Batch Usage on JURECA
Introduction to Slurm

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Batch System Concepts

**Resource Manager** is the software responsible for managing the resources of a cluster, usually controlled by a scheduler.

- It manages resources like tasks, nodes, CPUs, memory, network, etc.
- It handles the execution of the jobs on the compute nodes.
- It makes sure that jobs are not overlapping on the resources.

**Scheduler** is the software that controls user's jobs on a cluster according to policies. It receives and handles jobs from the users and controls the resource manager. It offers many features like:

- Scheduling mechanisms (backfill, fifo, etc)
- Partitions, queues and QoS to control jobs according to policies/limits
- Interfaces for defining work-flows (jobscript) or job dependencies and commands for managing the jobs (submit, cancel, etc)

**Batch-System/Workload-Manager** is the combination of a scheduler and a resource manager. It combines all the features of these two parts in an efficient way.
JSC Batch Model

- **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. Currently each user (uid) 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.

- Accounted CPU-Quotas/job = Number-of-nodes x Walltime (x cores/node)

- Contingent/CPU-Quota states for the projects (for monthly mode): normal, low-contingent, no-contingent.

- Contingent priorities: *normal* > *lowcont* > *nocont*. Users without contingent get some penalties for their jobs, but they are still allowed to submit and run jobs.
Slurm Introduction (1)

- **Slurm** is the chosen Batch System (Workload Manager) that is used on JURECA. Slurm is an open-source project developed by SchedMD. For our clusters *psslurm*, which is a plug-in of *psid* daemon and part of the Parastation Cluster tools, is replacing *slurmd* on the compute nodes. *psslurm* is under development by ParTec and JSC in the context of our collaboration.

- Slurm's configuration on JURECA:
  - High-availability for the main daemons slurmd and slurmdbd.
  - Backfilling scheduling algorithm.
  - No node-sharing.
  - Job scheduling according to priorities.
  - Accounting mechanism: slurmdbd with MySQL/MariaDB database.
  - User and job limits configured by QoS and Partitions.
  - No preemption configured. Running jobs cannot be preempted.
  - Prologue and Epilogue, with pshealthcheck from Parastation.
  - Generic resources (GRES) for different types of resources on the nodes.
Slurm groups the compute nodes into Partitions (similar to queues from Moab). Some limits and policies can be configured for each Partition:

- allowed users, groups or accounts
- max. nodes and max. wall-time limit per job

Other limits are enforced also by the Quality-of-Services (QoS), according to the contingent of user's group, e.g. max. wall-time limit, max number or queued or running jobs per user, etc...

Default limits/settings are used when not given by the users, like: number of nodes, number of tasks per node, wall-time limit, etc.

According to group's contingent user jobs are given certain QoS:

- **normal**: group has contingent, high job priorities.
- **lowcont**: this months contingent was used.
  penalty -> lower job priorities, max. wall-time limit and max. running jobs
- **nocont**: all contingent of the 3 months time-frame was used.
  penalty -> lowest job priorities, lower max. wall-time limit and max. jobs
- **suspended**: the group's project has ended; user cannot submit jobs
# JURECA Partitions

<table>
<thead>
<tr>
<th>Partition</th>
<th>Nodes</th>
<th>Resources</th>
<th>Walltime</th>
<th>MaxNodes</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>batch</code></td>
<td>1712</td>
<td>24(48) CPU Cores 128/256 GB RAM</td>
<td>Default: 1 hour Max.: 24 hours</td>
<td>256</td>
<td>Default partition, normal compute nodes (some with 256GB RAM)</td>
</tr>
<tr>
<td><code>devel</code></td>
<td>20</td>
<td>24(48) CPU Cores 128 GB RAM</td>
<td>Default: 30 mins Max.: 2 hours</td>
<td>8</td>
<td>Partition mainly for development (interactive jobs)</td>
</tr>
<tr>
<td><code>large</code></td>
<td>1712</td>
<td>24(48) CPU Cores 128/256 GB RAM</td>
<td>Default: 1 hour Max.: 24 hours</td>
<td>UNLIMITED</td>
<td>Same as batch targeting big jobs (currently down)</td>
</tr>
<tr>
<td><code>mem256</code></td>
<td>128</td>
<td>24(48) CPU Cores 256 GB RAM</td>
<td>Default: 1 hour Max.: 24 hours</td>
<td>128</td>
<td>Fat compute nodes with 256GB RAM (also in batch)</td>
</tr>
<tr>
<td><code>mem512</code></td>
<td>64</td>
<td>24(48) CPU Cores 512 GB RAM</td>
<td>Default: 1 hour Max.: 24 hours</td>
<td>32</td>
<td>Fat compute nodes with 512GB RAM</td>
</tr>
<tr>
<td><code>mem1024</code></td>
<td>2</td>
<td>24(48) CPU Cores 1 TB RAM</td>
<td>Default: 1 hour Max.: 24 hours</td>
<td>2</td>
<td>Fat compute nodes with 1TB RAM</td>
</tr>
<tr>
<td><code>gpus</code></td>
<td>70</td>
<td>24(48) CPU Cores 128 GB RAM 2x Nvidia K80</td>
<td>Default: 1 hour Max.: 24 hours</td>
<td>32</td>
<td>Compute nodes with 4 GPUs available (CUDA apps) (4 GPUs visible)</td>
</tr>
<tr>
<td><code>develgpus</code></td>
<td>5</td>
<td>24(48) CPU Cores 128 GB RAM 2x Nvidia K80</td>
<td>Default: 30 mins Max.: 2 hours</td>
<td>2</td>
<td>Partition for testing and development on the GPUs</td>
</tr>
<tr>
<td><code>vis</code></td>
<td>10</td>
<td>24(48) CPU Cores 512/1024 GB RAM 2x Nvidia K40</td>
<td>Default: 1 hour Max.: 24 hours</td>
<td>4</td>
<td>Compute nodes with 2 GPUs, mainly for visualization SW</td>
</tr>
</tbody>
</table>

