

~ JUROPATEST ~

User's Manual for the Batch System

[Slurm integrated with Parastation]

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1 Cluster Information

1.1 Introduction

JUROPATEST is test system which allows users of JSC's current general purpose supercomputer JUROPA to port and optimize their applications for the new Haswell CPU architecture. Moreover, the system also serves as a development platform for system software and hardware with respect to the JUROPA collaboration. This cluster's main purpose is to prepare users and administrators for a smooth transition to the next JUROPA installation.

1.2 Cluster Nodes

For JUROPATEST cluster we have: 2 Master, 2 Login and 70 Compute nodes.

Type (Node Num.)	Hostname	CPU	Cores(SMT)	RAM	Description
Login (2)	juropatest1.zam.kfa-juelich.de (j3l03) juropatest2.zam.kfa-juelich.de (j3l04)	2x Intel Xeon E5-2695 v3 (Haswell) @ 2.3GHz	28 (56)	256 GB DDR4	Login Nodes
Master (2)	-	2x Intel Xeon E5-2695 v3 (Haswell) @ 2.3GHz	28 (56)	256 GB DDR4	Master Nodes
Compute (70)	j3c[061-130]	2x Intel Xeon E5-2695 v3 (Haswell) @ 2.3GHz	28 (56)	128 GB DDR4	Compute Nodes

The Operating System on JUROPATEST cluster is Scientific Linux release 6.5 (Carbon). For the management of the system we use the ParaStation Cluster Management software. Regarding the network, we use FDR Infiniband with a non-blocking Fat Tree topology.

1.3 Data Management - Filesystems

On JUROPATEST we provide GPFS shared filesystems. We provide home, scratch and archive filesystems, which have different purposes. The home filesystems are supposed to be used for user's data storage with the safety of backups (TSM backup), the scratch filesystem should be used as a fast storage for the data produced by the jobs (no backup and purged regularly) and the archive ones are to be used for long-term data archiving. Here is a small matrix with all filesystems available to the users:

Filesystem	Mount Point	Description
GPFS \$WORK	/work	Scratch filesystem – without backup
GPFS \$HOME	/homea /homeb /homec	Home filesystems – with TSM backup
GPFS \$ARCH	/arch /arch2	Archiving filesystems – with TSM backup. Available only on the login nodes.
GPFS \$DATA	/data	Special filesystem used only by certain groups – with TSM backup
User local binaries (GPFS)	/usr/local	Software repository available via module commands

The GPFS filesystems on JUROPATEST are mounted from JUST storage cluster. JUQUEEN and JUDGE users should be aware that they will work in the same \$HOME and \$WORK directories as on these production machines. The JUROPA successor will use the same home filesystems as JUROPATEST. Please note that JSC will do an automatic migration of all user data from Lustre to GPFS for \$HOME and \$WORK directories as soon as the JUROPA successor system is starting production. Therefore there is no need to copy any data ahead of this time. Only those data should be copied which is needed for the code porting and optimization process on JUROPATEST.

1.4 Access to the Cluster

Users can have access to the login nodes of the system only through SSH connections. As we described above there are two available login nodes. There is not a configured round-robin shared hostname between the login nodes. Users must explicitly define the login node they want to have access to. For example, to connect to the system, users must execute from their workstation the following command:

```
$ ssh username@juropatest1.fz-juelich.de
```

or

```
$ ssh username@juropatest2.fz-juelich.de
```

It is not possible to login by supplying username/password credentials. Instead, password-free login based on SSH key exchange is required. The public/private ssh key pair has to be generated on the workstation you are using for accessing JUROPATEST. On Linux or UNIX-based systems, the key pair can be generated by executing:

```
$ ssh-keygen -t [dsa|rsa]
```

It is required to protect the SSH key with a non-trivial pass phrase to fulfill the FZJ security policy. The generated public ssh key contained in the file “id_dsa.pub” or “id_rsa.pub” on user's workstation must be uploaded through the web interface from Dispatch when initially applying for a user account on JUROPA system. This SSH key afterwards will be automatically stored in the file “\$HOME/.ssh/authorized_keys” on the cluster.

1.5 Shell Environment

The default shell for all users on JUROPATEST is BASH (/bin/bash). After a successful login, user's shell environment is defined in files “\$HOME/.bash_profile” and “\$HOME/.bashrc”. Since the GPFS filesystems are shared between different clusters in JSC, that means the users' home directories are also shared on all system where the users have access to. This makes it more difficult for the users to create the correct or desired shell environment for each system. In order to solve this issue, a file has been created on all systems which contains a string with the system's name. The file is:

```
/etc/FZJ/systemname
```

This file is available on all login and compute nodes. The users can read this file and depending on the system they are logged-in they can set the desired environment. On JUROPATEST the string that is stored in that file is “juropatest”.

1.6 Modules

The installed software on JUROPATEST is organized through a hierarchy of modules. Loading a module adapts your environment variables to give you access to a specific set of software and its dependencies. The hierarchical organization of the modules ensures that you get a consistent set of dependencies, for example all built with the same compiler version or all relying on the same implementation of MPI. The module hierarchy is built upon toolchains. Toolchains modules in the lowest level contain just a compiler suite (like Intel compilers `icc` and `ifort`). Toolchains in the second level contain a compiler suite and a compatible implementation of MPI. The third and highest level contains "full toolchains", with a compiler suite, an MPI implementation, and compatible mathematical libraries such as SCALAPACK. An application is only accessible to the user when its module is loaded. You can load the application module only when the toolchain modules containing its dependencies are loaded first.

Modules and Toolchains hierarchy

If you know the dependencies of the application you would like to run, you can simply load a Toolchain module bundle from one of the three levels: Compilers, Compilers+MPI, or FullToolchains.

Here is a quick reference to the tools provided by each toolchain module:

Type	Modules available
Compilers	GCC: Gnu compilers with frontends for C, C++, Objective-C, Fortran, Java & Ada ifort: Intel Fortran compiler icc: Intel C and C++ compilers iccifort: icc/ifort (Intel C and Fortran compilers together)
Compilers+MPI	gpsmpi2: GCC + Parastation MPICH MPI ipsmpi2: icc/ifort + Parastation MPICH MPI iimpi: icc/ifort + Intel MPI
FullToolchains	gpsolf: gpsmpi + OpenBLAS, FFTW and ScaLAPACK intel-para: ipsmpi2 + Intel Math Kernel Library (imkl) intel: iimpi + Intel Math Kernel Library (imkl)

Using the module command

Users should load, unload and query modules through the module command. Several useful module commands are:

Command	Description
<code>module avail</code>	Shows the available toolchains and what modules are compatible to load right now according to the currently loaded toolchain.
<code>module load <modname>/<modversion></code>	Loads a specific module. Default version if it is not given.
<code>module list</code>	Lists what modules are currently loaded.
<code>module unload <modname>/<modversion></code>	Unloads a module.
<code>module purge</code>	Unloads all modules
<code>module spider <modname></code>	Finds the location of a module within the module hierarchy.

As we said above, in order to load a desired application module it is necessary first to load the correct toolchain. Therefore, preparing the module environment includes two steps:

1. First, load one of the available toolchains. The intel-para toolchain (from the Fulltoolchains) has the most supported software at this moment.
2. Second, load other application modules, which were built with currently loaded toolchain.

Following we will give some examples of the module command:

List the available toolchains:

```
$ module avail
----- /usr/local/software/juropatest/TC/FullToolchains -----
gpsolf/2014.11      intel/2014.11      intel-para/2014.11
----- /usr/local/software/juropatest/TC/Compilers+MPI -----
gpsmpi/2014.11     iimpi/7.1.2       ipsmpi/2014.11
----- /usr/local/software/juropatest/TC/Compilers -----
GCC/4.9.1          icc/2015.0.090    ifort/2015.0.090
...
```

Load a toolchain:

```
$ module load intel-para/2014.11
```

List all loaded modules from the current toolchain:

```
$ module list
Currently Loaded Modules:
 1) binutils/2.24      4) popt/1.14        7) iccifort/2015.0.090
 2) icc/2015.0.090    5) pscom/5.0.44-1  8) imkl/11.2.0.090
 3) ifort/2015.0.090  6) psmpi/5.1.0-1   9) intel-para/2014.11
```

List all application modules available for the current toolchain:

```
$ module avail
-- /usr/local/software/juropatest/Stagel/modules/all/MPI/intel/2015.0.090/psmpi/5.1.0-1 ---
Bison/2.7                Python/2.7.5
Bison/3.0.2              (D)  Python/2.7.8
Boost/1.53.0              (D)  Python/3.4.1
Boost/1.56.0              (D)  Qt/4.8.4
CMake/2.8.4               (D)  Qt/4.8.5
CMake/3.0.0               (D)  QuantumESPRESSO/5.1
Cube/4.2.3                SCOTCH/5.1.12b_esmumps
...
```

Get information about a package:

```
$ module spider Boost # or module spider Boost/1.56.0
```

Load an application module:

```
$ module load Boost/1.56.0
```

Unload all currently loaded modules:

```
$ module purge
```


1.7 Compilers

On JUROPATEST we offer some wrappers to the users, in order to compile and execute parallel jobs using MPI. Different wrappers are provided depending on the MPI version that is used. Users can choose the compiler's version using the module command (see the modules section).

The following table shows the names of the MPI wrapper procedures for the Intel compilers as well as the names of compilers themselves. The wrappers build up the MPI environment for your compilation task, so please always use the wrappers instead of the compilers:

Programming Language	Compiler	Parastation MPI Wrapper	Intel MPI Wrapper
<i>Fortran 90</i>	ifort	mpif90	mpiifort
<i>Fortran 77</i>	ifort	mpif77	mpiifort
C++	icpc	mpicxx	mpicpc
C	icc	mpicc	mpiicc

In the following table we present some useful compiler options that are commonly used:

Option	Description
-openmp	Enables the parallelizer to generate multi-threaded code based on the OpenMP directives.
-g	Creates debugging information in the object files. This is necessary if you want to debug your program.
-O[0-3]	Sets the optimization level.
-L	A path can be given in which the linker searches for libraries
-D	Defines a macro.
-U	Undefines a macro.
-I	Allows to add further directories to the include file search path.
-H	Gives the include file order. This options is very useful if you want to find out which directories are used and in which order they are applied.
-sox	Stores useful information like compiler version, options used etc. in the executable.
-ipo	Inter-procedural optimization.
-axCORE-AVX2	Indicates the processor for which code is created.
-help	Gives a long list of quite a big amount of options.

Compilation Examples

Compile an MPI program in C++:

```
$ mpicxx -O2 -o mpi_prog program.cpp
```

Compile a hybrid MPI/OpenMP program in C:

```
$ mpicc -openmp -o mpi_prog program.c
```

1.8 Batch model & Accounting

Following, we present the main policies concerning the batch model and accounting that are applied on JUROPATEST:

- Job scheduling according to priorities. The jobs with the highest priorities will be scheduled next.
- Backfilling scheduling algorithm. The scheduler checks the queue and may schedule jobs with lower priorities that can fit in the gap created by freeing resources for the next highest priority jobs.
- No node-sharing. The smallest allocation for jobs is one compute node. Running jobs do not disturb each other.
- For each project a Linux group is created where the users belong to. Each user has available contingent from one project only.
- CPU-Quota modes: monthly and fixed. The projects are charged on a monthly base or get a fixed amount until it is completely used.
- Contingent/CPU-Quota states for the projects: normal, low-contingent, no-contingent.
- Contingent priorities: normal > lowcont > nocont. Users without contingent get a penalty to the priorities of their jobs, but they are still allowed to submit and run jobs.

2 Batch System – Slurm

2.1 Slurm Overview

Slurm is the Batch System (Workload Manager) of JUROPATEST cluster. Slurm (Simple Linux Utility for Resource Management) is a free open-source resource manager and scheduler. It is a modern, extensible batch system that is widely deployed around the world on clusters of various sizes. A Slurm installation consists of several programs and daemons.

The Slurm control daemon (**slurmctld**) is the central brain of the batch system, responsible for monitoring the available resources and scheduling batch jobs. The **slurmctld** runs on an administrative node with a special setup to ensure availability of the services in case of hardware failures. Most user programs such as *srun*, *sbatch*, *salloc* and *scontrol* interact with the **slurmctld**. For the purpose of job accounting **slurmctld** communicates with Slurm database daemon (**slurmdbd**).

Slurm stores all the information about users, jobs and accounting data in its own database. The functionality of accessing and managing these data is implemented in **slurmdbd**. In our case, **slurmdbd** is configured to use a MySQL database as the back-end storage. To interact with **slurmdbd** and get information from the accounting database, Slurm provides commands like *sacct* and *sacctmgr*.

In contrast to the Moab/Torque combination where Moab provides scheduling and Torque performs resource management (like batch job start or node health monitoring) Slurm combines the functionality of the batch system and resource management. For this purpose Slurm provides the **slurmd** daemon which runs on the compute nodes and interacts with **slurmctld**. For the executing of user processes, **slurmstepd** instances are spawned by **slurmd** to shepherd the user processes. On JUROPATEST cluster no **slurmd**/**slurmstepd** daemons are running on the compute nodes. Instead the process management is performed by **psid** the management daemon from the Parastation Cluster Suite which has a proven track record on the JUROPA system. Similar to the architecture of the JUROPA resource management system, where a **psid** plugin called **psmom** replaces the Torque daemon on the compute nodes, a plugin of **psid** called **psslurm** replaces **slurmd** on the compute nodes of JUROPATEST. Therefore only one daemon is required on the compute nodes for the resource management which minimizes jitter (which can affect large-scale applications). For the end-users, there is no real difference visible because of this integration between Slurm and Parastation. Currently, **psslurm** is under active development by ParTec and JSC in the context of the JuRoPA collaboration.

