



COMP SUPERSCALAR

COMPSS at BSC

Supercomputers Manual

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**Barcelona
Supercomputing
Center**

Centro Nacional de Supercomputación

This manual only provides information about the COMPSs usage at MareNostrum. Specifically, it details the available COMPSs modules, how to load them and how to create and track COMPSs jobs.

If you want to install COMPSs on your local machine please refer to the *COMPSs Installation Manual* available at our webpage <http://compss.bsc.es>.

For further information about the application's execution please refer to the *COMPSs User Manual: Application execution guide* available at <http://compss.bsc.es> .

For further information about the application's development please refer to the *COMPSs User Manual: Application development guide* available at <http://compss.bsc.es/> .

For full COMPSs example application (codes, execution commands, results, logs, etc.) please refer to the *COMPSs Sample Applications* available at <http://compss.bsc.es/>

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1 COMP Superscalar (COMPSs)

COMP Superscalar (COMPSs) is a programming model which aims to ease the development of applications for distributed infrastructures, such as Clusters, Grids and Clouds. COMP Superscalar also features a runtime system that exploits the inherent parallelism of applications at execution time.

For the sake of programming productivity, the COMPSs model has four key characteristics:

- **Sequential programming:** COMPSs programmers do not need to deal with the typical duties of parallelization and distribution, such as thread creation and synchronization, data distribution, messaging or fault tolerance. Instead, the model is based on sequential programming, which makes it appealing to users that either lack parallel programming expertise or are looking for better programmability.
- **Infrastructure unaware:** COMPSs offers a model that abstracts the application from the underlying distributed infrastructure. Hence, COMPSs programs do not include any detail that could tie them to a particular platform, like deployment or resource management. This makes applications portable between infrastructures with diverse characteristics.
- **Standard programming languages:** COMPSs is based on the popular programming language Java, but also offers language bindings for Python and C/C++ applications. This facilitates the learning of the model, since programmers can reuse most of their previous knowledge.
- **No APIs:** In the case of COMPSs applications in Java, the model does not require to use any special API call, pragma or construct in the application; everything is pure standard Java syntax and libraries. With regard the Python and C/C++ bindings, a small set of API calls should be used on the COMPSs applications.

2 Common usage

2.1 Available COMPSs modules

COMPSs is configured as a Linux Module. As shown in next Figure, the users can type the `module available COMPSs` command to list the supported COMPSs modules in the supercomputer. The users can also execute the `module load COMPSs/<version>` command to load an specific COMPSs module.

```
$ module available COMPSs
----- /apps/modules/modulefiles/tools -----
COMPSs/1.3
COMPSs/1.4
COMPSs/2.0
COMPSs/2.1
COMPSs/2.2

COMPSs/release(default)
COMPSs/trunk

$ module load COMPSs/release
load java/1.8.0u66 (PATH, MANPATH, JAVA_HOME, JAVA_ROOT, JAVA_BINDIR,
                  SDK_HOME, JDK_HOME, JRE_HOME)
load MKL/11.0.1 (LD_LIBRARY_PATH)
load PYTHON/2.7.3 (PATH, MANPATH, LD_LIBRARY_PATH, C_INCLUDE_PATH)
load COMPSs/release (PATH, MANPATH, COMPSS_HOME)
```

The following command can be run to check if the correct COMPSs version has been loaded:

```
$ enqueue_comps --version
COMPSs version <version>
```

2.2 Configuration

The COMPSs module contains **all** the COMPSs dependencies, including Java, Python and MKL. Modifying any of these dependencies can cause execution failures and thus, we **do not** recomend to change them. Before running any COMPSs job please check your environment and, if needed, comment out any line inside the `.bashrc` file that loads custom COMPSs, Java, Python and/or MKL modules.

The COMPSs module needs to be loaded in all the nodes that will run COMPSs jobs. Consequently, the `module load` **must** be included in your `.bashrc` file. To do so, please run the following command with the corresponding COMPSs version:

```
$ cat "module load COMPSs/release" >> ~/.bashrc
```

Log out and back in again to check that the file has been correctly edited. The next listing shows an example of the output generated by well loaded COMPSs installation.

```

$ exit
$ ssh USER@SC
load java/1.8.0u66 (PATH, MANPATH, JAVA_HOME, JAVA_ROOT, JAVA_BINDIR,
                  SDK_HOME, JDK_HOME, JRE_HOME)
load MKL/11.0.1 (LD_LIBRARY_PATH)
load PYTHON/2.7.3 (PATH, MANPATH, LD_LIBRARY_PATH, C_INCLUDE_PATH)
load COMPSs/release (PATH, MANPATH, COMPSS_HOME)

$ enqueue_compss --version
COMPSs version <version>

```

Please remember that COMPSs runs in several nodes and your current environment is not exported to them. Thus, all the needed environment variables **must** be loaded through the *.bashrc* file.

