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Information about the batch system (SLURM)

Please confer /etc/slurm/README.

The documentation of Slurm can be found ?here.

Overview

Slurm offers interactive and batch jobs (scripts submitted into the system). The relevant commands are srun and sbatch. The srun command can be used to spawn processes (**please do not use mpiexec**), both from the frontend and from within a batch script. You can also get a shell on a node to work locally there (e.g. to compile your application natively for a special platform).

Available Partitions

Please note that there is no default partition configured. In order to run a job, you have to specify one of the following partitions, using the --partition=... switch:

Name	Description
dp-cn	dp-cn[01-50], DEEP-EST Cluster nodes (Xeon Skylake)
dp-dam	dp-dam[01-16], DEEP-EST Dam nodes (Xeon Cascadelake + 1 V100 + 1 Stratix 10)
dp-dam-ext	dp-dam[09-16], DEEP-EST Dam nodes connected with Extoll Tourmalet
dp-esb	dp-esb[01-25], DEEP-EST Esb nodes (Xeon Cascadelake + 1 V100)
dp-sdv-esb	dp-sdv-esb[01-02], DEEP-EST ESB Test nodes (Xeon Cascadelake + 1 V100)
ml-gpu	ml-gpu[01-03], GPU test nodes for ML applications (up to 4 V100 cards)
sdv	deeper-sdv[01-16], cluster test nodes with Xeon Haswell CPU
extoll	deeper-sdv[01-16] (these nodes use an Extoll Tourmalet fabric)
knl	knl[01,04-06], KNL nodes
knl256	knl[01,05], KNL nodes with 64 cores
knl272	knl[04,06], KNL nodes with 68 cores
snc4	knl[05], KNL node in snc4 memory mode
psgw-cluster	gateway test node
psgw-booster	gateway test node
debug	all compute nodes (no gateways)

Anytime, you can list the state of the partitions with the sinfo command. The properties of a partition can be seen using

scontrol show partition <partition>

Remark about environment

By default, Slurm passes the environment from your job submission session directly to the execution environment. Please be aware of this when running jobs with srun or when submitting scripts with sbatch. This behavior can be controlled via the --export option. Please refer to the <u>?Slurm</u> <u>documentation</u> to get more information about this.

In particular, when submitting job scripts, it is recommended to load the necessary modules within the script and submit the script from a clean environment.

An introductory example

Suppose you have an mpi executable named hello_mpi. There are three ways to start the binary.

From a shell on a node

First, start a shell on a node. You would like to run your mpi task on 4 machines with 2 tasks per machine:

```
[kreutzl@deepv /p/project/cdeep/kreutz1/Temp]$ srun -A deep -p dp-cn -N 4 -n 8 -t 00:30:00 --pty /bin/bash -i
[kreutzl@dp-cn01 /p/project/cdeep/kreutz1/Temp]$
```

The environment is transported to the remote shell, no .profile, .bashrc, ... are sourced (especially not the modules default from /etc/profile.d/modules.sh). As of March 2020, an account has to be specified using the --account (short -A) option, which is "deep" for the project members. For people not included in the DEEP-EST project, please use the "Budget" name you received along with your account creation.

Once you get to the compute node, start your application using srun. Note that the number of tasks used is the same as specified in the initial srun command above (4 nodes with two tasks each):

```
[kreutzl@deepv Temp]$ srun -A deep -p dp-cn -N 4 -n 8 -t 00:30:00 --pty /bin/bash -i
[kreutzl@dp-cn01 Temp]$ srun ./MPI_HelloWorld
Hello World from rank 3 of 8 on dp-cn02
Hello World from rank 7 of 8 on dp-cn04
Hello World from rank 2 of 8 on dp-cn04
Hello World from rank 6 of 8 on dp-cn04
Hello World from rank 0 of 8 on dp-cn01
Hello World from rank 4 of 8 on dp-cn03
Hello World from rank 1 of 8 on dp-cn01
Hello World from rank 5 of 8 on dp-cn03
```

You can ignore potential warnings about the cpu binding. ParaStation will pin your processes.