The Batch System manages the compute **nodes**, which are the main resource entity of the cluster. Slurm groups the compute nodes into **partitions**. These partitions are the equivalent of queues in Moab. It is possible for different partitions to overlap, which means that the compute nodes can belong to multiple partitions. Also partitions can be configured with certain limits for the **jobs** that will be executed. Jobs are the allocations of resources by the users in order to execute tasks on the cluster for a specified period of time. Slurm introduces also the concept of **job-steps**, which are sets of (possibly parallel) tasks within the jobs. One can imagine job-steps as smaller allocations or jobs within the job, which can be executed sequentially or in parallel during the main job allocation.

In **Figure 1** we present the architecture of the daemons and their interactions with the user commands of Slurm.

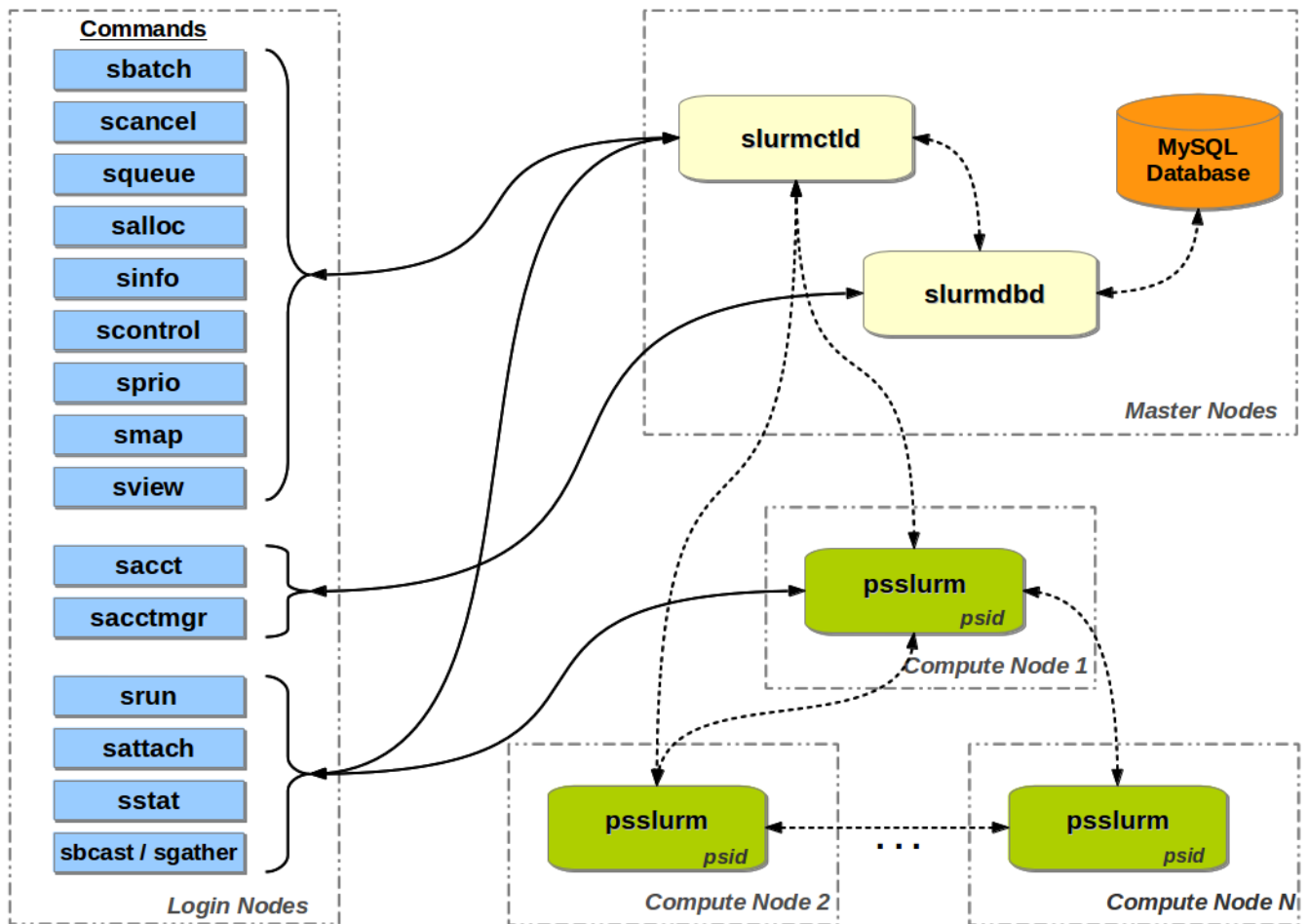


Figure 1.

2.2 Slurm Configuration

- High-Availability for the main controllers `slurmctld` and `slurmdbd`.
- Backfilling scheduling algorithm.
- No node-sharing.
- Job scheduling according to priorities.
- Accounting mechanism: `slurmdbd` with MySQL database as back-end storage.
- User and job limits enforced by QoS (Quality of Service) and some hard-limits configured in the partition settings. There is a QoS for each contingent state: normal, lowcont, nocont and suspended. Users without contingent are set to a different QoS and get a penalty for their job priorities.
- No preemption configured. Running jobs cannot be preempted.
- Prologue and Epilogue, with `pshealthcheck` from Parastation. The prologue checks the status of the nodes at job start and Epilogue cleans up the nodes after job completion.
- Same limits/configurations for batch and interactive jobs (no difference between batch and interactive jobs for Slurm, different behavior than Moab).

2.3 Partitions

In Slurm multiple nodes can be grouped into partitions which are sets of nodes with associated limits (for wall-clock time, job size, etc.). These limits are hard-limits for the jobs and can not be overruled by the specified limits in QoS's. Partitions may overlap and nodes may belong to more than one partition, making **partitions serve as general purpose queues**, like queues in Moab. The following table shows the partitions on JUROPATEST and the configured maximum limits and default values:

Partition	Limit	Value
<i>batch (default)</i>	Maximum wall-clock time for each job	2 hours
	Default wall-clock time for each job	30 minutes
	Minimum/Maximum number of nodes per job	1 / 8 nodes
	Default number of nodes for each job	1 node
	Maximum number of running jobs per user	4 jobs
<i>large</i>	Maximum wall-clock time for each job	1 hour
	Default wall-clock time for each job	30 minutes
	Minimum/Maximum number of nodes per job	9 / 64 nodes
	Default number of nodes for each job	9 nodes
	Maximum number of running jobs per user	4 jobs

The batch partition is intended for code optimization and small-scale performance tests. The large partition is intended for short running scalability tests. Jobs using the large partition only run in fixed time-slots (usually once per week depending on demand). The default partition is batch.

2.4 Slurm's Accounting Database

Slurm manages its own data in two different ways. First, there is a runtime engine in memory, backed-up with state files that is managed by `slurmctld` and second, there is the MySQL database that is managed by `slurmdbd`. Slurm stores all the important information in its MySQL database, like: cluster information, events, accounts, users, associations, QoS and job history. An association is the combination of cluster, account, user and partition. Associations are stored in a tree-like hierarchical structure starting with the root node with the accounts as its children and users as children of the accounts. In each association it is possible to specify fair-share, job limits and QoS.

To interact with `slurmdbd` and get accounting information from the database Slurm provides the commands `sacct` and `sacctmgr`.

2.5 Job Limits – QoS

As we describe above, the limits of the partitions are the hard-limits that put an upper limit for the jobs. However, the actual job limits are enforced by the limits specified in both partitions and Quality-of-Services, which means that first the QoS limits are checked/enforced, but these limits can never go over the partition limits.

One QoS is configured for each possible contingent status: *normal*, *lowcont*, *nocont*. These QoSs play the most important role to define the job priorities. By defining those QoSs the available range of priorities is separated into three sub-ranges, one for each contingent mode. Also one more QoS is defined with the name *suspended* which will be given to all associations that belong to users/projects that have ended and/or are not allowed to submit jobs anymore. Following we present the list with the configured Quality-of-Services:

Name	Priority	Flags	MaxNodes/Job	MaxWall/Job	MaxJobs/User	MaxSubmittedJobs
<i>normal</i>	100,000	DenyOnLimit	*Partition*	*Partition*	4	12
<i>lowcont</i>	50,000	DenyOnLimit	*Partition*	*Partition*	4	12
<i>nocont</i>	0	DenyOnLimit	*Partition*	*Partition*	4	12
<i>suspended</i>	0	DenyOnLimit	0	-	0	0

Note: For the entries that have **Partition** as value, it means that the limits are inherited from the Partitions where the jobs are running.

Each association in Slurm's database belongs to one user only. In each association there are two entries regarding the QoSs. One entry with the list of available QoSs and another entry with the Default-QoS (used when QoS is not specified with options). In every association only one available QoS is defined (same as default) for each user depending on the contingent status. This is implemented in JSC's accounting mechanism and the users are not allowed to change their QoS. The limits are enforced to the users by setting the correct QoS for their association according to their contingent. Job limits are enforced by that QoS in combination with the partition limits. If the users request allocations over the limits then the submission will fail (flag DenyOnLimit).

2.6 Priorities

Slurm schedules the jobs according to their priorities, which means that the jobs with the highest priorities will be executed next. With the backfilling algorithm though, jobs (usually small) with lower priorities can be schedule next if they can fit and run on the available resources before the next high-priority job is scheduled to start. Slurm has a very simple and well defined priority mechanism that allows us to define exactly the batch model we want. Following, we present how Slurm calculates the priorities for each job:

```
Job_priority = (PriorityWeightAge) * (age_factor) +
               (PriorityWeightFairshare) * (fair-share_factor) +
               (PriorityWeightJobSize) * (job_size_factor) +
               (PriorityWeightPartition) * (partition_factor) +
               (PriorityWeightQOS) * (QOS_factor)
```

Slurm uses five factors to calculate the job priorities: Age, Fairshare, Job-Size, Partition and QoS. The possible range of values for the factors is between 0.0 (min) and 1.0 (max). For each factor we have defined a weight that is used in the job-priority equation. Following is the list of weights we have configured:

Weight	Value
<i>WeightQOS</i>	100,000
<i>WeightAge</i>	32,500
<i>WeightJobSize</i>	14,500
<i>WeightFairshare</i>	3,000
<i>WeightPartition</i>	0

It is clear now that QoS plays an important role for the calculation of the priorities. With the different QoSs that have been defined, it is possible to create different priority range according to the contingent of the users. Below follows a table with the priority ranges for each contingent mode:

Contingent Status	Priority Ranges
<i>normal</i>	100,001 – 150,000
<i>lowcont</i>	50,001 – 100,000
<i>nocont</i>	0 – 50,000
<i>suspended</i>	–

For each contingent state the available range for priorities is 50k and is calculated from three factors: a) job age, b) job size and c) fair-share. In current setup, the partition factor is not used which means no difference in the priorities between different partitions.

2.7 Job Environment

On the compute nodes the whole shell environment is passed to the jobs during submission. With some options of the allocation commands, users can change this default behavior. The users can load modules and prepare the desired environment before job submission, and then this environment will be passed to the jobs that will be submitted. Of course, a good practice is to include module commands inside the job-scripts, in order to have full control of the environment of the jobs.

2.8 SMT

Similar to the Intel Nehalem processors in JUROPA, the Haswell processors in JUROPATEST offer the possibility of Simultaneous Multithreading (SMT) in the form of the Intel Hyper-Threading (HT) Technology. With HT enabled each (physical) processor core can execute two threads or tasks simultaneously. The operating system thus lists a total of 56 logical cores or Hardware Threads (HWT). Therefore a maximum of 56 processes can be executed on each compute node without overbooking.

Each compute node on JUROPATEST consists of two CPUs, located on socket zero and one, with 14 physical cores. These cores are numbered 0 to 27 and the hardware threads are named 0 to 55 in a round-robin fashion. **Figure 2** depicts a node schematically and illustrates the naming convention.

Node

Socket 0				Socket 1			
Core 0	Core 1	Core 2	Core 3	Core 14	Core 15	Core 16	Core 17
HWT 0	HWT 1	HWT 2	HWT 3	HWT 14	HWT 15	HWT 16	HWT 17
HWT 28	HWT 29	HWT 30	HWT 31	HWT 42	HWT 43	HWT 44	HWT 45
Core 4	Core 5	Core 6	Core 7	Core 18	Core 19	Core 20	Core 21
HWT 4	HWT 5	HWT 6	HWT 7	HWT 18	HWT 19	HWT 20	HWT 21
HWT 32	HWT 33	HWT 34	HWT 35	HWT 46	HWT 47	HWT 48	HWT 49
Core 8	Core 9	Core 10	Core 11	Core 22	Core 23	Core 24	Core 25
HWT 8	HWT 9	HWT 10	HWT 11	HWT 22	HWT 23	HWT 24	HWT 25
HWT 36	HWT 37	HWT 38	HWT 39	HWT 50	HWT 51	HWT 52	HWT 53
Core 12	Core 13			Core 26	Core 27		
HWT 12	HWT 13			HWT 26	HWT 27		
HWT 40	HWT 41			HWT 54	HWT 55		

Figure 2.

In chapter 4/section 4.1, there are examples (commands and job scripts) on how to use SMT.

3 Slurm User Commands

In this section we will give first a list of all commands with a short description and then later we will describe with more details the functionality of each command, giving also some examples.

