, when running the simulations in the cloud computing environments, sometimes the simulator provides native visualization interface that allows for viewing the progress for the simulation experiments. However, since the simulators are running in the docker containers, some simulators do not expose a web-server where it is possible to see the simulation progress. These simulators are not necessarily built to run in the cloud native environments. As such we had to find alternate solutions to make the simulation visualization be exposed to the remote users. Thus, there was a need to find an external software driver to support the streaming of the visualization interface of the simulator running in the cloud computing environments. After testing various different remote visualization drivers, the virtual network connection (VNC) viewer, allowed us to fulfill this capability of streaming real time visualization of the simulator running on the cloud computing infrastructure to the end-users.

Running the co-simulations in the docker container context also requires creating special packaging of the inputs files needed to run inside the docker container. Also, there was a need to capture the log files generated from the simulation itself. Since, the federates of the simulation could be running across the physical host, there was a need to aggregate the logs running from all these federates at a centralized location. Thus, we were able to explore the network file sharing facility (NFS) to mount the distributed file systems to the federate docker container and associate it with a distinct tag. This tag represents the federation and the experimental run to uniquely identify and aggregate the simulation logs and results at a centralized place.

In summary, there are many operational challenges involved in successfully running these simulations in the cloud computing context, which we were able to address in the EXPPO platform thereby having faster and seamless execution of the CPS co-simulations in the cloud computing environment.

## 9. Conclusion & Future Work

Resource allocation plays a critical role in co-simulation performance. However, the end user is not necessarily well-equipped to determine what resource

allocations work best for their co-simulation jobs given the various resource configuration options (and associated costs) available from the cloud provider. To address these challenges, this paper presents EXPPO, which is a Co-simulation-as-a-Service (CaaS) platform for executing distributed co-simulations in cloud computing environments. EXPPO provides performance profiling capabilities for federates which help in the understanding of the relationship between resource allocations and the simulation performance. Similarly, it addresses performance profiling challenges for a simulation job comprising heterogeneous computation tasks or federates deployed across distributed systems. Furthermore, EXPPO selects the resource configurations for these federates in a way that not only minimizes the makespan of the co-simulation, but also satisfies the cost budget of the user.

Future work will address the following:

- The assumption that federate computations have identical wall-clock execution times across all iterations restricts the generality of the approach to a subset of application use cases. For hybrid simulations or simulations with non-linear dynamics, the execution times of the involved simulation steps will vary at each iteration. Future work must explore dynamic resource allocation for federates which have different work loads for different computation steps. Reinforcement learning approaches which can monitor the federates and dynamically adjust resource allocations based on the varying demand of a federate over time offer a promising approach that remains to be explored.
- EXPPO only considers workloads that are CPU bound. It assumes constant communication costs during each computation step for all the federates. Future work must consider the different communication costs for simulations which may need to be constantly updating states or sending messages for triggering discrete events.
- When other applications are co-located in the cluster environments, the effects of noisy-neighbors [38, 39] can also affect federation performance. Thus, future work could consider the effects of noisy-neighbors for resource scheduling and simulation placement in the cluster.
- EXPPO assumes the use of homogeneous servers for running simulations to simplify its algorithms, and could be extended to include heterogeneous systems. Also, with the growing relevance of edge



computing and digital twin techniques, efficient resource allocation and scheduling of the simulations at the edge will be necessary.

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