Running directly from the front ends

You can run the application directly from the frontend, bypassing the shell:

```
[kreutzl@deepv Temp]$ srun -A deep -p dp-cn -N 4 -n 8 -t 00:30:00 ./MPI_HelloWorld
Hello World from rank 7 of 8 on dp-cn04
Hello World from rank 6 of 8 on dp-cn02
Hello World from rank 2 of 8 on dp-cn02
Hello World from rank 4 of 8 on dp-cn03
Hello World from rank 0 of 8 on dp-cn01
Hello World from rank 1 of 8 on dp-cn01
Hello World from rank 5 of 8 on dp-cn03
```

In this case, it can be useful to create an allocation which you can use for several runs of your job:

[kreutzl@deepv Temp]\$ salloc -A deep -p dp-cn -N 4 -n 8 -t 00:30:00 salloc: Granted job allocation 69263 [kreutzl@deepv Temp]\$ srun ./MPI_HelloWorld Hello World from rank 7 of 8 on dp-cn04 Hello World from rank 3 of 8 on dp-cn02 Hello World from rank 6 of 8 on dp-cn02 Hello World from rank 2 of 8 on dp-cn03 Hello World from rank 1 of 8 on dp-cn01 Hello World from rank 4 of 8 on dp-cn03 Hello World from rank 0 of 8 on dp-cn01 ... # several more runs

```
[kreutzl@deepv Temp]$ exit
exit
salloc: Relinquishing job allocation 69263
```

Batch script

Given the following script hello_cluster.sh:

```
#!/bin/bash
#SBATCH --partition=dp-cn
#SBATCH -A deep
#SBATCH -N 4
#SBATCH -n 8
#SBATCH -o /p/project/cdeep/kreutz1/hello_cluster-%j.out
#SBATCH -e /p/project/cdeep/kreutz1/hello_cluster-%j.err
#SBATCH --time=00:10:00
srun ./MPI_HelloWorld
```

This script requests 4 nodes with 8 tasks, specifies the stdout and stderr files, and asks for 10 minutes of walltime. Submit:

```
[kreutzl@deepv Temp]$ sbatch hello_cluster.sh
Submitted batch job 69264
```

Check what it's doing:

[kreutz1@deepv Te	mp]\$ squeue	-u \$USER					
JOBID	PARTITION	NAME	USER	ST	TIME	NODES	NODELIST(REASON)
69264	dp-cn ł	nello_cl	kreutz1	CG	0:04	4	dp-cn[01-04]

Check the result:

```
[kreutzl@deepv Temp]$ cat /p/project/cdeep/kreutzl/hello_cluster-69264.out
Hello World from rank 6 of 8 on dp-cn04
Hello World from rank 3 of 8 on dp-cn02
Hello World from rank 4 of 8 on dp-cn01
Hello World from rank 4 of 8 on dp-cn03
Hello World from rank 7 of 8 on dp-cn04
Hello World from rank 5 of 8 on dp-cn03
Hello World from rank 1 of 8 on dp-cn01
```

Heterogeneous jobs

As of version 17.11 of Slurm, heterogeneous jobs are supported. For example, the user can run:

```
srun --account=deep --partition=dp-cn -N 1 -n 1 hostname : --partition=dp-dam -N 1 -n 1 hostname
dp-cn01
dp-dam01
```

Please notice the : separating the definitions for each sub-job of the heterogeneous job. Also, please be aware that it is possible to have more than two sub-jobs in a heterogeneous job.

The user can also request several sets of nodes in a heterogeneous allocation using salloc. For example:

```
salloc --partition=dp-cn -N 2 : --partition=dp-dam -N 4
```

In order to submit a heterogeneous job via sbatch, the user needs to set the batch script similar to the following one:

#!/bin/bash #SBATCH -- job-name=imb_execute_1 #SBATCH --account=deep *#SBATCH --mail-user=* #SBATCH --mail-type=ALL #SBATCH --output=job.out #SBATCH --error=job.err #SBATCH --time=00:02:00 *#SBATCH --partition=dp-cn* #SBATCH --nodes=1 #SBATCH --ntasks=12 #SBATCH --ntasks-per-node=12 #SBATCH --cpus-per-task=1 #SBATCH packjob #SBATCH --partition=dp-dam *#SBATCH --constraint=* #SBATCH --nodes=1 #SBATCH --ntasks=12 #SBATCH --ntasks-per-node=12 #SBATCH --cpus-per-task=1 srun ./app_cn : ./app_dam

Here the packjob keyword allows to define Slurm parameters for each sub-job of the heterogeneous job. Some Slurm options can be defined once at the beginning of the script and are automatically propagated to all sub-jobs of the heterogeneous job, while some others (i.e. --nodes or --ntasks) must be defined for each sub-job. You can find a list of the propagated options on the ?Slurm documentation.