3.1 List of Commands

Slurm offers a variety of user commands for all the necessary actions concerning the jobs. With these commands the users have a rich interface to allocate resources, query job status, control jobs, manage accounting information and to simplify their work with some utility commands.

Here is the list of all Slurm's user commands:

salloc is used to request interactive jobs/allocations. When the job is started a shell (or other program specified on the command line) is started on the submission host (login node). From the shell *srun* can be used to interactively spawn parallel applications. The allocation is released when the user exits the shell.

sattach is used to attach standard input, output, and error plus signal capabilities to a currently running job or job step. One can attach to and detach from jobs multiple times.

sbatch is used to submit a batch script (which can be a bash, Perl or Python script). The script will be executed on the first node in the allocation chosen by the scheduler. The working directory coincides with the working directory of the *sbatch* directory. Within the script one or multiple *srun* commands can be used to create job steps and execute (MPI) parallel applications. Note: *mpiexec* is not supported on JUROPATEST. *srun* is the only supported method to spawn MPI applications. In the future the *mpirun* command from Intel MPI may be supported.

scancel is used to cancel a pending or running job or job step. It can also be used to send an arbitrary signal to all processes associated with a running job or job step.

sbcast is used to transfer a file to all nodes allocated for a job. This command can be used only inside a job script.

sgather is used to transfer a file from all allocated nodes to the currently active job. This command can be used only inside a job script.

scontrol is primarily used by the administrators to view or modify Slurm configuration, like partitions, nodes, reservations, jobs, etc. However it provides also some functionality for the users to manage jobs or query and get some information about the system configuration.

sinfo is used to retrieve information about the partitions, reservations and node states. It has a wide variety of filtering, sorting, and formatting options.

smap graphically shows the state of the partitions and nodes using a curses interface. We recommend *llview* as an alternative which is supported on all JSC machines.

sprio can be used to query job priorities.

squeue allows to query the list of pending and running jobs. By default it reports the list of pending jobs sorted by priority and the list of running jobs sorted separately according to the job priority.

srun is used to initiate job steps mainly within a job or start an interactive job. *srun* has a wide variety of options to specify resource requirements. A job can contain multiple job steps executing sequentially or in parallel on independent or shared nodes within the job's node allocation.

sshare is used to retrieve fair-share information for each user.

sstat allows to query status information about a running job.

sview is a graphical user interface to get state information for jobs, partitions, and nodes.

sacct is used to retrieve accounting information about jobs and job steps. For older jobs *sacct* queries the accounting database.

sacctmgr is primarily used by the administrators to view or modify accounting information in Slurm's database. However, it allows also the users to query some information about their accounts and other accounting information.

Note: Man pages exist for all daemons, commands, and API functions. The command option “--help” also provides a brief summary of the available options.

3.2 Allocation Commands

sbatch & salloc

The commands *sbatch* and *salloc* can be used to allocate resources. *sbatch* is used for batch jobs. The arguments for the *sbatch* command is the allocation options followed by the jobscript. *sbatch* gets the allocation options either from the command line or from the job script (using #SBATCH directives). *salloc* is used to allocate resources for interactive jobs.

Command format:

```
sbatch [options] jobscript [args...]  
salloc [options] [<command> [command args]]
```

Here we present some useful options only for *sbatch* command:

Option	Description
-a <indexes> --array=<indexes>	Submit a job array (set of jobs). Each job can be identified by its index number.
--export=<env variables ALL NONE>	Specify which environment variables will be passed to the job. Default is ALL.
--ignore-pbs	Ignore any "#PBS" options in the job script.
--wrap=<command string>	Wraps a command in a simple "sh" shell script.
-d <dependency_list> --dependency=<dependency_list>	Delay the start of the job until the specified dependencies have been satisfied.

These three commands (*sbatch*, *salloc* and *srun*) share many allocation options. The most useful and commonly used allocation options are explained in following table:

Option	Description
--begin=<time>	Delay and schedule job after the specified time.
--cores-per-socket=<cores>	Allocate nodes with at least the specified number of cores per socket.
-c <ncpus> --cpus-per-task=<ncpus>	Number of logical CPUs (hardware threads) per task. This option is only relevant for hybrid/OpenMP jobs.
-D <directory>	Set the working directory of the job.
-e <filename pattern> --error=<filename pattern>	Path to the job's standard error. Slurm supports format strings containing replacement symbols such as %j (job ID).
-H --hold	Job will be submitted in a held state (zero priority). Can be released with "scontrol release <job_id>".
-i <filename pattern> --input=<filename pattern>	Connect the jobscript's standard input directly to the specified file.
-J <jobname> --job-name=<jobname>	Set the name of the job.
--mail-user	Define the mail address to receive mail notification.
--mail-type	Define when to send a mail notifications. Valid options: BEGIN, END, FAIL, REQUEUE or ALL.
-N <minnodes[-maxnodes]> --nodes=<minnodes[-maxnodes]>	Number of compute nodes used by the job. Can be omitted if --ntasks and --ntasks-per-node is given.
-n <number> --ntasks=<number>	Number of tasks (MPI processes). Can be omitted if --nodes and --ntasks-per-node is given.
--ntasks-per-core=<ntasks>	Number of tasks that will run on each CPU.
--ntasks-per-node=<ntasks>	Number of tasks per compute node.
-o <filename pattern> --output=<filename pattern>	Path to the job's standard output. Slurm supports format strings containing replacement symbols such as %j (job ID).
-p <partition_names> --partition=<partition_names>	Partition to be used. The argument can be either <i>batch</i> or <i>large</i> on JUROPATEST. If omitted, <i>batch</i> is the default.
--reservation=<name>	Allocate resources from the specified reservation.
-t <time> --time=<time>	Maximal wall-clock time of the job.
--tasks-per-node=<n>	Same as --ntasks-per-node.

Note: *srun* can also be used to start interactive jobs but we suggest to use *salloc*. *srun* should be used only to start job steps and spawn the processes (like MPI tasks) inside an allocation.

Implied allocation options

Depending on the combination of the allocation options that are used during submission, some other allocation options can be omitted because they are implied and the system calculates them automatically or the default values are used. Following there is table with these combinations:

Used options	Implied options (can be omitted)
--nodes & --ntasks	--ntasks-per-node
--nodes & --ntasks-per-node	--ntasks
--nodes	--ntasks (default is 1 task per node)
--ntasks	--nodes & --ntasks-per-node

Examples:

Submit a job requesting 2 nodes for 1 hour, with 28 tasks per node (implied value of ntasks: 56):

```
sbatch -N2 --ntasks-per-node=28 --time=1:00:00 jobscript
```

Submit a job-script allocating 4 nodes with 16 tasks in total (implied: 4 tasks per node) for 30 minutes:

```
sbatch -N4 -n16 -t 30 jobscript
```

Submit a job array of 4 jobs with 1 node per job, with the default walltime:

```
sbatch --array=0-3 -N1 jobscript
```

Submit a job-script in the large partition requesting 70 nodes for 2 hours:

```
sbatch -N70 -p large -t 2:00:00 jobscript
```

Submit a job without a job-script but wrapping a shell command:

```
sbatch -N4 -n4 --wrap="srun hostname"
```

Submit a job requesting the execution to start after the specified date:

```
sbatch --begin=2015-01-11T12:00:00 -N2 --time 2:00:00 jobscript
```

Submit a job requesting all available mail notifications to the specified email address:

```
sbatch -N2 --mail-user=myemail@address.com --mail-type=ALL jobscript
```

Specify a job name and the standard output/error files:

```
sbatch -N1 -J myjob -o MyJob-%j.out -e MyJob-%j.err jobscript
```

Start an interactive job and allocate 4 nodes for 1 hour:

```
salloc -N4 --time=60
```

Start an interactive job with *srun* and allocate 1 node for 10 minutes:

```
srun -N1 -t 10 --pty -u /bin/bash -i
```

3.3 Spawning commands

srun

With *srun* the users can spawn any kind of application, process or task inside a job allocation. It can be a shell command, any single-/multi-threaded executable in binary or script format, MPI application or hybrid application with MPI and OpenMP. When no allocation options are defined with *srun* command the options from *sbatch* or *salloc* are inherited.

srun should be used either,

1. Inside a job script submitted by *sbatch*.
2. Or after calling *salloc*.

Note: To start an application with Parastation MPI, the users should use only *srun* and not *mpiexec*. For Intel MPI, *mpirun* is not supported yet but it will be later.

Command format:

```
srun [options...] executable [args...]
```

The allocation options of *srun* for the job-steps are (almost) the same as for *sbatch* and *salloc* (please see the table above with allocation options). There are also some useful options only for *srun*:

Option	Description
--forward-x	Enable X11 forwarding <u>only for interactive jobs</u> .
--multi-prog <filename>	Run different programs with different arguments for each task specified in a text file.
--pty	Execute the first task in pseudo terminal mode.
-r <num> --relative=<num>	Execute a jobstep inside allocation with relative index of a node.
--exclusive	Allocate distinct cores for each task.

Examples:

Spawn 56 tasks on 4 nodes (14 tasks per node) for 30 minutes:

```
srun -N4 -n56 -t 30 executable
```

Spawn 8 tasks on 2 nodes (4 tasks per node), specifying in a file the executables for each task:

```
srun -n8 -N2 --multi-prog ./tasks.conf
---
./tasks.conf:
  0-3  hostname
  4-7  ./executable2
---
```

Inside a job-script, execute 4 tasks on 1 node without sharing cores with other job-steps:

```
srun --exclusive -n 4 -N1 mpi-prog
```

3.4 Query Commands

squeue

With *squeue*, we can see the current status information of the queued and running jobs.

Command format:

```
squeue [OPTIONS...]
```

Some of the most useful *squeue* options are:

Option	Description
-A <account_list> --account=<account_list>	List jobs for the specified accounts.
-a --all	Show information about jobs and job-steps for all partitions.
-r --array	Optimized display for job arrays.
-h --noheader	Do not print the header of the output.
-i <seconds> --iterate=<seconds>	Repeatedly print information at the specified interval.
-l --long	Report more information.
-o <output_format> --format=<output_format>	Specify the information that will be printed (columns). Please read the man pages for more information.
-p <part_list> --partition=<part_list>	List jobs only from the specified partitions.
-R <reservation_name> --reservation <reservation_name>	List jobs only for the specified reservation.
-S <sort_list> --sort=<sort_list>	Specify the order of the listed jobs.
--start	Print the expected start time for each job in the queue.
-t <state_list> --states=<state_list>	List jobs only with the specified state (failed, pending, running, etc).
-u <user_list> --user=<user_list>	Print the jobs of the specified user.

Examples:

Repeatedly print queue status every 4 seconds:

```
squeue -i 4
```

Show jobs in the large partition:

```
squeue -p large
```

Show jobs that belong to a specific user:

```
squeue -u user01
```

Print queue status with a custom format, showing only job ID, partition, user and job state:

```
squeue --format="%.18i %.9P %.8u %.2t"
```

Normally, the jobs will pass through several states during their life-cycle. Typical job states from submission until completion are: PENDING (PD), RUNNING (R), COMPLETING (CG) and COMPLETED (CD). However there are plenty of possible job states for Slurm. The following table describes the most common states:

State Code	State Name	Description
CA	CANCELLED	Job was explicitly cancelled by the user or an administrator. The job may or may not have been initiated.
CD	COMPLETED	Job has terminated all processes on all nodes.
CF	CONFIGURING	Job has been allocated resources, but is waiting for them to become ready for use.
CG	COMPLETING	Job is in the process of completing. Some processes on some nodes may still be active. Usually Slurm is running job's epilogue during this state.
F	FAILED	Job terminated with non-zero exit code or other failure condition.
NF	NODE_FAIL	Job terminated due to failure of one or more allocated nodes.
PD	PENDING	Job is awaiting resource allocation.
R	RUNNING	Job currently has an allocation. <u>Note</u> : Slurm is always running the prologue at the beginning of each job before the actual execution of user's application.
TO	TIMEOUT	Job terminated upon reaching its walltime limit.

sview

With *sview*, we get a graphical overview of the cluster. It shows information about system configuration, partitions, nodes, jobs, reservations. Some actions also are possible through the GUI. No options are available for *sview*. Users can just call the command and they will get the graphical window.

sinfo

With *sinfo*, we can get information and check the current state of partitions, nodes and reservations. This command is useful for checking the availability of the nodes.

Command format:

```
sinfo [OPTIONS...]
```

Some of the most useful *sinfo* options are:

Option	Description
-a --all	Show information about all partitions.
-d --dead	Show information only for the non-responding (dead) nodes.
-i <seconds> --iterate=<seconds>	Repeatedly print information at the specified interval.
-l --long	Report more information.
-n <nodes> --nodes=<nodes>	Show information only about the specified nodes.
-N --Node	Show information in a node-oriented format.
-o <output_format> --format=<output_format>	Specify the information that will be printed (columns). Please read the man pages for more information.
-p <partition> --partition=<partition>	Show information in a node-oriented format.
-r --responding	Show information only for the responding nodes.
-R --list-reasons	List the reasons why nodes are not in a healthy state.
-s --summarize	List partitions without many details for the nodes.
-t <states> --states=<states>	List nodes only with the specified state (e.g. allocated, down, drain, idle, maint, etc).
-T --reservation	Show information about the reservations.

Examples:

Show information about nodes in idle state:

```
sinfo -t idle
```

Show information about partitions and nodes in a summarized way:

```
sinfo -s
```

List all reservations:

```
sinfo -R
```

Show jobs that belong to a specific user:

```
sinfo -T
```

Show information for partition large:

```
sinfo -p large
```


Depending on the options, the *srun* command will print the states of the partitions and the nodes. The partitions may be in state UP, DOWN or INACTIVE. The UP state means that a partition will accept new submissions and the jobs will be scheduled. The DOWN state allows submissions to a partition but the jobs will not be scheduled. The INACTIVE state means that not submissions are allowed.