* Other partitions: `largegpus`, `largevis` (same as `gpus` and `vis` but with no max. nodes limit)
## Generic Resources

- Slurm provides the functionality to define generic resources (GRES) for each node type. These generic resources can be used during job submissions in order to allocate nodes with specific resources or features. Users can request GRES with the “--gres” submission option.

- We have configured GRES for different resource types like memory or GPUs:
  - **Memory**: mem128, mem256, mem512, mem1024
  - **GPUs**: gpu:[1-4] (for visualization nodes: gpu:[1-2])

- The following table shows the list of GRES available on each partition:

<table>
<thead>
<tr>
<th>Partitions</th>
<th>GRES list</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>devel</code></td>
<td>mem128</td>
</tr>
<tr>
<td><code>batch ,large</code></td>
<td>mem128, mem256</td>
</tr>
<tr>
<td><code>mem256</code></td>
<td>mem256</td>
</tr>
<tr>
<td><code>mem512</code></td>
<td>mem512</td>
</tr>
<tr>
<td><code>gpus, largegpus</code></td>
<td>mem128+gpu:4</td>
</tr>
<tr>
<td><code>vis, largevis</code></td>
<td>mem512+gpu:2, mem1024+gpu:2</td>
</tr>
<tr>
<td><code>mem1024</code></td>
<td>mem1024+gpu:2</td>
</tr>
</tbody>
</table>
System Usage – Modules

- The installed software of the clusters 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.

- Preparing the module environment includes different steps:
  1. Load a compiler
  2. Load an MPI runtime (optional)
  3. Then load other application modules, which were built with currently loaded modules (compiler, MPI or other libraries)

- Useful commands:

<table>
<thead>
<tr>
<th>Task</th>
<th>Command</th>
</tr>
</thead>
<tbody>
<tr>
<td>List available modules</td>
<td><code>$ module avail</code></td>
</tr>
<tr>
<td>Load compiler and MPI</td>
<td><code>$ module load Intel ParaStationMPI</code></td>
</tr>
<tr>
<td>List all loaded modules</td>
<td><code>$ module list</code></td>
</tr>
<tr>
<td>List available modules</td>
<td><code>$ module avail</code></td>
</tr>
<tr>
<td>Check a package</td>
<td><code>$ module spider GROMACS</code></td>
</tr>
<tr>
<td>Load a module</td>
<td><code>$ module load GROMACS/&lt;version&gt;</code></td>
</tr>
<tr>
<td>Unload a module</td>
<td><code>$ module unload GROMACS/&lt;version&gt;</code></td>
</tr>
<tr>
<td>Unload all loaded modules</td>
<td><code>$ module purge</code></td>
</tr>
</tbody>
</table>
Slurm – User Commands (1)

- **salloc** is used to request interactive jobs/allocations.
- **sattach** is used to attach standard input, output, and error plus signal capabilities to a currently running job or job step.
- **sbatch** is used to submit a batch script (which can be a bash, Perl or Python script).
- **scancel** is used to cancel a pending or running job or job step.
- **sbcast** is used to transfer a file to all nodes allocated for a job.
- **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** 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.
- **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.
Slurm – User Commands (2)

- **sprio** can be used to query job priorities.
- **squeue** allows to query the list of pending and running jobs.
- **srun** is used to initiate *job-steps* mainly within a job or start an interactive jobs. 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 in Slurm's database.
- **sacctmgr** allows also the users to query some information about their accounts and other accounting information in Slurm's database.