When submitting a heterogeneous job with this colon notation using ParaStationMPI, a unique MPI_COMM_WORLD is created, spanning across the two partitions. If this is not desired, one can use the --pack-group key to submit independent job steps to the different node-groups of a heterogeneous allocation:

srun --pack-group=0 ./app_cn ; srun --pack-group=1 ./app_dam

Using this configuration implies that inter-communication must be established manually by the applications during run time, if needed.

For more information about heterogeneous jobs please refer to the <u>?relevant page</u> of the Slurm documentation.

Heterogeneous jobs with MPI communication across modules

In order to establish MPI communication across modules using different interconnect technologies, some special Gateway nodes must be used. On the DEEP-EST system, MPI communication across gateways is needed only between Infiniband and Extoll interconnects.

Attention: Only ParaStation MPI supports MPI communication across gateway nodes.

This is an example job script for setting up an Intel MPI benchmark between a Cluster and a DAM node using a IB \leftrightarrow Extoll gateway for MPI communication:

#!/bin/bash
Script to launch IMB PingPong between DAM-CN using 1 Gateway
Use the gateway allocation provided by SLURM
Use the packjob feature to launch separately CM and DAM executable
General configuration of the job

```
#SBATCH --job-name=modular-imb
#SBATCH --account=deep
#SBATCH --time=00:10:00
#SBATCH --output=modular-imb-%j.out
#SBATCH --error=modular-imb-%j.err
# Configure the gateway daemon
#SBATCH --gw_num=1
#SBATCH --gw_psgwd_per_node=1
# Configure node and process count on the CM
#SBATCH --partition=dp-cn
#SBATCH --nodes=1
#SBATCH --ntasks-per-node=1
#SBATCH packjob
# Configure node and process count on the DAM
#SBATCH --partition=dp-dam-ext
#SBATCH --nodes=1
#SBATCH --ntasks-per-node=1
# Echo job configuration
echo "DEBUG: SLURM_JOB_NODELIST=$SLURM_JOB_NODELIST"
echo "DEBUG: SLURM_NNODES=$SLURM_NNODES"
echo "DEBUG: SLURM_TASKS_PER_NODE=$SLURM_TASKS_PER_NODE"
# Set the environment to use PS-MPI
module --force purge
module use $OTHERSTAGES
module load Stages/Devel-2019a
module load Intel
module load ParaStationMPI
# Show the hosts we are running on
srun hostname : hostname
# Execute
APP="./IMB-MPI1 Uniband"
```

srun \${APP} : \${APP}

Attention: During the first part of 2020, only the DAM nodes will have Extoll interconnect, while the CM and the ESB nodes will be connected via Infiniband. This will change later during the course of the project (expected Summer 2020), when the ESB will be equipped with Extoll connectivity (Infiniband will be removed from the ESB and left only for the CM).

A general description of how the user can request and use gateway nodes is provided at <u>?this section</u> of the JURECA documentation.

Attention: some information provided on the JURECA documentation do not apply for the DEEP system. In particular:

- as of 31/03/2020, the DEEP system has 2 gateway nodes.
- As of 09/01/2020 the gateway nodes are exclusive to the job requesting them. Given the limited number of gateway nodes available on the system, this may change in the future.
- As of 09/04/2020 the xenv utility (necessary on JURECA to load modules for different architectures Haswell and KNL) is not needed any more on DEEP when using the latest version of ParaStationMPI (currently available in the Devel-2019a stage and soon available on the default production stage).

Workflows

The new version of the installed slurm now supports workflows. The idea is to have an overlap between the dependent jobs so that they can communicate the data over the network instead of writing and reading it on storage. We have provided two ways to achieve a workflow. One way is to

use the new delay switch provided in sbatch command. While the other method is to submit jobs with dependencies of type afterok and later the independent job changes the dependency type of the dependent job to after using our provided shared library (explained below). Jacopo has developed an example project <u>?https://gitlab.version.fz-juelich.de/deamicis1/mpi_connect_test/-/tree/test_zia_workflows</u> that uses all the features discussed here.