The nodes also can be in various states. Node state code may be shortened according to the size of the printed field. The following table shows the most common node states:

Shortened State	State Name	Description
<i>alloc</i>	ALLOCATED	The node has been allocated.
<i>comp</i>	COMPLETING	The job associated with this node is in the state of COMPLETING.
<i>down</i>	DOWN	The node is unavailable for use.
<i>drain</i>	DRAINING & DRAINED	While in DRAINING state any running job on the node will be allowed to run until completion. After that and in DRAIN state the node will be unavailable for use.
<i>idle</i>	IDLE	The node is not allocated to any jobs and is available for use.
<i>maint</i>	MAINT	The node is currently in a reservation with a flag of "maintenance".
<i>resv</i>	RESERVED	The node is in an advanced reservation and not generally available.

smap

With *smap*, we can get a graphical overview of the cluster. It shows information about the nodes and the jobs that are running on them.

Command format:

```
smap [OPTIONS...]
```

Some of the most useful *smap* options are:

Option	Description
-c --commandline	Send output to the command-line, without using curses.
-D <option> --display=<option>	Define the display mode of smap. Please read the man pages for more information.
-h --noheader	Do not print the header of the output.
-H --show_hidden	Show information about hidden partitions and their jobs.
-i <seconds> --iterate=<seconds>	Repeatedly print information at the specified interval.
-n <node_list> --nodes <node_list>	Show information only for the specified nodes.

sprio

With *sprio*, we can check the priorities of all pending jobs in the queue.

Command format:

```
sprio [OPTIONS...]
```

Some of the most useful *sprio* options are:

Option	Description
-h --noheader	Do not print the header of the output.
-j <job_id_list> --jobs=<job_id_list>	Show information only about the requested jobs.
-l --long	Report more information.
-n --norm	Print the the normalized priority factors of the jobs.
-o <output_format> --format=<output_format>	Specify the information that will be printed (columns). Please read the man pages for more information.
-u <user_list> --user=<user_list>	Show information about the jobs of the specified users.
-w --weights	Print the configured weights for each factor.

Examples:

Show information about priorities of all queued jobs in a long format:

```
sprio -l
```

Show priority information for job 777:

```
sprio -j 777
```

Show the priorities of all jobs that belong to the specified user:

```
sprio -u user1
```

Show priority information in a custom format, printing only job ID, priority and user:

```
sprio -o "%.7i %.10Y  %.8u"
```

scontrol

This command is primarily used by the administrators to manage Slurm's configuration. However it provides also some functionality for the users to manage jobs or query and get some information about the system configuration. Here we present the way to query and get various information with *scontrol*:

Command format:

```
scontrol [OPTIONS...] [COMMAND...]
```

Some of the most useful *scontrol* query commands are:

Command	Description
<code>show hostlist <host_list></code>	Return a compressed regular expression for the given comma separated host list.
<code>show hostlistsorted <host_list></code>	Return a compressed and sorted regular expression for the given comma separated host list.
<code>show hostnames <host_regex></code>	Expand the given regular expression to a full list of hosts.
<code>show job [<job_id>]</code>	Show information about all jobs or about the specified job.
<code>show node [<node_name>]</code>	Show information about all nodes or about the specified node.
<code>show partition [<partition_name>]</code>	Show information about all partitions or about the specified one.
<code>show reservation [<reservation_name>]</code>	Show information about all reservations or about the specified one.
<code>show step [<step_id>]</code>	Show information about all jobsteps or about the specified one.

Examples:

Expand and print a list of hostnames for the specified range:

```
scontrol show hostname j3c[061-070]
```

Show information about the job 777:

```
scontrol show job 777
```

Show information about the node j3c069:

```
scontrol show node j3c069
```

Show information about the partition batch:

```
scontrol show partition batch
```

sshare

With *sshare*, we can retrieve fairshare information and check the current value of the fairshare factor that is used to calculate the priorities of the jobs.

Command format:

```
sshare [OPTIONS...]
```

Some of the most useful options of *sshare* are:

Option	Description
<code>-A <account_list></code> <code>--accounts=<account_list></code>	Show information for the specified accounts. By default users belong only to one account.
<code>-h</code> <code>--noheader</code>	Do not display the header in the beginning of the output.
<code>-l</code> <code>--long</code>	Show more information.
<code>-p</code> <code>--parsable</code>	Print information in a parsable way. Delimit output with “ ”, with a “ ” in the end.
<code>-P</code> <code>--parsable2</code>	Print information in a parsable way. Delimit output with “ ”, without a “ ” in the end.

Examples:

Print information about the user's shares in a long format:

```
sshare -l
```

Print information about the user's shares in a parsable way:

```
sshare -P
```

Print information about the user's shares without the initial header in the output:

```
sshare -n
```

3.5 Job Control Commands

scancel

With *scancel*, we can signal or cancel jobs, job arrays or job steps.

Command format:

```
scancel [OPTIONS...] [job_id[_array_id][.step_id]...]
```

Some of the most useful options of the *scancel* command are:

Option	Description
-A <account> --account=<account>	Restrict the operation only to the jobs under the specified account.
-b --batch	Send a signal to the batch job shell and its child processes.
-i --interactive	Enables interactive mode. User must confirm for each operation.
-n <job_name> --name=<job_name>	Cancel a job with the specified name.
-p <partition_name> --partition=<partition_name>	Restrict the operation only to the jobs that are running in the specified partition.
-R <reservation_name> --reservation=<reservation_name>	Restrict the operation only to the jobs that are running using the specified reservation.
-s <signal_name> --signal=<signal_name>	Send a signal to the specified job(s).
-t <job_state_name> --state=<job_state_name>	Restrict the operation only to the jobs that have the specified state. Please check the man page.
-u <user_name> --user=<user_name>	Cancel job(s) only from the specified user. If no job ID is given then cancel all jobs of this user.

Examples:

Cancel jobs with ID 777 and 778:

```
scancel 777 778
```

Cancel jobs with the specified names:

```
scancel -n testjob1 testjob2
```

Cancel all jobs in queue (pending, running, etc.) from user1:

```
scancel -u user1
```

Cancel all jobs in partition large that belong to user1:

```
scancel -p large -u user1
```

Cancel all jobs from user1 that are in pending state:

```
scancel -t PENDING -u user1
```

scontrol

The *scontrol* command can be also used to manage and do some actions on the jobs:

Command	Description
hold <job_list>	Prevent a <u>pending</u> job from being started.
release <job_list>	Release a previously held job, so it can start.
notify <job_id> <message>	Send message to the standard error (stderr) of a job.

Examples:

Put jobs 777 and 778 in hold:

```
scontrol hold 777 778
```

Release job 777 from hold:

```
scontrol release 777
```

3.6 Job Utility Commands

sattach

With *sattach*, we can attach to a running job-step and get or manage the IO streams of the tasks in that job-step. By default (without options) it attaches to the standard output/error streams.

Command format:

```
sattach [options] <jobid.stepid>
```

Some of the most useful options of *sattach* are:

Option	Description
--input-filter[=]<task number> --output-filter[=]<task number> --error-filter[=]<task number>	Transfer the standard input or print the standard output/error only from the specified task.
-l --label	Add the task number in the beginning of each line of standard output/error.
--layout	Print the task layout information of the job-step without attaching to its I/O streams.
--pty	Run task number zero in pseudo terminal.

Examples:

Attach to the output of job 777 and job-step 1:

```
sattach 777.1
```

Attach to the output of job 777 and job-step 2, adding the task ID in the beginning of each line:

```
sattach -l 777.2
```

sstat

With sstat, we can get various status information about running job-steps, for example minimum, maximum and average values for metrics like CPU time, Virtual Memory (VM) usage, Resident Set Size (RSS), Disk I/O, Tasks number, etc.

Command format:

```
sstat [OPTIONS...]
```

Some of the most useful options of *sstat* are:

Option	Description
-a --allsteps	Show information about all steps for the specified job.
-e --helpformat	Show the list of fields that can be specified with the "--format" option.
-i --pidformat	Show information about the pids for each jobstep.
-j <job(.step)> --jobs <job(.step)>	Show information for the specified jobs or jobsteps.
-n --noheader	Do not display the header in the beginning of the output.
-o <field_list> --format=<field_list> --fields=<field_list>	Specify the comma separated list of fields that will be displayed in the output. Available fields can be found with "-e" option or in the man pages.
-p --parsable -P --parsable2	Print information in a parsable way. Output will be delimited with " ".

Examples:

Display default status information for job 777:

```
sstat -j 777
```

Display the defined metrics for job 777 in parsable format:

```
sstat -P --format=JobID,AveCPU,AvePages,AveRSS,AveVMSize -j 777
```

3.7 Job Accounting Commands

sacct

With *sacct*, we can get accounting information and data for the jobs and jobsteps that are stored in Slurm's accounting database. Slurm stores the history of all jobs in the database but each user has permissions to check only his/her own jobs.

Command format:

```
sacct [OPTIONS...]
```

Some of the most useful options of *sacct* are:

Option	Description
-b --brief	Show a brief listing, with the fields: jobid, status and exitcode.
-e --helpformat	Show the list of fields that can be specified with the “--format” option.
-E <end_time> --endtime=<end_time>	List jobs with any state (or with specified states using option “--state”) before the given date. Please check the man pages for the available time formats.
-j <job(.step)> --jobs=<job(.step)>	Show information only for the specified jobs/job-steps.
-l --long	Show full report with all available fields for each reported job/job-step.
-n --noheader	Do not display the header in the beginning of the output.
-N <node_list> --nodelist=<node_list>	Show information only for jobs that ran on the specified nodes.
--name=<jobname_list>	Show information about jobs with the specified names.
-o <field_list> --format=<field_list>	Specify the list of fields that will be displayed in the output. Available fields can be found with “-e” option or in the man pages.
-r <partition_name> --partition=<partition_name>	Show information only for jobs that ran in the specified partitions. Default is all partitions.
-s <state_list> --state=<state_list>	Filter and show information only about jobs with the specified states, like completed, cancelled, failed, etc. Please check the man pages for the full list of states.
-S <start_time> --starttime=<start_time>	List jobs with any state (or with specified states using option “--state”) after the given date. The default value is 00:00:00 of current date. Check man page for date formats.
-X --allocations	Show information only for jobs and not for job-steps.

Examples:

Show job information in long format for default period (starting from 00:00 today until now):

```
sacct -l
```

Show job only information (without jobsteps) starting from the defined date until now:

```
sacct -S 2014-10-01T07:33:00 -X
```

Show job and jobstep information printing only the specified fields:

```
sacct -S 2014-10-01 --format=jobid,elapsed,nnodes,state
```

sacctmgr

The *sacctmgr* command is mainly used by the administrators to view or modify accounting information and data in the accounting database. This command provides also an interface with limited permissions to the users for some querying actions.

Command format:

```
sacctmgr [OPTIONS...] [COMMAND...]
```

Some of the most useful commands for *sacctmgr* are:

Command	Description
show/list* cluster	Show cluster information.
show association [where user=<name>]	List all visible associations or the ones for the specified user.
show event [where node=<node_name>]	List all events for all or for the specified nodes.
show qos [where name=<qos_name>]	List all or the specified QoS.
show user	Show some user information, like privileges, etc.

* “show” and “list” commands are the same for *sacctmgr*.

Examples:

Show cluster information:

```
sacctmgr show cluster
```

Show the association of user1:

```
sacctmgr show association where user=user1
```

Print all QoSs:

```
sacctmgr show qos
```

Show the privileges of my user:

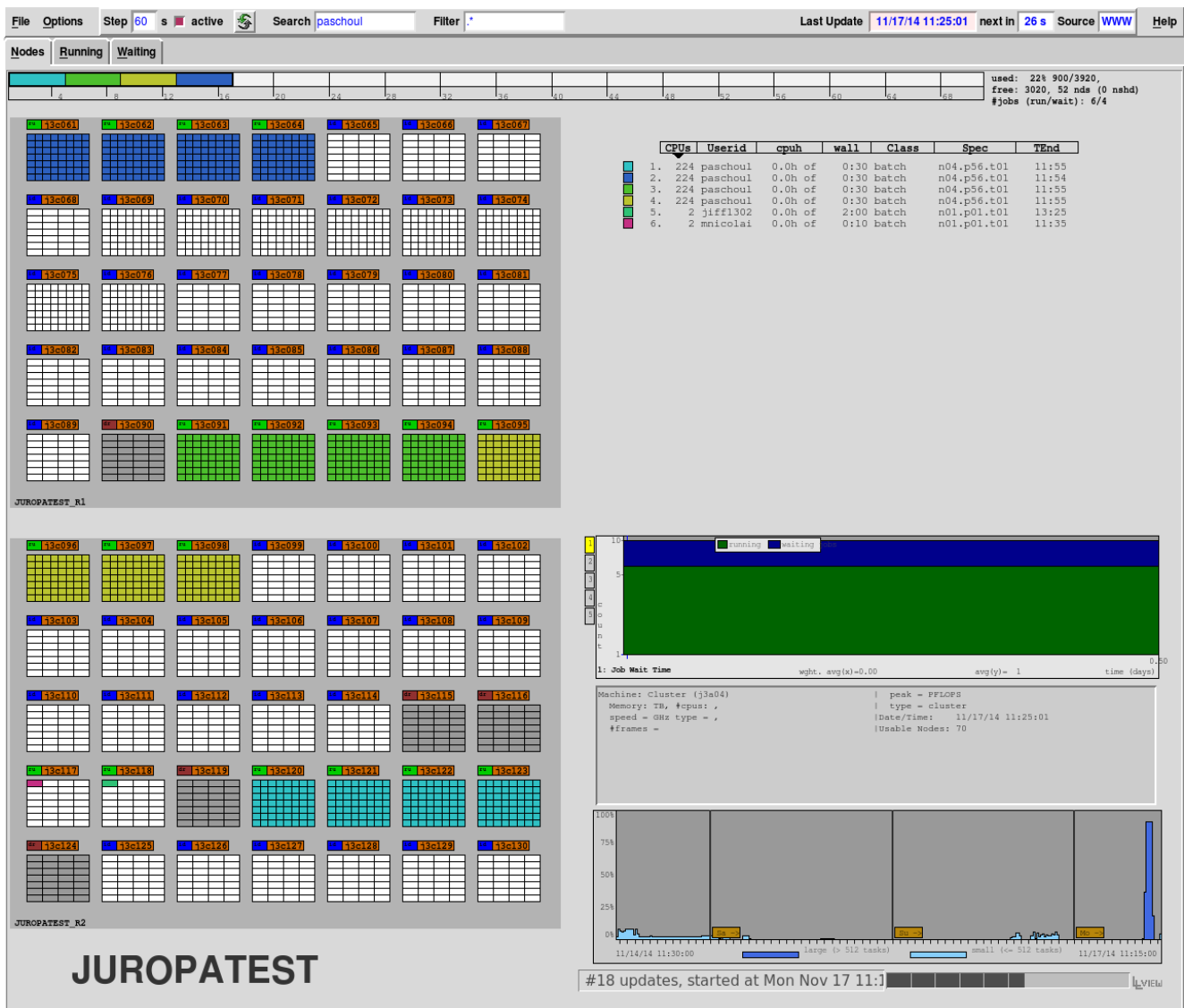
```
sacctmgr show user
```

3.8 Custom commands from JSC

llview

llview is a cluster monitoring tool implemented in JSC that shows a graphical overview of the cluster. The nodes are grouped and presented per rack, and different coloring is used per job for each allocation on the nodes. The GUI shows the list of all current jobs in the queue, and gives also information about the utilization of the cluster.

Below in **Figure 3** there is a screenshot of *llview*:



q_cpuquota

Job accounting is done via a central database in JSC and the information about JUROPATEST jobs will be completed once per day around midnight, based on information obtained from Slurm Accounting Database. Users get information about their current quota-test status or the usage of single jobs by using the command *q_cpuquota*.

Command format:

```
q_cpuquota [OPTIONS...]
```

Some useful options of the *q_cpuquota* command are:

Option	Description
-?	Print usage information.
-h <cluster>	Show information for the specified system (e.g. JUROPATEST).
-j <jobID>	Show accounting information for the specified job.
-t <time>	Show information about jobs in the specified time period.
-d <number>	Show information about jobs of the last specified days.

4 Batch Jobs

Users submit batch applications (usually bash scripts) using the *sbatch* command. In the job scripts, in order to define the *sbatch* parameters #SBATCH directives must be used. The script is executed on the first compute node in the allocation. To execute parallel MPI tasks users call *srun* within their script. With *srun* users can also create job-steps. A job step can allocate the whole or a subset of the already allocated resources from *sbatch*. With these commands Slurm offers a mechanism to allocate resources for a certain walltime and then run many parallel jobs in that frame. The following table describes the most common or necessary allocation options that can be defined in a job script:

Option	Default value	Description
#SBATCH --nodes=<number> #SBATCH -N <number>	1	Number of nodes for the allocation.
#SBATCH --ntasks=<number> #SBATCH -n <number>	1	Number of tasks (MPI processes). Can be omitted if --nodes and --ntasks-per-node are given.
#SBATCH --ntasks-per-node=<num> #SBATCH --tasks-per-node=<num>	1	Number of tasks per node. If keyword omitted the default value is used, but there are still available maximum 56 CPUs per node for current allocation.
#SBATCH --cpus-per-task=<num> #SBATCH -c <num>	1	Number of threads/VCores per task. Used only for OpenMP or hybrid jobs.
#SBATCH --output=<path> #SBATCH -o <path>	slurm-<jobID>.out	Path to the file for the standard output.
#SBATCH --error=<path> #SBATCH -e <path>	slurm-<jobID>.out	Path to the file for the standard error.
#SBATCH --time=<walltime> #SBATCH -t <walltime>	30 minutes	Requested walltime limit for the job.
#SBATCH --partition=<name> #SBATCH -p <name>	batch	Partition to run the job. Currently available: batch and large partitions.
#SBATCH --mail-user=<email>	username	Email address for notifications.
#SBATCH --mail-type=<mode>	NONE	Event types for email notifications.
#SBATCH --job-name=<jobname> #SBATCH -J <jobname>	jobscript's name	Job name.

Multiple *srun* calls can be placed in a single batch script. Options such as --nodes, --ntasks and --ntasks-per-node are by default taken from the *sbatch* arguments but can be overwritten for each *srun* invocation. If --ntasks-per-node is omitted or set to a value higher than 28 then SMT (simultaneous multi-threading) will be enabled. Each compute node has 28 physical cores and 56 logical cores.

As we described before, the job script is submitted using:

```
sbatch [OPTIONS] <jobscript>
```

On success, *sbatch* writes the job ID to standard out.

Note: In case some allocation options are defined in both command-line and inside the job-script, then the options that were given as arguments in the command-line will be used and the options in the job-script will be ignored.

4.1 Job script examples

Serial job

Example 1: Here is a simple example where some system commands are executed inside the job script. This job will have the name “TestJob”. One compute node will be allocated for 30 minutes. Output will be written in the defined files. The job will run in the default partition *batch*.

```
#!/bin/bash
#SBATCH -J TestJob
#SBATCH -N 1
#SBATCH -o TestJob-%j.out
#SBATCH -e TestJob-%j.err
#SBATCH --time=30

sleep 5
hostname
```

Parallel job

In order to start a parallel job, users have to use the *srun* command that will spawn processes on the allocated compute nodes of the job. Options given to *srun* will override the allocation option from *sbatch*. In case of no *srun* options the defined options (with #SBATCH) or the defaults will be used.

Example 2: Here is a simple example of a job script where we allocate 4 compute nodes for 1 hour. Inside the job script, with the *srun* command we request to execute on 2 nodes with 1 process per node the system command *hostname* in a time-frame of 10 minutes.

```
#!/bin/bash
#SBATCH -J TestJob
#SBATCH -N 4
#SBATCH -o TestJob-%j.out
#SBATCH -e TestJob-%j.err
#SBATCH --time=60

srun -N2 --ntasks-per-node=1 -t 10 hostname
```

OpenMP job

Example 3: In this example the job will execute an OMP application named “*omp-prog*”. The allocation is for 1 node and by default, since there is no node-sharing, all CPUs of the node are available for the application. The output filenames are also defined and a walltime of 2 hours is requested. Note: It is important to define and export the variable `OMP_NUM_THREADS` that will be used by the executable.

```
#!/bin/bash
#SBATCH -J TestOMP
#SBATCH -N 1
#SBATCH -o TestOMP-%j.out
#SBATCH -e TestOMP-%j.err
#SBATCH --time= 02:00:00

export OMP_NUM_THREADS=56

/home/user/test/omp-prog
```

MPI job

Example 4: In the following example, an MPI application will start 112 tasks on 4 nodes running 28 tasks per node (no SMT) requesting a walltime limit of 15 minutes in batch partition. Each MPI task will run on a separate core of the CPU.

```
#!/bin/bash

#SBATCH --nodes=4
#SBATCH --ntasks=112
#SBATCH --output=mpi-out.%j
#SBATCH --error=mpi-err.%j
#SBATCH --time=00:15:00
#SBATCH --partition=batch

srun -N4 --ntasks-per-node=28 ./mpi-prog
```

MPI jobs with SMT

On each node there are 28 real cores available and, with SMT enabled, 56 virtual cores. In order to enable SMT the users just have to request from Slurm to allocate more than 28 CPUs on each compute node. Following there are some examples where SMT is enabled:

Example 5: In this example we have an MPI application starting 1792 tasks in total on 32 nodes using 56 logical CPUs (hardware threads) per node (SMT enabled) requesting a time period of 20 minutes. The *large* partition is used.

```
#!/bin/bash -x

#SBATCH --nodes=32
#SBATCH --ntasks=1792
#SBATCH --ntasks-per-node=56 # can be omitted #
#SBATCH --output=mpi-out.%j
#SBATCH --error=mpi-err.%j
#SBATCH --time=00:20:00
#SBATCH --partition=large

srun ./mpi-prog
```

Example 6: In this example, the job script will start the program “mpi-prog” on 4 nodes using 56 MPI tasks per node, where two MPI tasks will be executed on each physical core.

```
#!/bin/bash

#SBATCH --nodes=4
#SBATCH --ntasks=224 # can be omitted #
#SBATCH --ntasks-per-node=56
#SBATCH --output=mpi-out.%j
#SBATCH --error=mpi-err.%j
#SBATCH --time=00:15:00
#SBATCH --partition=batch

srun ./mpi-prog
```

Hybrid Jobs

Example 7: In this example, a hybrid MPI/OpenMP job is presented. This job will allocate 5 compute nodes for 2 hours. The job will have 35 MPI tasks in total, 7 tasks per node and 4 OpenMP threads per task. On each node 28 cores will be used (no SMT enabled). Note: It is important to define the environment variable `OMP_NUM_THREADS` and this must match with the value that was given to the option “`--cpus-per-task`”.

```
#!/bin/bash
#SBATCH -J TestJob
#SBATCH -N 5
#SBATCH -o TestJob-%j.out
#SBATCH -e TestJob-%j.err
#SBATCH --time= 02:00:00
#SBATCH --partition=large

export OMP_NUM_THREADS=4

srun -N 5 --ntasks-per-node=7 --cpus-per-task=4 ./hybrid-prog
```

Example 8: In this example, there is a hybrid application which will start 4 tasks per node on 2 allocated nodes and starting 14 threads per node (no SMT). In order to set the environment variable “`OMP_NUM_THREADS`”, Slurm's variable “`SLURM_CPUS_PER_TASK`” is used which is defined by the option “`--cpus-per-task`”.

```
#!/bin/bash
#SBATCH -N 4
#SBATCH -n 8 # can be omitted #
#SBATCH --ntasks-per-node=2
#SBATCH --cpus-per-task=7
#SBATCH --output=mpi-out.%j
#SBATCH --error=mpi-err.%j
#SBATCH --time=00:20:00
#SBATCH --partition=batch

export OMP_NUM_THREADS=${SLURM_CPUS_PER_TASK}
srun ./hybrid-prog
```

Hybrid jobs with SMT

Example 9: This example shows a hybrid application that will start 4 tasks per node on 3 allocated nodes and starting 14 threads per task, using in total 56 cores per node (SMT enabled).

```
#!/bin/bash
#SBATCH --nodes=3
#SBATCH --ntasks=12
#SBATCH --ntasks-per-node=4
#SBATCH --cpus-per-task=14
#SBATCH --output=mpi-out.%j
#SBATCH --error=mpi-err.%j
#SBATCH --time=00:20:00

export OMP_NUM_THREADS=${SLURM_CPUS_PER_TASK}
srun ./hybrid-prog
```

Example 10: This example presents a hybrid application which will execute “hybrid-prog” on 3 nodes using 2 MPI tasks per node and 28 OpenMP threads per task (56 CPUs per node).

```
#!/bin/bash
#SBATCH --nodes=3
#SBATCH --ntasks=6
#SBATCH --ntasks-per-node=2
#SBATCH --cpus-per-task=28
#SBATCH --output=mpi-out.%j
#SBATCH --error=mpi-err.%j
#SBATCH --time=00:20:00
#SBATCH --partition=batch

export OMP_NUM_THREADS=${SLURM_CPUS_PER_TASK}
srun ./hybrid-prog
```

Intel MPI jobs

In order to run Intel MPI jobs user can use *srun*. The *mpirun* command is currently not supported. That means for now the users can not export and use the environment variables from Intel MPI, because *srun* does not work with them. Users will be informed when *mpirun* will be supported.

4.2 Job steps

In a previous chapter we described job-steps as small allocations or jobs inside the current job. Each call of *srun* will create a new job-step. It is up to the users to decide how they will create job-steps. It is possible to have one job-step after another using all the allocated nodes each time, or to have many job-steps running in parallel. Instead of submitting many single-node jobs, known as farming, it is suggested to the users to do farming using job-steps inside a single job. In this case, since all CPUs are available to the job, the only bounding factor is the memory per task (and the walltime). The users will be accounted for all the nodes of the allocation regardless if all nodes are used for job-steps or not.