*For more detailed info please check the online documentation and the man pages.*
There are 2 commands for job allocation: `sbatch` is used for batch jobs and `salloc` is used to allocate resource for interactive jobs. The format of these commands:

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

List of the most important submission/allocation options:

- `-A|--account` Charge CPU-Quota to specified account (budget ID).
- `-c|--cpus-per-task` Number of logical CPUs (hardware threads) per task.
- `-e|--error` Path to the job's standard error.
- `-i|--input` Connect the jobscript’s standard input directly to a file.
- `-J|--job-name` Set the name of the job.
- `--mail-user` Define the mail address for notifications.
- `--mail-type` When to send mail notifications. Options: BEGIN, END, FAIL, ALL
- `-N|--nodes` Number of compute nodes used by the job.
- `-n|--ntasks` Number of tasks (MPI processes).
- `--ntasks-per-node` Number of tasks per compute node.
- `-o|--output` Path to the job's standard output.
- `-p|--partition` Partition to be used from the job.
- `-t|--time` Maximum wall-clock time of the job.
- `--gres` Request nodes with specific Generic Resources.
Slurm - Submission filter and GRES

- Slurm is using a submission filter with the following functionality*:
  - Deny jobs requesting multiple partitions, we allow only one.
  - Disable the `--requeue` options. We do not allow users to requeue their jobs.
  - By default add the `memXXX` GRESs when missing, users can always specify the `memXXX` GRES if they want.
  - When a job is submitted in the partitions with GPUs then the submission is denied if no `gpu` GRES was specified.
  - Deny jobs with wrong `memXXX` GRESs, e.g. job submitted to `mem512` partition with GRES `mem128`.

* New policies starting from June 2017.

- Examples:
  - Submit a job in the `gpus` partition requesting 4 GPUs per node:
    ```bash
    sbatch -N 2 -p gpus --gres=gpu:4 <job-script>
    ```
  - Submit a job in the `mem512` partition:
    ```bash
    sbatch -N 4 -p mem512 --gres=mem512 <job-script>
    ```
Slurm – Spawning Command

- With **srun** the users can spawn any kind of application, process or task inside a job allocation. *srun* should be used either:
  1. Inside a job script submitted by `sbatch` (starts a job-step).
  2. After calling `salloc` (execute programs interactively).

- **Command format:**

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

- **srun** accepts almost all allocation options of `sbatch` and `salloc`. There are however some other unique options:

  ```bash
  --forward-x        Enable X11 forwarding only for interactive jobs.
  --pty              Execute a task in pseudo terminal mode.
  --multiprog <file> Run different programs with different arguments for each task specified in a text file.
  ```

- Note: In order to spawn the MPI applications, the users should always use *srun* and not *mpiexec.*
Instead of passing options to `sbatch` from the command-line, it is better to specify these options using the “`#SBATCH`” directives inside the job scripts which must be positioned in the very beginning of the job-script!

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.

```bash
#!/bin/bash

#SBATCH -J TestJob
#SBATCH -N 1
#SBATCH -o TestJob-%j.out
#SBATCH -e TestJob-%j.err
#SBATCH --time=30

sleep 5

hostname
```
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 4 nodes with 2 processes per node the system command `hostname`, requesting a walltime of 10 minutes. 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.

```
#!/bin/bash

#SBATCH -J TestJob
#SBATCH -N 4
#SBATCH -o TestJob-%j.out
#SBATCH -e TestJob-%j.err
#SBATCH --time=10

srun --ntasks-per-node=2 hostname
```
Job-Script – OpenMP Job

In this example the job will execute an OpenMP 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.

```bash
#!/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=48

/home/user/test/omp-prog
```
In the following example, an MPI application will start 96 tasks on 4 nodes running 24 tasks per node requesting a wall-time limit of 15 minutes in batch partition. Each MPI task will run on a separate core of the CPU. Users can change the modules also inside the jobscript.

```bash
#!/bin/bash

#SBATCH --nodes=4
#SBATCH --ntasks=96
#SBATCH --output=mpi-out.%j
#SBATCH --error=mpi-err.%j
#SBATCH --time=00:15:00
#SBATCH --partition=batch

module purge
module load Intel ParaStationMPI

srun ./mpi-prog   # implied --ntasks-per-node=24
```
In this example, a hybrid MPI/OpenMP job is presented. This job will allocate 5 compute nodes for 2 hours. The job will have 30 MPI tasks in total, 6 tasks per node and 4 OpenMP threads per task. On each node 24 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 of the option “--cpus-per-task/-c”.

```bash
#!/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=6 --cpus-per-task=4 ./hybrid-prog
```
The CPUs on our clusters support Simultaneous Multi-Threading (SMT). SMT is enabled by default for Slurm. In order to use SMT, the users must either allocate more than half of the Logical Cores on each Socket or by setting some specific CPU-Binding (Affinity) options.