Please remember that PyCOMPSs uses Python 2.7 by default. In order to use Python 3, the Python 2.7 module **must** be unloaded after loading COMPSs module, and then load the Python 3 module.

2.3 COMPSs Job submission

COMPSs jobs can be easily submitted by running the **enqueue_compss** command. This command allows to configure any **runcompss** option and some particular queue options such as the queue system, the number of nodes, the wallclock time, the master working directory, the workers working directory and number of tasks per node.

Next, we provide detailed information about the *enqueue_compss* command:

```

$ enqueue_compss -h

Usage: enqueue_compss [queue_system_options] [COMPSs_options]
       application_name [application_arguments]

* Options:

General:

  --help, -h                Print this help message

Queue system configuration:

  --sc_cfg=<name>            SuperComputer configuration file to use.
                             Must exist inside queues/cfgs/
                             Default: default

Submission configuration:

  --exec_time=<minutes>      Expected execution time of the application (in minutes)
                             Default: 10

```

<code>--num_nodes=<int></code>	Number of nodes to use Default: 2
<code>--num_switches=<int></code>	Maximum number of different switches. Select 0 for no restrictions. Maximum nodes per switch: 18 Only available for at least 4 nodes. Default: 0
<code>--queue=<name></code>	Queue name to submit the job. Depends on the queue system. For example (Nord3): bsc_cs bsc_debug debug interactive Default: default
<code>--reservation=<name></code>	Reservation to use when submitting the job. Default: disabled
<code>--constraints=<constraints></code>	Constraints to pass to queue system. Default: disabled
<code>--qos=<qos></code>	Quality of Service to pass to the queue system. Default: default
<code>--job_dependency=<jobID></code>	Postpone job execution until the job dependency has ended. Default: None
<code>--storage_home=<string></code>	Root installation dir of the storage implementation Default: null
<code>--storage_props=<string></code>	Absolute path of the storage properties file Mandatory if storage_home is defined

Launch configuration:

<code>--cpus_per_node=<int></code>	Available CPU computing units on each node Default: 48
<code>--gpus_per_node=<int></code>	Available GPU computing units on each node Default: 0
<code>--max_tasks_per_node=<int></code>	Maximum number of simultaneous tasks running on a node Default: -1
<code>--node_memory=<MB></code>	Maximum node memory: disabled <int> (MB) Default: disabled
<code>--network=<name></code>	Communication network for transfers: default ethernet infiniband data. Default: infiniband
<code>--prolog="<string>"</code>	Task to execute before launching COMPSs (Notice the quotes) If the task has arguments split them by " , " rather than spaces. This argument can appear multiple times for more than one prolog action Default: Empty
<code>--epilog="<string>"</code>	Task to execute after executing the COMPSs application (Notice the quotes) If the task has arguments split them by " , " rather than spaces. This argument can appear multiple times for more than one epilog action Default: Empty
<code>--master_working_dir=<path></code>	Working directory of the application Default: .
<code>--worker_working_dir=<name path></code>	Worker directory. Use: scratch gpfs <path> Default: scratch
<code>--worker_in_master_cpus=<int></code>	Maximum number of CPU computing units that the master node can run as worker. Cannot exceed cpus_per_node.