Example 11: In the following example it is presented how to execute MPI programs in different job-steps sequentially inside a job allocation. In total 4 nodes are allocated for 2 hours. In this job 3 job-steps will be created. The first job-step will run on 4 nodes having 1 MPI task per node for 20 minutes. After that the second job-step will be executed on 3 nodes with 28 MPI tasks per node for 1 hour. And in the end the last job-step will run on 4 nodes with 56 MPI tasks per node using all virtual cores on each node (SMT) and it will finish when the MPI application will be completed or will be canceled by the scheduler if it will reach the walltime limit.

```
#!/bin/bash

#SBATCH --nodes=4
#SBATCH --output=mpi-out.%j
#SBATCH --error=mpi-err.%j
#SBATCH --time=02:00:00

srun -N4 --ntasks-per-node=1 --time=00:20:00 ./mpi-prog1
srun -N3 --ntasks-per-node=28 --time=01:00:00 ./mpi-prog2
srun -N4 --ntasks-per-node=56 ./mpi-prog3
```


Example 12: In the following example we show a job script where two different job-steps are initiated within one job. In total 28 cores are allocated on two nodes. Each job step uses 14 cores on one of the compute nodes. Here the job-steps will be executed in parallel:

```
#!/bin/bash

#SBATCH --nodes=2
#SBATCH --ntasks=28
#SBATCH --ntasks-per-node=14
#SBATCH --output=mpi-out.%j
#SBATCH --error=mpi-err.%j
#SBATCH --time=00:20:00

srun -N1 -n 14 ./mpi-prog1 &
srun -N1 -n 14 ./mpi-prog2 &

wait
```

4.3 Dependency Chains

Slurm supports dependency chains which are collections of batch jobs with defined dependencies, similar to job chains of Moab on JUROPA. Job dependencies can be defined using the `--dependency` argument of *sbatch*. The format is:

```
sbatch --dependency=<type>:<jobID> <jobscrip>
sbatch -d <type>:<jobID> <jobscrip>
```

The available dependency types for job-chains are: *after*, *afterany*, *afternotok* and *afterok*. For more information please check the man page of *sbatch*.

Example 13: Below is an example of a job-script for the handling of job chains. The script submits a chain of “`$NO_OF_JOBS`”. A job will only start after successful completion of its predecessor. Please note that a job which exceeds its time-limit is not marked successful.

```
#!/bin/bash -x
# submit a chain of jobs with dependency

# number of jobs to submit
NO_OF_JOBS=<no of jobs>

# define jobscrip
JOB_SCRIPT=<jobscrip>

echo "sbatch ${JOB_SCRIPT}"
JOBID=$(sbatch ${JOB_SCRIPT} 2>&1 | awk '{print $(NF)}')

I=0
while [ ${I} -le ${NO_OF_JOBS} ]; do
    echo "sbatch -d afterok:${JOBID} ${JOB_SCRIPT}"
    JOBID=$(sbatch -d afterok:${JOBID} ${JOB_SCRIPT} 2>&1 | awk '{print $(NF)}')
    let I=${I}+1
done
```

4.4 Job Arrays

Slurm supports job-arrays and offers a mechanism to easily manage these collections of jobs. Job arrays are only supported for the *sbatch* command and, as we described previously, they can be defined using the options “--array” or “-a”. To address a job-array, Slurm provides a base array ID and an array index unique for each job. The format for specifying an array job is first the base array jobID followed by “_” and then the array index:

```
<base job id>_<array index>
```

Slurm exports two environment variables that can be used in the job script to identify each array-job:

```
SLURM_ARRAY_JOB_ID  # base array job ID
SLURM_ARRAY_TASK_ID # array index
```

Some additional options are available to specify the *stdin*, *stdout*, and *stderr* file names: option “%A” will be replaced by the value of `SLURM_ARRAY_JOB_ID` and option “%a” will be replaced by the value of `SLURM_ARRAY_TASK_ID`.

Also each job in an array has its own normal unique job ID. This ID is exported in the environment variable

```
SLURM_JOBID
```

Example 14: In the following example, the job-script will create a job array of 4 jobs with indices 0-3. Each job will run on 1 node with walltime of 1 hour and will execute a different bash script (`script_[0-3].sh`).

```
#!/bin/bash
#SBATCH --nodes=1
#SBATCH --output=prog-%A_%a.out
#SBATCH --error=prog-%A_%a.err
#SBATCH --time=01:00:00
#SBATCH --array=0-3

./script_${SLURM_ARRAY_TASK_ID}.sh
```

Example 15: In the following job-script a job array of 20 jobs will be submitted with indices 1-20. Each job will run on a separate node with 2 hours walltime limit. Some may be running and some may be waiting in the queue. For this job array all jobs will execute the same binary “prog” with different input files (`input_[1-20].txt`):

```
#!/bin/bash -x
#SBATCH --nodes=1
#SBATCH --output=prog-%A_%a.out
#SBATCH --error=prog-%A_%a.err
#SBATCH --time=02:00:00
#SBATCH --array=1-20
#SBATCH --partition=large

srun -N1 --ntasks-per-node=1 ./prog input_${SLURM_ARRAY_TASK_ID}.txt
```

4.5 MPMD

Slurm supports the MPMD model (Multiple Program Multiple Data Execution Model) that can be used for MPI applications, where multiple executables can have one common `MPI_COMM_WORLD` communicator. For this purpose Slurm provides the option “`--multi-prog`” for the `srun` command only. This option expects a configuration text file as an argument and the format is:

```
srun [OPTIONS..] --multi-prog <text-file>
```

Each line of the configuration file can have two or three possible fields separated by space and the format is like this:

```
<list of task ranks> <executable> [<possible arguments>]
```

In the first field is defined a comma separated list of ranks for the MPI tasks that will be spawned. Possible values are integer numbers or ranges of numbers. The second field is the path/name of the executable. And the third field is optional and defines the arguments of the program.

Example 16: In this example there is a simple configuration file with name “*multi.conf*”. This file defines three MPI programs. For the first executable `mpi-prog1` only one instance will be executed with rank 0 and one integer argument. For the second program `mpi-prog2` Slurm will create two tasks with ranks 4 and 6 and each one will have the path of a file as argument. For the third program `mpi-prog3` five MPI tasks will be executed with ranks 1, 2, 3, 5 and 7 without any arguments.

```
0          ./mpi-prog1 0
4,6        ./mpi-prog2 ./tmp.txt
1-3,5,7    ./mpi-prog3
```

Following is the job-script that will start this MPMD job. The job-script allocates 4 nodes for 1 hour. The command `srun` will start this MPMD application, where all 4 nodes will be used with 2 MPI tasks per node (8 tasks in total). It can be submitted with `sbatch`:

```
#!/bin/bash
#SBATCH --nodes=4
#SBATCH --time=01:00:00

srun -N4 --ntasks-per-node=2 --multi-prog ./multi.conf
```

The “`--multi-prog`” option can be used of course for any kind of binary and its usage is not restricted to MPI jobs only, but it is the only way to apply the MPMD model.

5 Interactive Jobs

5.1 Interactive Session

Interactive sessions can be allocated using the *salloc* command. The following command for example will allocate 2 nodes for 30 minutes:

```
salloc --nodes=2 --time=00:30:00
```

Once an allocation has been made, the *salloc* command will start a bash on the login node where the submission was done. After a successful allocation the users can execute *srun* from that shell and they can spawn interactively their applications. For example:

```
srun --ntasks=4 --ntasks-per-node=2 --cpus-per-task=7 ./hybrid-prog
```

The interactive session is terminated by exiting the shell. In order to obtain a shell on the first allocated compute nodes (like command “*msub -I*” from Moab), the users can start a remote shell from within the current session and connect it to a pseudo terminal (pty) using the *srun* command with a shell as an argument. For example:

```
srun --nodes=2 --pty /bin/bash
```

After gaining access to the remote shell it is possible to run *srun* again from that remote shell in order to execute interactively applications without any delays (no scheduling delays since the allocation has already been granted). Below follows a transcript of an exemplary interactive session:

```
$ salloc --nodes=2 --time=00:01:00
salloc: Pending job allocation 4749
salloc: job 4749 queued and waiting for resources
salloc: job 4749 has been allocated resources
salloc: Granted job allocation 4749

$ hostname
j3l03

$ srun --ntasks 2 --ntasks-per-node=2 hostname
j3c061
j3c062

$ srun --nodes=1 --ntasks=1 --pty /bin/bash -i

$ hostname
j3c061

$ logout

$ hostname
j3l03

$ exit
exit
salloc: Relinquishing job allocation 4749
salloc: Job allocation 4749 has been revoked.
```

5.2 X Forwarding

The X11 forwarding support has been implemented with the “`--forward-x`” option of the *srun* command. It is similar to the option “`msub -x`” from Moab. X11 forwarding is required for users who want to use applications or tools which provide a GUI.

Here is an example that shows how to use this feature:

```
$ salloc --nodes=1 --time=00:01:00
...

$ srun --nodes=1 --ntasks=1 --forward-x --pty /bin/bash -i

$ ./GUI-App
```

Note: User accounts will be charged per allocation whether the compute nodes are used or not. Batch submission is the preferred way to execute jobs.

6 From Moab/Torque to Slurm

On JUROPA we are using the combination of Moab and Torque for the Batch System. Moab works as the scheduler and Torque is the resource manager. However, on JUROPATEST and later on the next Juropa installation (Juropa Successor) we will use Slurm as scheduler and resource manager. In this chapter we will compare and give some information about these two solutions and we will try to help the users have an easier migration from Moab/Torque to Slurm.

6.1 Differences between the Systems

Here we will compare and declare some differences between Moab and Slurm:

	Moab	Slurm
Resource Management	Not supported. Needs an external Resource Manager (like Torque).	A flexible and capable resource manager (in our case psslurm on the nodes).
Nodes	It is possible to set nodes for batch and interactive jobs only, or both.	No difference between batch and interactive jobs for Slurm.
Queues	Partitions separate node into groups. Queues are used for job submission on one partition only.	Slurm defines only partitions. For Slurm the partitions are used as queues. Partitions can overlap and we can specify limits.
Priorities	Complex priorities mechanism.	Easy to configure, maintain and manage. The desired batch model from JSC can be easily applied.
Limits/Policy	Good support for limits and policies configuration.	Highly configurable: define limits and policies per partition/account/user. Enforce limits with QoS.
Job scripts	Define job-script options with #MSUB.	Define job-script options with #SBATCH.

In the following table you can see some of the differences between Torque and Slurm:

	Torque	Slurm
Scheduling	Integrates only a simple FIFO scheduler, needs external scheduler.	Slurm is a capable scheduler with support for backfilling algorithm.
Output files	With the default options, stores output locally on the nodes. Upon completion files are gathered at destination.	Standard output and error files are created in the final destination immediately.
Working directory	Must explicitly change to current working directory.	Jobs start to run in the directory where they were submitted from.
Job Steps	Not supported by Torque.	Flexible allocations within jobs.
Task Distribution	Possible to specify different number of tasks per set of nodes, e.g.: “-l nodes=1:ppn=2+nodes=4:ppn=8”	Possible to specify only the same number of tasks on all nodes with the allocation options.
Environment	If users want to export the whole shell environment, they must use the option “-V”.	The environment defined in user's shell during submission will be automatically exported to the job.

6.2 User Commands Comparison

The following table presents commands with similar functionality from Slurm, Moab and Torque:

User Commands	Slurm	Moab	Torque
Job Submission	sbatch	msub	qsub
Job deletion	scancel	canceljob	qdel
Job status	squeue scontrol show job	checkjob	qstat
Job hold	scontrol hold	mjobctl -h	qhold
Job release	scontrol release	mjobctl -u	qrls
Queue list	squeue	showq	qstat -Q
Cluster status	sinfo	---	qstat -a
Node list	scontrol show nodes	---	pbsnodes -l
GUI	sview	---	xpbsmon

The table below compares the allocation options of *msub* and *sbatch*:

Allocation option	Moab/Torque (msub)	Slurm (sbatch)
Number of nodes	-l nodes=<number>	--nodes=<number> -N <number>
Number of total tasks	None	--ntasks=<number> -n <number>
Number of tasks/cpus per node	-l ppn=<number>	--ntasks-per-node=<num> --tasks-per-node=<num>
Number of threads per task	-v tpt=<number>	--cpus-per-task=<num> -c <num>
File for the standard output	-o <path>	--output=<path> -o <path>
File for the standard error	-e <path>	--error=<path> -e <path>
Walltime limit	-l walltime=<time>	--time=<walltime> -t <walltime>
Partition/Queue selection	-q <queue>	--partition=<queue> -p <queue>
Email for notifications	-M <email>	--mail-user=<email>
Event types for notifications	-m <mode>	--mail-type=<mode>
Job name	-N <jobname>	--job-name=<jobname> -J <jobname>
Interactive jobs	-I	None (use salloc or srun)
Job dependencies	-W depend=<mode>:<jobID>	--dependency=<dependency_list> -d <dependency_list>

7 Known issues

The *psslurm* plugin is currently under active development and it is projected that a first release version will be available by the end of the year 2014. The Parastation consortium members monitor the progress of the development using an extensive regression test suite. Based on this test-suite the following list of known issues has been compiled. Users which observe problems or inconsistencies between the batch system on JUROPATEST and other Slurm installations are kindly asked to report their findings after checking with the list below whether the problem is already known.