The following simple example script helps understanding the mechanism of new delay switch for workflows.

```
[hudal@deepv scripts]$ cat test.sh
#!/bin/sh
NAME=$(hostname)
echo "$NAME: Going to sleep for $1 seconds"
sleep $1
echo "$NAME: Awake"
[hudal@deepv scripts]$ cat batch_workflow.sh
#!/bin/bash
#SBATCH -p sdv -N2 -t3
#SBATCH packjob
#SBATCH packjob
#SBATCH -p sdv -N1 -t3 --delay 2
srun test.sh 175
[hudal@deepv scripts]$
```

In the above sbatch script, the usage of --delay can be seen. It takes thee values in minutes. The idea is to delay the corresponding job of a heterogeneous job by the provided number of minutes from the start of the first job in the job pack. After submission of this job pack, slurm divides it into separate jobs at the time of the resource reservation. So you can see multiple jobs in the output of squeue command. Here is the example execution of this script.

```
[hudal@deepv scripts]$ sbatch batch_workflow.sh
Submitted batch job 81458
[hudal@deepv scripts]$ squeue -u hudal
JOBID PARTITION NAME USER ST TIME NODES NODELIST(REASON)
81458 sdv batch_wo hudal CF 0:01 2 deeper-sdv[02-03]
81459 sdv batch_wo hudal PD 0:00 1 (Reservation)
[hudal@deepv scripts]$
```

Here the second job(81458) will start 2 minutes after the start of the first job(81459). Similarly, the output files will be different for each separated job in the job pack. the final outputs are:

```
[hudal@deepv scripts]$ cat slurm-81458.out
deeper-sdv02: Going to sleep for 175 seconds
deeper-sdv03: Going to sleep for 175 seconds
deeper-sdv02: Awake
deeper-sdv03: Awake
[hudal@deepv scripts]$ cat slurm-81459.out
deeper-sdv01: Going to sleep for 175 seconds
deeper-sdv01: Awake
[hudal@deepv scripts]$
```

Another feature to note is that if there are multiple jobs in a job pack and any number of consecutive jobs have the same delay values, they are combined into a new heterogeneous job. Here is an example of such a script:

```
[huda1@deepv scripts]$ cat batch_workflow_complex.sh
#!/bin/bash
#SBATCH -p sdv -N 2 -t 3
#SBATCH -J first
#SBATCH packjob
#SBATCH -p sdv -N 1 -t 3 --delay 2
#SBATCH -J second
#SBATCH packjob
#SBATCH -p sdv -N 1 -t 2 --delay 2
#SBATCH -J second
#SBATCH packjob
#SBATCH -p sdv -N 2 -t 3 --delay 4
#SBATCH -J third
if [ "$SLURM_JOB_NAME" == "first" ]
then
       srun ./test.sh 150
elif [ "$SLURM_JOB_NAME" == "second" ]
then
       srun ./test.sh 150 : ./test.sh 115
elif [ "$SLURM_JOB_NAME" == "third" ]
then
       srun ./test.sh 155
fi
[hudal@deepv scripts]$
```

Note the delay values for the second and third job in the script are equal. Also, note the usage of the environment variable SLURM_JOB_NAME in the script to decide which command to run in which job. The example execution leads to the following:

```
[huda1@deepv scripts]$ sbatch batch_workflow_complex.sh
Submitted batch job 81460
[hudal@deepv scripts]$ squeue -u huda1
         JOBID PARTITION NAME
                                USER ST
                                            TIME NODES NODELIST(REASON)
        81461+0 sdv second hudal PD
                                            0:00
                                                  1 (Resources)
                  sdv second hudal PD
                                                    1 (Resources)
        81461+1
                                            0:00
                  sdv third hudal PD
         81463
                                            0:00
                                                    2 (Resources)
                  sdv first hudal PD
                                                    2 (Resources)
         81460
                                            0:00
[huda1@deepv scripts]$
```

Note that the submitted heterogeneous job has been divided into a single job (81460), a job pack (81461) and again a single job (81643). Similarly, three different output files will be generated, one for each new job.

```
[hudal@deepv scripts]$ cat slurm-81460.out
deeper-sdv03: Going to sleep for 150 seconds
deeper-sdv04: Going to sleep for 150 seconds
deeper-sdv03: Awake
deeper-sdv04: Awake
```