	Default: 24
<code>--worker_in_master_memory=<int> MB</code>	Maximum memory in master node assigned to the worker. Cannot exceed the <code>node_memory</code> . Mandatory if <code>worker_in_master_cpus</code> is specified. Default: 50000
<code>--jvm_worker_in_master_opts="<string>"</code>	Extra options for the JVM of the COMPSs Worker in the Master Node. Each option separated by "," and without blank spaces (Notice the quotes) Default:
<code>--container_image=<path></code>	Runs the application by means of a container engine image Default: Empty
<code>--container_comps_path=<path></code>	Path where compss is installed in the container image Default: /opt/COMPSs
<code>--container_opts="<string>"</code>	Options to pass to the container engine Default: empty
<code>--elasticity=<max_extra_nodes></code>	Activate elasticity specifying the maximum extra nodes (ONLY AVAILABLE FORM SLURM CLUSTERS WITH NIO ADAPTOR) Default: 0
Runcompss configuration:	
Tools enablers:	
<code>--graph=<bool>, --graph, -g</code>	Generation of the complete graph (<code>true/false</code>) When no value is provided it is set to <code>true</code> Default: <code>false</code>
<code>--tracing=<level>, --tracing, -t</code>	Set generation of traces and/or tracing level ([<code>true</code> basic] advanced <code>false</code>) True and basic levels will produce the same traces. When no value is provided it is set to <code>true</code> Default: <code>false</code>
<code>--monitoring=<int>, --monitoring, -m</code>	Period between monitoring samples (milliseconds) When no value is provided it is set to 2000 Default: 0
<code>--external_debugger=<int>, --external_debugger</code>	Enables external debugger connection on the specified port (or 9999 if empty) Default: <code>false</code>
Runtime configuration options:	
<code>--task_execution=<compss storage></code>	Task execution under COMPSs or Storage. Default: <code>compss</code>
<code>--storage_conf=<path></code>	Path to the storage configuration file Default: None
<code>--project=<path></code>	Path to the project XML file Default: /apps/COMPSs/2.3/Runtime/configuration/xml/projects/default_project.xml
<code>--resources=<path></code>	Path to the resources XML file Default: /apps/COMPSs/2.3/Runtime/configuration/xml/resources/default_resources.xml
<code>--lang=<name></code>	Language of the application (java/c/python) Default: Inferred is possible. Otherwise: java
<code>--summary</code>	Displays a task execution summary at the end of the application execution

	Default: <code>false</code>
<code>--log_level=<level>, --debug, -d</code>	Set the debug level: <code>off info debug</code> Default: <code>off</code>
Advanced options:	
<code>--extrae_config_file=<path></code>	Sets a custom extrae config file. Must be in a shared disk between all COMPSs workers. Default: <code>null</code>
<code>--comm=<ClassName></code>	Class that implements the adaptor <code>for</code> communications Supported adaptors: <code>es.bsc.compss.nio.master.NIOAdaptor</code> <code>es.bsc.compss.gat.master.GATAdaptor</code> Default: <code>es.bsc.compss.nio.master.NIOAdaptor</code>
<code>--conn=<className></code>	Class that implements the runtime connector <code>for</code> the cloud Supported connectors: <code>es.bsc.compss.connectors.DefaultSSHConnector</code> <code>es.bsc.compss.connectors.DefaultNoSSHConnector</code> Default: <code>es.bsc.compss.connectors.DefaultSSHConnector</code>
<code>--scheduler=<className></code>	Class that implements the Scheduler <code>for</code> COMPSs Supported schedulers: <code>es.bsc.compss.scheduler.fullGraphScheduler.FullGraphScheduler</code> <code>es.bsc.compss.scheduler.fifoScheduler.FIFOScheduler</code> <code>es.bsc.compss.scheduler.resourceEmptyScheduler.ResourceEmptyScheduler</code> Default: <code>es.bsc.compss.scheduler.loadBalancingScheduler.LoadBalancingScheduler</code>
<code>--scheduler_config_file=<path></code>	Path to the file which contains the scheduler configuration. Default: <code>Empty</code>
<code>--library_path=<path></code>	Non-standard directories to search <code>for</code> libraries (e.g. Java JVM library, Python library, C binding library) Default: <code>Working Directory</code>
<code>--classpath=<path></code>	Path <code>for</code> the application classes / modules Default: <code>Working Directory</code>
<code>--appdir=<path></code>	Path <code>for</code> the application class folder. Default: <code>/home/user/</code>
<code>--pythonpath=<path></code>	Additional folders or paths to add to the PYTHONPATH Default: <code>/home/user/</code>
<code>--base_log_dir=<path></code>	Base directory to store COMPSs log files (a <code>.COMPSs/</code> folder will be created inside this location) Default: <code>User home</code>
<code>--specific_log_dir=<path></code>	Use a specific directory to store COMPSs log files (the folder MUST exist and no sandbox is created) Warning: Overwrites <code>--base_log_dir</code> option Default: <code>Disabled</code>
<code>--uuid=<int></code>	Preset an application UUID Default: <code>Automatic random generation</code>
<code>--master_name=<string></code>	Hostname of the node to run the COMPSs master Default:
<code>--master_port=<int></code>	Port to run the COMPSs master communications. Only <code>for</code> NIO adaptor Default: <code>[43000,44000]</code>
<code>--jvm_master_opts="<string>"</code>	Extra options <code>for</code> the COMPSs Master JVM. Each option separated by <code>","</code> and without blank spaces (Notice the quotes) Default:

<code>--jvm_workers_opts=<string></code>	Extra options for the COMPSs Workers JVMs. Each option separated by "," and without blank spaces (Notice the quotes) Default: <code>-Xms1024m,-Xmx1024m,-Xmn400m</code>
<code>--cpu_affinity=<string></code>	Sets the CPU affinity for the workers Supported options: disabled, automatic, user defined map of the form <code>"0-8/9,10,11/12-14,15,16"</code> Default: automatic
<code>--gpu_affinity=<string></code>	Sets the GPU affinity for the workers Supported options: disabled, automatic, user defined map of the form <code>"0-8/9,10,11/12-14,15,16"</code> Default: automatic
<code>--task_count=<int></code>	Only for C/Python Bindings. Maximum number of different functions/methods, invoked from the application, that have been selected as tasks Default: 50
<code>--input_profile=<path></code>	Path to the file which stores the input application profile Default: Empty
<code>--output_profile=<path></code>	Path to the file to store the application profile at the end of the execution Default: Empty
<code>--PyObject_serialize=<bool></code>	Only for Python Binding. Enable the object serialization to string when possible (<code>true/false</code>). Default: <code>false</code>
<code>--persistent_worker_c=<bool></code>	Only for C Binding. Enable the persistent worker in c (<code>true/false</code>). Default: <code>false</code>
<code>--enable_external_adaptation=<bool></code>	Enable external adaptation. This option will disable the Resource Optimizer. Default: <code>false</code>
* Application name:	
For Java applications: Fully qualified name of the application	
For C applications: Path to the master binary	
For Python applications: Path to the .py file containing the main program	
* Application arguments:	
Command line arguments to pass to the application. Can be empty.	

3 MareNostrum 4

3.1 Basic queue commands

The MareNostrum supercomputer uses the SLURM (Simple Linux Utility for Resource Management) workload manager. The basic commands to manage jobs are listed below:

- **sbatch** Submit a batch job to the SLURM system
- **scancel** Kill a running job
- **squeue -u <username>** See the status of jobs in the SLURM queue

For more extended information please check the *SLURM: Quick start user guide* at <https://slurm.schedmd.com/quickstart.html>.

3.2 Tracking COMPSs jobs

When submitting a COMPSs job a temporal file will be created storing the job information. For example:

```
$ enqueue_comps \
  --exec_time=15 \
  --num_nodes=3 \
  --cpus_per_node=16 \
  --master_working_dir=. \
  --worker_working_dir=gpfs \
  --lang=python \
  --log_level=debug \
  <APP> <APP_PARAMETERS>

SC Configuration:      default.cfg
Queue:                 default
Reservation:           disabled
Num Nodes:             3
Num Switches:          0
GPUs per node:         0
Job dependency:        None
Exec-Time:             00:15
Storage Home:          null
Storage Properties:    null
Other:
  --sc_cfg=default.cfg
  --cpus_per_node=48
  --master_working_dir=.
  --worker_working_dir=gpfs
  --lang=python
  --classpath=.
  --library_path=.
  --comm=es.bsc.compss.nio.master.NIOAdaptor
  --tracing=false
  --graph=false
  --pythonpath=.
  <APP> <APP_PARAMETERS>
Temp submit script is: /scratch/tmp/tmp.pBG5yfFxEO

$ cat /scratch/tmp/tmp.pBG5yfFxEO
#!/bin/bash
#
#SBATCH --job-name=COMPSs
```

```
#SBATCH --workdir=.
#SBATCH -o compss-%J.out
#SBATCH -e compss-%J.err
#SBATCH -N 3
#SBATCH -n 144
#SBATCH --exclusive
#SBATCH -t00:15:00
...
```

In order to trac the jobs state users can run the following command:

```
$ squeue
JOBID  PARTITION  NAME      USER  TIME_LEFT  TIME_LIMIT  START_TIME  ST  NODES  CPUS  NODELIST
474130  main       COMPSs    XX    0:15:00    0:15:00     N/A        PD   3     144    -
```

The specific COMPSs logs are stored under the `~/.COMPSs/` folder; saved as a local *runcompss* execution. For further details please check *COMPSs User Manual: Application Execution* available at our webpage <http://compss.bsc.es> .