The link [here](#) presents the updated list of all known issues for the current installed version of the psslurm plugin. The URL of the web page is:

<http://www.fz-juelich.de/ias/jsc/EN/Expertise/Supercomputers/JUROPATEST/UserInfo/Batch.html>

8 Examples

8.1 Template job-scripts

Template MPI job-script:

```
#!/bin/bash
#SBATCH -J <jobname>
#SBATCH -N <number>
#SBATCH -n <number> # can be omitted
#SBATCH --ntasks-per-node=<number>
#SBATCH -o <jobname>-%j.out
#SBATCH -e <jobname>-%j.err
#SBATCH --mail-type=<BEGIN, END, FAIL, or ALL>
#SBATCH --mail-user=<email>
#SBATCH --partition=<batch | large>
#SBATCH --time=<time>

# run MPI application below (with srun)
```

Template Hybrid job-script:

```
#!/bin/bash
#SBATCH -J <jobname>
#SBATCH -N <number>
#SBATCH -n <number> # can be omitted
#SBATCH --ntasks-per-node=<number>
#SBATCH --cpus-per-task=<number>
#SBATCH -o <jobname>-%j.out
#SBATCH -e <jobname>-%j.err
#SBATCH --mail-type=<BEGIN, END, FAIL, or ALL>
#SBATCH --mail-user=<email>
#SBATCH --partition=<batch | large>
#SBATCH --time=<time>

export OMP_NUM_THREADS=${SLURM_CPUS_PER_TASK}

# run Hybrid application below (with srun)
```

8.2 Modules

Check loaded modules:

```
$ module list
No modules loaded
```

Check available Toolchains:

```
$ module avail

----- /usr/local/software/juopatest/TC/FullToolchains -----
gpsolf/2014.11      intel/2014.11      intel-para/2014.11

----- /usr/local/software/juopatest/TC/Compilers+MPI -----
gpsmpi/2014.11     iimpi/7.1.2        ipsmpi/2014.11-mt  ipsmpi/2014.11 (D)

----- /usr/local/software/juopatest/TC/Compilers -----
GCC/4.9.1         Java/1.7.0_71       icc/2015.0.090     iccifort/2015.0.090  ifort/2015.0.090

----- /usr/local/software/juopatest/Stage1/modules/tools/Core -----
AllineaPerformanceReports/4.2-PR-39422  Inspector/2015      (D)      binutils/2.24
```

```
EasyBuild/1.15.2          VTune/2015_update1
Inspector/2015_update1    VTune/2015          (D)
```

```
----- /usr/local/software/juopatest/Devel -----
software_devel (S)
```

Where:

```
(S): Module is Sticky, requires --force to unload or purge
(D): Default Module
```

Use "module spider" to find all possible modules.

Use "module keyword key1 key2 ..." to search for all possible modules matching any of the "keys".

Load a Toolchain and check loaded modules:

```
$ module load intel-para/2014.11
$ module list
```

Currently Loaded Modules:

```
1) binutils/2.24      4) popt/1.14      7) iccifort/2015.0.090
2) icc/2015.0.090    5) pscom/5.0.44-1 8) imkl/11.2.0.090
3) ifort/2015.0.090 6) psmpi/5.1.0-1  9) intel-para/2014.11
```

Check available packages:

```
$ module avail
```

```
----- /usr/local/software/juopatest/Stage1/modules/all/MPI/intel/2015.0.090/psmpi/5.1.0-1 -----
ABINIT/7.8.2          Qt/4.8.4
ASE/3.6.0.2515-Python-2.7.3 Qt/4.8.5          (D)
ASE/3.8.0.3420-Python-2.7.3 (D) QuantumESPRESSO/5.1
Autoconf/2.69         SCOTCH/5.1.12b_esmumps
Automake/1.13.4       SCOTCH/6.0.0_esmumps (D)
BioPerl/1.6.1-Perl-5.20.0 SIONlib/1.5.2
Bison/2.6.5           Scalasca/2.1
Bison/2.7             ScientificPython/2.8-Python-2.7.3
Bison/3.0.2          (D) Score-P/1.2.3
Boost/1.49.0-Python-2.7.3 SuiteSparse/3.7.0-withpammetis
Boost/1.53.0          Szzip/2.1
Boost/1.56.0          (D) Tcl/8.5.16
CMake/2.8.4           UDUNITS/2.1.24
CMake/3.0.0           (D) UltraScan3/3.3.1868
Cube/4.2.3            VampirTrace/5.14.4
Doxygen/1.8.2         XML-LibXML/2.0018-Perl-5.20.0
Doxygen/1.8.7         (D) arpack-ng/3.1.3
ELPA/2014.06-generic-simple bzip2/1.0.4
FFTW/3.3.1            bzip2/1.0.5
FFTW/3.3.4           (D) bzip2/1.0.6          (D)
FIAT/1.0.0-Python-2.7.3 cURL/7.37.1
GDB/7.8              flex/2.5.37
GLib/2.34.3          flex/2.5.39        (D)
GPAW/0.10.0.11364-Python-2.7.3 freetype/2.5.2
GSL/1.15             gettext/0.18.2     (D)
GSL/1.16            (D) imkl/11.2.0.090
HDF5/1.8.10-gpfs     inputproto/2.3
HDF5/1.8.10          kbproto/1.0.6
HDF5/1.8.12          libICE/1.0.8
HDF5/1.8.13         (D) libSM/1.2.1
HPL/2.1              libX11/1.6.1
Harminv/1.3.1        libXaw/1.0.12
Hypre/2.8.0b         libXmu/1.1.2
IOR/2.10.3-mpiio     libXpm/3.5.11
JasPer/1.900.1       libXt/1.1.4
LWM2/1.0             libffi/3.0.13      (D)
Libint/1.1.4         libjpeg-turbo/1.3.1
LinkTest/1.1p5       libpng/1.6.12
M4/1.4.16            libreadline/6.2
M4/1.4.17           (D) libreadline/6.3     (D)
METIS/5.0.2          libtool/2.4.2
```

```

MUMPS/4.10.0-parmetis          libunistring/0.9.3
MethPipe/3.0.1                  libxc/2.0.1
NASM/2.07                        libxc/2.0.2 (D)
NASM/2.11.05                     (D) libxcb/1.8-Python-2.7.3
OPARI2/1.1.2                     libxml2/2.9.2-Python-3.4.1
OTF/1.12.5                       (D) libxml2/2.9.2 (D)
OTF2/1.2.1                       ncurses/5.9
OTF2/1.4                         (D) ncview/2.1.1
OpenFOAM/2.3.0                   ncview/2.1.2
OpenSSL/1.0.1i                   ncview/2.1.3 (D)
PAPI/5.2.0                       netCDF/4.2.1.1
PAPI/5.3.2                       (D) netCDF-Fortran/4.2
PDT/3.19                         (D) pkg-config/0.27.1
PETSc/3.3-p2-Python-2.7.3       tcsh/6.18.01
ParMETIS/3.2.0                  xcb-proto/1.7-Python-2.7.3
ParMETIS/4.0.2                  (D) xextproto/7.2.1
Perl/5.20.0                      xproto/7.0.23
Python/2.7.3                     xtrans/1.2
Python/2.7.5                     zlib/1.2.7
Python/2.7.8                     (D) zlib/1.2.8
Python/3.4.1                     (D)

----- /usr/local/software/juopatest/Stage1/modules/all/Compiler/intel/2015.0.090 -----
OTF/1.12.5    PDT/3.19    impi/5.0.1.035    popt/1.14    psmpi/5.1.0-1-mt
PAPI/5.3.2    gettext/0.18.2    libffi/3.0.13    pscom/5.0.44-1    psmpi/5.1.0-1 (D)

----- /usr/local/software/juopatest/TC/FullToolchains -----
gpsolf/2014.11    intel/2014.11    intel-para/2014.11

----- /usr/local/software/juopatest/TC/Compilers+MPI -----
gpsmpi/2014.11    iimpi/7.1.2    ipsmpi/2014.11-mt    ipsmpi/2014.11 (D)

----- /usr/local/software/juopatest/TC/Compilers -----
GCC/4.9.1    Java/1.7.0_71    icc/2015.0.090    iccifort/2015.0.090    ifort/2015.0.090

----- /usr/local/software/juopatest/Stage1/modules/tools/Core -----
AllineaPerformanceReports/4.2-PR-39422    Inspector/2015 (D)    binutils/2.24
EasyBuild/1.15.2    VTune/2015_update1
Inspector/2015_update1    VTune/2015 (D)

----- /usr/local/software/juopatest/Devel -----
software_devel (S)

Where:
(S): Module is Sticky, requires --force to unload or purge
(D): Default Module

Use "module spider" to find all possible modules.
Use "module keyword key1 key2 ..." to search for all possible modules matching any of the "keys".

```

Load a module:

```

$ module load OpenFOAM/2.3.0
$ module list

Currently Loaded Modules:
 1) binutils/2.24      5) pscom/5.0.44-1      9) intel-para/2014.11  13) OpenFOAM/2.3.0
 2) icc/2015.0.090    6) psmpi/5.1.0-1      10) libreadline/6.2
 3) ifort/2015.0.090  7) iccifort/2015.0.090 11) SCOTCH/6.0.0_esmumps
 4) popt/1.14         8) imkl/11.2.0.090    12) ncurses/5.9

```

Purge all modules:

```

$ module purge
$ module list
No modules loaded

```

Check a package:

```
$ module spider Boost
```

```
-----  
Boost:
```

```
-----  
Description:
```

```
Boost provides free peer-reviewed portable C++ source libraries. - Homepage:  
http://www.boost.org/
```

```
-----  
Versions:
```

```
Boost/1.49.0-Python-2.7.3  
Boost/1.53.0  
Boost/1.56.0
```

```
-----  
To find detailed information about Boost please enter the full name.  
For example:
```

```
$ module spider Boost/1.56.0  
-----
```

Check a specific version of a package:

```
$ module spider Boost/1.56.0
```

```
-----  
Boost: Boost/1.56.0
```

```
-----  
Description:
```

```
Boost provides free peer-reviewed portable C++ source libraries. - Homepage:  
http://www.boost.org/
```

```
This module can only be loaded through the following modules:
```

```
GCC/4.9.1, psmapi/5.1.0-1  
Stages/.software_devel_Stagel, GCC/4.9.1, psmapi/5.1.0-1  
Stages/.software_devel_Stagel, icc/2015.0.090, impi/5.0.1.035  
Stages/.software_devel_Stagel, icc/2015.0.090, psmapi/5.1.0-1  
Stages/.software_devel_Stagel, ifort/2015.0.090, impi/5.0.1.035  
Stages/.software_devel_Stagel, ifort/2015.0.090, psmapi/5.1.0-1  
icc/2015.0.090, impi/5.0.1.035  
icc/2015.0.090, psmapi/5.1.0-1  
ifort/2015.0.090, impi/5.0.1.035  
ifort/2015.0.090, psmapi/5.1.0-1  
software_devel, GCC/4.9.1, psmapi/5.1.0-1  
software_devel, icc/2015.0.090, impi/5.0.1.035  
software_devel, icc/2015.0.090, psmapi/5.1.0-1  
software_devel, ifort/2015.0.090, impi/5.0.1.035  
software_devel, ifort/2015.0.090, psmapi/5.1.0-1
```

8.3 Compilation

MPI program example (file *mpi.c*):

```
#include <stdio.h>  
#include <mpi.h>  
  
int main ( int argc, char** argv )  
{  
  
    int rank, size;  
    char processor_name [MPI_MAX_PROCESSOR_NAME];  
    int name_len;
```

```

// Initialize the MPI environment.
MPI_Init( &argc, &argv );

// Get the number of processes.
MPI_Comm_size ( MPI_COMM_WORLD, &size);

// Get the rank of the process.
MPI_Comm_rank ( MPI_COMM_WORLD, &rank );

// Get the name of the processor.
MPI_Get_processor_name ( processor_name, &name_len );

// Print out.
printf( "Hello world from processor %s, rank %d out of %d processors.\n", processor_name,
rank, size);

// Finalize the MPI environment.
MPI_Finalize();

return 10;
}

```

Hybrid program example (file *hybrid.c*):

```

#include <stdio.h>
#include <mpi.h>
#include "mpi.h"

#define _NUM_THREADS 16

int main ( int argc, char** argv )
{
    int rank, size, count, total;
    char processor_name [MPI_MAX_PROCESSOR_NAME];
    int name_len;

//    omp_set_num_threads(_NUM_THREADS);

// Initialize the MPI environment.
MPI_Init( &argc, &argv );

// Get the number of processes.
MPI_Comm_size ( MPI_COMM_WORLD, &size);

// Get the rank of the process.
MPI_Comm_rank ( MPI_COMM_WORLD, &rank );

// Get the name of the processor.
MPI_Get_processor_name ( processor_name, &name_len );

count = 0;

#pragma omp parallel reduction(+:count)
{
    count = count + omp_get_num_threads();
    total = omp_get_num_threads();
}

// Print out.
printf( "Hello world from processor %s, rank %d out of %d processors. OpenMP threads: %d\n",
processor_name, rank, size, total);

// Finalize the MPI environment.
MPI_Finalize();

return 0;
}

```

Compile the MPI program:

```
$ mpicc -o mpi-prog mpi.c
```

Compile the Hybrid program:

```
$ mpicc -openmp -o hybrid-prog hybrid.c
```

8.4 Job submission

Job-script for an MPI job (file *mpiscript.sh*):

```
#!/bin/bash
#SBATCH -J mpitest
#SBATCH -N 4
#SBATCH --ntasks-per-node=28
#SBATCH -o mpitest-%j.out
#SBATCH -e mpitest-%j.err
#SBATCH --mail-type=END
#SBATCH --mail-user=c.paschoulas@fz-juelich.de
#SBATCH --partition=large
#SBATCH --time=00:30:00

# run MPI application below (with srun)
srun -N 4 --ntasks-per-node=28 ./mpi-prog
```

Submit the MPI job-script:

```
$ sbatch ./mpiscript.sh
```

Job-script for a Hybrid job (file *hybridtest.sh*):

```
#!/bin/bash
#SBATCH -J hybridtest
#SBATCH -N 4
#SBATCH --ntasks-per-node=28
#SBATCH --cpus-per-task=2
#SBATCH -o hybridtest-%j.out
#SBATCH -e hybridtest-%j.err
#SBATCH --mail-type=END
#SBATCH --mail-user=c.paschoulas@fz-juelich.de
#SBATCH --partition=batch
#SBATCH --time=00:30:00

export OMP_NUM_THREADS=${SLURM_CPUS_PER_TASK}

# run Hybrid application below (with srun)
srun -N 4 --ntasks-per-node=28 -c ${SLURM_CPUS_PER_TASK} ./hybrid-prog
```