This example presents a hybrid application which will execute “hybrid-prog” on 3 nodes using 2 MPI tasks per node and 24 OpenMP threads per task (48 CPUs per node).

```
#!/bin/bash

#SBATCH --ntasks=6
#SBATCH --ntasks-per-node=2
#SBATCH --cpus-per-task=24
#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
```
Slurm introduces the concept of **job-steps**. A job-step can be viewed as a smaller job or allocation inside the current allocation. Job-steps can be started only with the `srun` command.

The following example shows the usage of job-steps. With `sbatch` we allocate 32 compute nodes for 6 hours. Then we spawn 3 job-steps. The first step will run on 16 compute nodes for 50 minutes, the second step on 2 nodes for 10 minutes and the third step will use all 32 allocated nodes for 5 hours.

```bash
#!/bin/bash

#SBATCH -N 32
#SBATCH --time=06:00:00
#SBATCH --partition=batch

srun -N 16 -n 32 -t 00:50:00 ./mpi-prog1
srun -N 2 -n 4 -t 00:10:00 ./mpi-prog2
srun -N 32 --ntasks-per-node=2 -t 05:00:00 ./mpi-prog3
```
Job Dependencies & Job-Chains

- Slurm supports dependency chains which are collections of batch jobs with defined dependencies. Job dependencies can be defined using the "--dependency" or "-d" option of `sbatch`. The format is:

  `sbatch -d <type>:<jobID> <jobscript>

  Available dependency types: afterany, afternotok, afterok`

- Below is an example of a bash script for starting a chain of jobs. The script submits a chain of "$NO_OF_JOBS". Each job will start only after successful completion of its predecessor.

```bash
#!/bin/bash

NO_OF_JOBS=<no-of-jobs>
JOB_SCRIPT=<jobscript-name>

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

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

- Slurm supports job-arrays which can be defined using the option “-a” or “--array” of `sbatch` command. To address a job-array, Slurm provides a base array ID and an array index for each job. The syntax for specifying an array-job is: `--array=<range of indices>`

- Slurm exports also 2 env. variables that can be used in the job scripts:
  
  ```
  SLURM_ARRAY_JOB_ID : base array job ID
  SLURM_ARRAY_TASK_ID : array index
  ```

- Some additional options are available to specify the std-in/-out/-err file names in the job scripts: “%A” will be replaced by `SLURM_ARRAY_JOB_ID` and “%a” will be replaced by `SLURM_ARRAY_TASK_ID`.

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

srun -N 1 --ntasks-per-node=1  ./prog input_${SLURM_ARRAY_TASK_ID}.txt
```
CUDA-aware MPI Jobs

- JURECA offers the environment for CUDA-aware MPI Jobs. For this purpose certain modules are provided, like “MVAPICH2” which is built with PGI compiler.
- Following there is an example of a job-script that is running a CUDA MPI job with multiple tasks per node. 4 nodes will be allocated in the gpus partition, also 4 GPUs per node are requested:

```bash
#!/bin/bash
#SBATCH --nodes=4
#SBATCH --time=02:00:00
#SBATCH -p gpus
#SBATCH --gres=gpu:4
module purge
module load PGI/16.9-GCC-5.4.0 MVAPICH2/2.2-GDR
srun -N 4 --ntasks-per-node=24 ./cuda_aware_mpi
```

**Note:** For jobs with more than one task per node users can enable the CUDA MPS server on the GPU nodes with the submission option “--cuda-mps” which is expected to improve the performance (currently there are some issues with current versions of PGI compiler and MVAPICH2 runtime).
Interactive Jobs

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

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

After a successful allocation, `salloc` will start a shell on the login node where the submission happened. After the allocation the users can execute `srun` in order to spawn interactively their applications on the compute nodes. For example:

```
srun -N 4 --ntasks-per-node=2 -t 00:10:00 ./mpi-prog
```

The interactive session is terminated by exiting the shell. It is possible to obtain a remote shell on the compute nodes, after `salloc`, by running `srun` with the pseudo-terminal “--pty” option and a shell as argument:

```
srun --cpu_bind=none -N 2 --pty /bin/bash
```

It is also possible to start an interactive job and get a remote shell on the compute nodes with `srun` (not recommended without `salloc`):

```
srun --cpu_bind=none -N 1 -n 1 -t 01:00:00 --pty /bin/bash -i
```
Further Information

- Updated status of the systems:
  - See "Message of the day" at login.
- Get recent status updates by subscribing to the system high-messages:
  http://juelich.de/jsc/CompServ/services/high_msg.html
- JURECA online documentation:
  http://www.fz-juelich.de/ias/jsc/jureca
- User support at FZJ:
  - Email: sc@fz-juelich.de
  - Phone: +49 2461 61-2828
Questions?