4 MinoTauro

4.1 Basic queue commands

The MinoTauro supercomputer uses the SLURM (Simple Linux Utility for Resource Management) workload manager. The basic commands to manage jobs are listed below:

- **sbatch** Submit a batch job to the SLURM system
- **scancel** Kill a running job
- **squeue -u <username>** See the status of jobs in the SLURM queue

For more extended information please check the *SLURM: Quick start user guide* at <https://slurm.schedmd.com/quickstart.html>.

4.2 Tracking COMPSs jobs

When submitting a COMPSs job a temporal file will be created storing the job information. For example:

```
$ enqueue_comps \
  --exec_time=15 \
  --num_nodes=3 \
  --cpus_per_node=16 \
  --master_working_dir=. \
  --worker_working_dir=gpfs \
  --lang=python \
  --log_level=debug \
  <APP> <APP_PARAMETERS>

SC Configuration:      default.cfg
Queue:                 default
Reservation:           disabled
Num Nodes:             3
Num Switches:          0
GPUs per node:         0
Job dependency:        None
Exec-Time:             00:15
Storage Home:          null
Storage Properties:    null
Other:
  --sc_cfg=default.cfg
  --cpus_per_node=16
  --master_working_dir=.
  --worker_working_dir=gpfs
  --lang=python
  --classpath=.
  --library_path=.
  --comm=es.bsc.compss.nio.master.NIOAdaptor
  --tracing=false
  --graph=false
  --pythonpath=.
  <APP> <APP_PARAMETERS>
Temp submit script is: /scratch/tmp/tmp.pBG5yfFxEO

$ cat /scratch/tmp/tmp.pBG5yfFxEO
#!/bin/bash
#
#SBATCH --job-name=COMPSs
```

```
#SBATCH --workdir=.
#SBATCH -o compss-%J.out
#SBATCH -e compss-%J.err
#SBATCH -N 3
#SBATCH -n 48
#SBATCH --exclusive
#SBATCH -t00:15:00
...
```

In order to trac the jobs state users can run the following command:

```
$ squeue
JOBID PARTITION NAME USER ST TIME NODES NODELIST (REASON)
XXXX projects COMPSs XX R 00:02 3 nvb[6-8]
```

The specific COMPSs logs are stored under the `~/.COMPSs/` folder; saved as a local *runcompss* execution. For further details please check *COMPSs User Manual: Application Execution* available at our webpage <http://compss.bsc.es> .

5 Nord 3

5.1 Basic queue commands

The Nord3 supercomputer uses the LSF (Load Sharing Facility) workload manager. The basic commands to manage jobs are listed below:

- **bsub** Submit a batch job to the LSF system
- **bkill** Kill a running job
- **bjobs** See the status of jobs in the LSF queue
- **bqueues** Information about LSF batch queues

For more extended information please check the *IBM Platform LSF Command Reference* at https://www.ibm.com/support/knowledgecenter/en/SSETD4_9.1.2/lsf_kc_cmd_ref.html.