Submit the Hybrid job-script

```
$ sbatch ./hybridscrip.sh
```

8.5 Job Control

Hold a job:

```
$ scontrol hold 14900
$ squeue
```

JOBID	PARTITION	NAME	USER	ST	TIME	NODES	NODELIST(REASON)
14896	batch	hybridte	paschoul	CG	0:08	4	j3c[065-068]
14898	batch	hybridte	paschoul	PD	0:00	4	(QOSResourceLimit)
14899	batch	hybridte	paschoul	PD	0:00	4	(QOSResourceLimit)
14901	batch	hybridte	paschoul	PD	0:00	4	(QOSResourceLimit)
14900	batch	hybridte	paschoul	PD	0:00	4	(JobHeldUser)
14894	batch	hybridte	paschoul	R	0:08	4	j3c[120-123]
14895	batch	hybridte	paschoul	R	0:08	4	j3c[061-064]

Release a job:

```
$ scontrol release 14900
$ squeue
```

JOBID	PARTITION	NAME	USER	ST	TIME	NODES	NODELIST(REASON)
14900	batch	hybridte	paschoul	R	0:01	4	j3c[120-123]
14800	large	job	esmil702	PD	0:00	32	(PartitionDown)

Cancel a job:

```
$ scancel 14905
```

8.6 Query Commands

Check the Queue

```
$ squeue
```

JOBID	PARTITION	NAME	USER	ST	TIME	NODES	NODELIST(REASON)
14905	batch	hybridte	paschoul	CG	0:05	4	j3c[099-102]
14902	batch	hybridte	paschoul	CG	0:07	4	j3c[120-123]
14903	batch	hybridte	paschoul	CG	0:08	4	j3c[091-094]
14904	batch	hybridte	paschoul	CG	0:08	4	j3c[095-098]
14800	large	job	esmil702	PD	0:00	32	(PartitionDown)

Check the Queue for one user:

```
$ squeue -u paschoul
```

JOBID	PARTITION	NAME	USER	ST	TIME	NODES	NODELIST(REASON)
14910	batch	mpitest	paschoul	PD	0:00	4	(QOSResourceLimit)
14911	batch	mpitest	paschoul	PD	0:00	4	(QOSResourceLimit)
14912	batch	hybridte	paschoul	R	0:02	4	j3c[120-123]
14913	batch	hybridte	paschoul	R	0:02	4	j3c[091-094]
14908	batch	mpitest	paschoul	R	0:02	4	j3c[095-098]
14909	batch	mpitest	paschoul	R	0:02	4	j3c[099-102]

Check partitions and nodes:

```
$ sinfo
```

PARTITION	AVAIL	TIMELIMIT	NODES	STATE	NODELIST
batch*	up	2:00:00	5	drain	j3c[090,115-116,119,124]
batch*	up	2:00:00	65	idle	j3c[061-089,091-114,117-118,120-123,125-130]
large	down	1:00:00	5	drain	j3c[090,115-116,119,124]
large	down	1:00:00	65	idle	j3c[061-089,091-114,117-118,120-123,125-130]

Check off-line nodes:

```
$ sinfo -R
REASON          USER      TIMESTAMP      NODELIST
#523 - ipc objects l root    2014-11-17T09:47:55 j3c116
#494 - MCE-Errors: p root    2014-11-14T13:49:10 j3c090
#495 - MCE-Errors: p root    2014-11-14T13:49:49 j3c115
#496 - MCE-Errors: v root    2014-11-14T13:48:34 j3c119
#497 - MCE-Errors: p root    2014-11-14T13:53:22 j3c124
#494 - MCE-Errors: p root    2014-11-14T13:49:10 j3c090
#495 - MCE-Errors: p root    2014-11-14T13:49:49 j3c115
#494 - MCE-Errors: p root    2014-11-14T13:49:10 j3c090
```

Check reservations:

```
$ sinfo -T
RESV_NAME      STATE      START TIME      END TIME      DURATION      NODELIST
test           ACTIVE    2014-11-14T15:24:47  2015-10-01T00:00:00  320-07:35:13  j3c128
```

Check one partition:

```
$ scontrol show partition batch
PartitionName=batch
  AllowGroups=ALL AllowAccounts=ALL AllowQos=ALL
  AllocNodes=j3l03,j3l04 Default=YES
  DefaultTime=00:30:00 DisableRootJobs=YES GraceTime=0 Hidden=NO
  MaxNodes=4 MaxTime=02:00:00 MinNodes=1 LLN=NO MaxCPUsPerNode=56
  Nodes=j3c[061-130]
  Priority=1 RootOnly=NO ReqResv=NO Shared=NO PreemptMode=OFF
  State=UP TotalCPUs=3920 TotalNodes=70 SelectTypeParameters=N/A
  DefMemPerNode=125952 MaxMemPerNode=125952
```

Check one node:

```
$ scontrol show node j3c130
NodeName=j3c130 Arch=x86_64 CoresPerSocket=14
  CPUAlloc=0 CPUErr=0 CPUTot=56 CPULoad=0.01 Features=normal
  Gres=(null)
  NodeAddr=j3c130 NodeHostName=j3c130 Version=5.0.12
  OS=Linux RealMemory=128952 AllocMem=0 Sockets=2 Boards=1
  State=IDLE ThreadsPerCore=2 TmpDisk=0 Weight=1
  BootTime=2014-10-15T14:22:58 SlurmdStartTime=2014-11-17T09:04:37
  CurrentWatts=0 LowestJoules=0 ConsumedJoules=0
  ExtSensorsJoules=n/s ExtSensorsWatts=0 ExtSensorsTemp=n/s
```

Check the shares:

```
$ sshare
Account      User Raw Shares Norm Shares      Raw Usage Effectv Usage      FairShare
-----
zam          paschoul      3000      0.000037      317994      0.002611      0.000000
```

Check the priorities:

```
$ sprio
JOBID      PRIORITY      AGE      FAIRSHARE      JOBSIZE      QOS
15979      2882          2727          0          155          0
16004      2881          2726          0          155          0
16005      2933          2726          0          207          0
16008      2881          2726          0          155          0
16013      2881          2726          0          155          0
16017      2933          2726          0          207          0
```


8.7 Accounting Commands

Check user association:

```
$ sacctmgr show assoc where user=paschoul
```

Cluster	Account	User	Partition	Share	GrpJobs	GrpNodes	GrpCPUs	GrpMem	GrpSubmit
GrpWall	GrpCPUMins	MaxJobs	MaxNodes	MaxCPUs	MaxSubmit	MaxWall	MaxCPUMins		QOS
Def	QOS	GrpCPURunMins							
juropatest			zam	paschoul					3000
normal	normal								

Check all QoSs:

```
$ sacctmgr show qos
```

Name	Priority	GraceTime	Preempt	PreemptMode	Flags			
UsageThres	UsageFactor	GrpCPUs	GrpCPUMins	GrpCPURunMins	GrpJobs	GrpMem	GrpNodes	GrpSubmit
GrpWall	MaxCPUs	MaxCPUMins	MaxNodes	MaxWall	MaxCPUsPU	MaxJobsPU	MaxNodesPU	MaxSubmitPU
normal	100000	00:00:00		cluster	DenyOnLimit			
1.000000								
4	70	12						
lowcont	50000	00:00:00		cluster	DenyOnLimit			
1.000000								
4	70	12						
nocont	0	00:00:00		cluster	DenyOnLimit			
1.000000								
4	70	12						
suspended	0	00:00:00		cluster	DenyOnLimit			
1.000000								
0	00:00:00	0	0	0				
nolimits	100000	00:00:00		cluster	DenyOnLimit			
1.000000								

Check one QoS:

```
$ sacctmgr show qos where name=normal
```

Name	Priority	GraceTime	Preempt	PreemptMode	Flags			
UsageThres	UsageFactor	GrpCPUs	GrpCPUMins	GrpCPURunMins	GrpJobs	GrpMem	GrpNodes	GrpSubmit
GrpWall	MaxCPUs	MaxCPUMins	MaxNodes	MaxWall	MaxCPUsPU	MaxJobsPU	MaxNodesPU	MaxSubmitPU
normal	100000	00:00:00		cluster	DenyOnLimit			
1.000000								
4	70	12						

Check old jobs history:

```
$ sacct -X -u paschoul
```

JobID	JobName	Partition	Account	AllocCPUS	State	ExitCode
14797	TestJob	batch	zam	56	COMPLETED	0:0
14798	sleepscri+	batch	zam	56	COMPLETED	0:0
14799	sleepscri+	batch	zam	56	COMPLETED	0:0
14801	TestJob	batch	zam	224	FAILED	127:0
14802	TestJob	batch	zam	224	FAILED	127:0
14803	TestJob	batch	zam	224	FAILED	127:0
14804	TestJob	batch	zam	224	FAILED	127:0
14805	TestJob	batch	zam	224	FAILED	127:0

14806	sleepscri+	batch	zam	56	COMPLETED	0:0
14807	sleepscri+	batch	zam	56	COMPLETED	0:0
14814	sleepscri+	batch	zam	56	COMPLETED	0:0
14816	TestJob	batch	zam	224	FAILED	127:0
14817	TestJob	batch	zam	224	FAILED	127:0
14818	TestJob	batch	zam	224	FAILED	127:0
14820	sleepscri+	batch	zam	56	COMPLETED	0:0
14821	sleepscri+	batch	zam	56	COMPLETED	0:0
14881	hybridtest	batch	zam	224	COMPLETED	0:0
14882	hybridtest	batch	zam	224	COMPLETED	0:0
14883	hybridtest	batch	zam	224	COMPLETED	0:0
14884	mpitest	batch	zam	224	FAILED	10:0
14885	mpitest	batch	zam	224	CANCELLED+	0:0
14886	mpitest	batch	zam	224	FAILED	10:0
14887	mpitest	batch	zam	224	FAILED	10:0
14888	hybridtest	batch	zam	224	COMPLETED	0:0
14889	hybridtest	batch	zam	224	COMPLETED	0:0
14890	hybridtest	batch	zam	224	COMPLETED	0:0
14891	hybridtest	batch	zam	224	COMPLETED	0:0
14892	hybridtest	batch	zam	224	COMPLETED	0:0
14893	hybridtest	batch	zam	224	COMPLETED	0:0
14894	hybridtest	batch	zam	224	COMPLETED	0:0
14895	hybridtest	batch	zam	224	COMPLETED	0:0
14896	hybridtest	batch	zam	224	COMPLETED	0:0
14897	hybridtest	batch	zam	224	COMPLETED	0:0
...						

Check old jobs with different format and specified time frame:

```
$ sacct -X -u paschoul --format="jobid,user,nnodes,nodelist,state,exit" -S 2014-11-15T00:00:00 -E 2014-11-17T18:00:00
```

JobID	User	NNodes	NodeList	State	ExitCode
14797	paschoul	1	j3c117	COMPLETED	0:0
14798	paschoul	1	j3c118	COMPLETED	0:0
14799	paschoul	1	j3c117	COMPLETED	0:0
14801	paschoul	4	j3c[120-123]	FAILED	127:0
14802	paschoul	4	j3c[091-094]	FAILED	127:0
14803	paschoul	4	j3c[095-098]	FAILED	127:0
14804	paschoul	4	j3c[091-094]	FAILED	127:0
14805	paschoul	4	j3c[061-064]	FAILED	127:0
14806	paschoul	1	j3c129	COMPLETED	0:0
14807	paschoul	1	j3c130	COMPLETED	0:0
14814	paschoul	1	j3c117	COMPLETED	0:0
14816	paschoul	4	j3c[093-096]	FAILED	127:0
14817	paschoul	4	j3c[093-096]	FAILED	127:0
14818	paschoul	4	j3c[093-096]	FAILED	127:0
14820	paschoul	1	j3c127	COMPLETED	0:0
14821	paschoul	1	j3c129	COMPLETED	0:0
14823	paschoul	1	j3c130	COMPLETED	0:0
14824	paschoul	4	j3c[093-096]	FAILED	127:0
14880	paschoul	4	j3c[120-123]	FAILED	10:0
14881	paschoul	4	j3c[120-123]	COMPLETED	0:0
14882	paschoul	4	j3c[120-123]	COMPLETED	0:0
14883	paschoul	4	j3c[120-123]	COMPLETED	0:0
14884	paschoul	4	j3c[120-123]	FAILED	10:0
14885	paschoul	4	j3c[061-064]	CANCELLED+	0:0
14886	paschoul	4	j3c[065-068]	FAILED	10:0
14887	paschoul	4	j3c[077-080]	FAILED	10:0
14888	paschoul	4	j3c[120-123]	COMPLETED	0:0
14889	paschoul	4	j3c[061-064]	COMPLETED	0:0
14890	paschoul	4	j3c[065-068]	COMPLETED	0:0
14891	paschoul	4	j3c[077-080]	COMPLETED	0:0
14892	paschoul	4	j3c[120-123]	COMPLETED	0:0
14893	paschoul	4	j3c[061-064]	COMPLETED	0:0
14894	paschoul	4	j3c[120-123]	COMPLETED	0:0
14895	paschoul	4	j3c[061-064]	COMPLETED	0:0
14896	paschoul	4	j3c[065-068]	COMPLETED	0:0
14897	paschoul	4	j3c[077-080]	COMPLETED	0:0
14898	paschoul	4	j3c[120-123]	COMPLETED	0:0
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