[hudal@deepv scripts]\$ cat slurm-81461.out deeper-sdv01: Going to sleep for 150 seconds deeper-sdv02: Going to sleep for 115 seconds deeper-sdv02: Awake deeper-sdv01: Awake [hudal@deepv scripts]\$ cat slurm-81463.out deeper-sdv01: Going to sleep for 155 seconds deeper-sdv02: Going to sleep for 155 seconds deeper-sdv01: Awake deeper-sdv02: Awake [hudal@deepv scripts]\$

If a job exits earlier than the allocated time asked by the user, the corresponding reservation for this job is deleted 5 minutes after the end of the job, automatically and the resources become available for the other jobs. However, users should be careful with the requested time when submitting workflows as the larger time values can delay the scheduling of the workflows depending on the situation of the resources.

The workflows created using delay switch ensure overlap between the applications. The second method that includes dependencies among jobs, does not ensure an overlap but avoids users to guess the time a job will take and how much should be the delay between jobs. The process is simple. A user submits a job and later a dependent job with a dependency of type afterok. Inside the first (independent) job, the application running calls the function provided in slurm_workflow library, that changes the dependency type of the dependent job to after. This enables the dependent job to be eligible for allocation by slurm immediately. However, the allocation of resources depends upon the situation of resources available in the system. The following script helps to submit jobs in the form of a chain with a provided dependency type.

```
[huda1@deepv scripts]$ cat chain_jobs.sh
#!/usr/bin/env bash
if [ $# -lt 3 ]
then
   echo "$0: ERROR (MISSING ARGUMENTS)"
   exit 1
fi
LOCKFILE=$1
DEPENDENCY_TYPE=$2
shift 2
SUBMITSCRIPT=$*
if [ -f $LOCKFILE ]
then
   if [[ "$DEPENDENCY_TYPE" =~ ^(after afterany afterok afternotok)$ ]]; then
       DEPEND_JOBID=`head -1 $LOCKFILE`
       echo "sbatch --dependency=${DEPENDENCY_TYPE}:${DEPEND_JOBID} $SUBMITSCRIPT"
       JOBID=`sbatch --dependency=${DEPENDENCY_TYPE}:${DEPEND_JOBID} $SUBMITSCRIPT`
   else
       echo "$0: ERROR (WRONG DEPENDENCY TYPE: choose among 'after', 'afterany', 'afterok' or 'afternotok')"
   fi
else
   echo "sbatch $SUBMITSCRIPT"
   JOBID=`sbatch $SUBMITSCRIPT`
fi
echo "RETURN: $JOBID"
# the JOBID is the last field of the output line
echo ${JOBID##* } > $LOCKFILE
exit 0
```

Here is the example of submission.

```
[hudal@deepv scripts]$ ./chain_jobs.sh lockfile afterok simple_job.sh
sbatch simple job.sh
RETURN: Submitted batch job 98626
[hudal@deepv scripts]$ ./chain_jobs.sh lockfile afterok simple_job.sh
sbatch --dependency=afterok:98626 simple_job.sh
RETURN: Submitted batch job 98627
[hudal@deepv scripts]$ ./chain_jobs.sh lockfile afterok simple_job.sh
sbatch --dependency=afterok:98627 simple_job.sh
RETURN: Submitted batch job 98628
[hudal@deepv scripts]$ squeue -u huda1
                                                    TIME NODES NODELIST(REASON)
           JOBID PARTITION
                              NAME
                                       USER ST
           98627
                      sdv simple_j hudal PD
                                                     0:00
                                                               2 (Dependency)
           98628
                      sdv simple_j hudal PD
                                                     0:00
                                                               2 (Dependency)
                                     hudal R
           98626
                      sdv simple j
                                                     0:21
                                                               2 deeper-sdv[01-02]
[hudal@deepv scripts]$ scontrol show job 98628 | grep Dependency
 JobState=PENDING Reason=Dependency Dependency=afterok:98627
[huda1@deepv scripts]$ cat lockfile
98628
```

Note that the lockfile contains the id of last submitted job.

slurm_workflow Library

We have developed a library that developers can use to change the reservation beginning times or dependency type of the dependent jobs in a workflow. This library is called slurm_workflow. The library has two functions.