5.2 Tracking COMPSs jobs

When submitting a COMPSs job a temporal file will be created storing the job information. For example:

```
$ enqueue_compss \
--exec_time=15 \
--num_nodes=3 \
--cpus_per_node=16 \
--master_working_dir=. \
--worker_working_dir=gdfs \
--lang=python \
--log_level=debug \
<APP> <APP_PARAMETERS>

SC Configuration:      default.cfg
Queue:                 default
Reservation:           disabled
Num Nodes:             3
Num Switches:          0
GPUs per node:         0
Job dependency:        None
Exec-Time:             00:15
Storage Home:          null
Storage Properties:    null
Other:
  --sc_cfg=default.cfg
  --cpus_per_node=16
  --master_working_dir=.
  --worker_working_dir=gdfs
  --lang=python
  --classpath=.
  --library_path=.
  --comm=es.bsc.compss.nio.master.NIOAdaptor
  --tracing=false
  --graph=false
  --pythonpath=.
  <APP> <APP_PARAMETERS>
Temp submit script is: /scratch/tmp/tmp.pBG5yfFxEO
```



```

$ cat /scratch/tmp/tmp.pBG5yfFxEO
#!/bin/bash
#
#BSUB -J COMPSs
#BSUB -cwd .
#BSUB -oo compss-%J.out
#BSUB -eo compss-%J.err
#BSUB -n 3
#BSUB -R "span[ptile=1]"
#BSUB -W 00:15
...

```

In order to track the jobs state users can run the following command:

```

$ bjobs
JOBID  USER   STAT  QUEUE  FROM_HOST  EXEC_HOST  JOB_NAME  SUBMIT_TIME
XXXX   bscXX  PEND  XX     login1     XX         COMPSs    Month Day Hour

```

The specific COMPSs logs are stored under the `~/.COMPSs/` folder; saved as a local *runcompss* execution. For further details please check *COMPSs User Manual: Application Execution* available at our webpage <http://compss.bsc.es>.

6 Enabling COMPSs Monitor

6.1 Configuration

As supercomputer nodes are connection restricted, the better way to enable the *COMPSs Monitor* is from the users local machine. To do so please install the following packages:

- COMPSs Runtime
- COMPSs Monitor
- sshfs

For further details about the COMPSs packages installation and configuration please refer to the *COMPSs Installation Manual* available at our webpage <http://compss.bsc.es> . If you are not willing to install COMPSs in your local machine please consider to download our Virtual Machine available at our webpage.

Once the packages have been installed and configured, users need to mount the sshfs directory as follows. The `SC_USER` stands for your supercomputer's user, the `SC_ENDPOINT` to the supercomputer's public endpoint and the `TARGET_LOCAL_FOLDER` to the local folder where you wish to deploy the supercomputer files):

```
compss@bsc:~$ scp $HOME/.ssh/id_dsa.pub ${SC_USER}@mn1.bsc.es:~/id_dsa_local.pub
compss@bsc:~$ ssh SC_USER@SC_ENDPOINT
"cat ~/id_dsa_local.pub >> ~/.ssh/authorized_keys;
rm ~/id_dsa_local.pub"
compss@bsc:~$ mkdir -p TARGET_LOCAL_FOLDER/.COMPSs
compss@bsc:~$ sshfs -o IdentityFile=$HOME/.ssh/id_dsa -o allow_other
SC_USER@SC_ENDPOINT:~/.COMPSs
TARGET_LOCAL_FOLDER/.COMPSs
```

Whenever you wish to unmount the sshfs directory please run:

```
compss@bsc:~$ sudo umount TARGET_LOCAL_FOLDER/.COMPSs
```

6.2 Execution

Access the COMPSs Monitor through its webpage (<http://localhost:8080/compss-monitor> by default) and log in with the `TARGET_LOCAL_FOLDER` to enable the COMPSs Monitor for MareNostrum.

Please remember that to enable **all** the COMPSs Monitor features applications must be ran with the `-m` flag. For further information please check the *COMPSs User Manual: Application Execution* available at our webpage <http://compss.bsc.es>.

Figure 1 illustrates how to login and Figure 2 shows the COMPSs Monitor main page for an application run inside a Supercomputer.



Figure 1: COMPSs Monitor login for Supercomputers

Applications

2435288

Refresh

Resources information

Tasks information

Current tasks graph

Complete tasks graph

Load chart

Runtime log

Execution information

Statistics

Status	Resource Name	CPU Computing Units	GPU Computing Units	FPGA Computing Units	OTHER Computing Units	Memory Size	Disk Size	Provider	Image	Running Actions
	s04/2b83	16	-	-	-	28.0 GB	-GB	-	-	195 203 387 395 355 443 411 403 379 324 363 427 435 371 419 348
	s04/2b80	16	-	-	-	28.0 GB	-GB	-	-	226 210 218 275 315 299 339 307 283 259 267 331 243 251 235 292

Figure 2: COMPSs Monitor main page for a test application at Supercomputers

Please find more details on the COMPSs framework at
`http://compss.bsc.es`