The first function moves all the reservations of the remaining workflow jobs to an earlier time when the workflow is created using --delay switch.

```
/*
IN: number of minutes from now. The start time of the next reservation of the workflow is moved to this time if doable.
OUT: 0 successful, non zero unsuccessful.slurm_wf_error is set.
*/
int slurm_wf_move_all_res(uint32_t t);
```

The second function changes the dependencies type of all jobs dependent on the current job from afterok: job_id to after: job_id.

```
/*
OUT: 0 successful, error no otherwise.
*/
int slurm_change_dep();
```

Call the above function to change all afterok:\$(SLURM_JOBID) dependencies into {{{after:\$(SLURM_JOBID)}} dependencies. This enables the jobs in workflow eligible for allocation by Slurm.

The header file can be included using #include <slurm/slurm_workflow.h> and should be linked using -lslurm_workflow and -lslurm.

Information on past jobs and accounting

The sacct command can be used to enquire the Slurm database about a past job.

[kreutz1@dee]	[kreutz1@deepv Temp]\$ sacct -j 69268							
JobID	JobName	Partition	Account	AllocCPUS	State 1	ExitCode		
69268+0	bash	dp-cn	deepest-a+	96	COMPLETED	0:0		
69268+0.0	MPI_Hello+		deepest-a+	2	COMPLETED	0:0		
69268+1	bash	dp-dam	deepest-a+	384	COMPLETED	0:0		

On the Cluster (CM) nodes it's possible to query the consumed energy for a certain job:

```
[kreutz1@deepv kreutz1]$ sacct -o ConsumedEnergy,JobName,JobID,CPUTime,AllocNodes -j 69326
ConsumedEnergy JobName JobID CPUTime AllocNodes
496.70K hpl_MKL_0+ 69326 16:28:48 1
0 batch 69326.batch 16:28:48 1
496.70K xlinpack_+ 69326.0 08:10:24 1
```

This feature will also be for the ESB nodes.

FAQ

Is there a cheat sheet for all main Slurm commands?

Yes, it is available ?here.

Why's my job not running?

You can check the state of your job with

scontrol show job <job id>

In the output, look for the Reason field.

You can check the existing reservations using

scontrol show res

How can I check which jobs are running in the machine?

Please use the squeue command (the "-u \$USER" option to only list jobs belonging to your user id).

How do I do chain jobs with dependencies?

Please confer the sbatch/srun man page, especially the

-d, --dependency=<dependency_list>

entry.

Also, jobs can be chained after they have been submitted using the scontrol command by updating their Dependency field.

How can check the status of partitions and nodes?

The main command to use is sinfo. By default, when called alone, sinfo will list the available partitions and the number of nodes in each partition in a given status. For example:

[deamicis1@d	eepv hy	bridhello]\$	sinfo		
PARTITION	AVAIL	TIMELIMIT	NODES	STATE NODELIST	
sdv	up	20:00:00	11	idle deeper-sdv[06-16]	
knl	up	20:00:00	1	drain knl01	
knl	up	20:00:00	3	idle knl[04-06]	
knl256	up	20:00:00	1	drain knl01	
knl256	up	20:00:00	1	idle knl05	
knl272	up	20:00:00	2	idle knl[04,06]	
snc4	up	20:00:00	1	idle knl05	
extoll	up	20:00:00	11	idle deeper-sdv[06-16]	
ml-gpu	up	20:00:00	3	idle ml-gpu[01-03]	
dp-cn	up	20:00:00	1	drain dp-cn33	
dp-cn	up	20:00:00	5	resv dp-cn[09-10,25,49-50]	
dp-cn	up	20:00:00	44	idle dp-cn[01-08,11-24,26-32,34-48]	18]

dp-dam	up	20:00:00	1	drain*	dp-dam08
dp-dam	up	20:00:00	2	drain	dp-dam[03,07]
dp-dam	up	20:00:00	3	resv	dp-dam[05,09-10]
dp-dam	up	20:00:00	2	alloc	dp-dam[01,04]
dp-dam	up	20:00:00	8	idle	dp-dam[02,06,11-16]
dp-dam-ext	up	20:00:00	2	resv	dp-dam[09-10]
dp-dam-ext	up	20:00:00	б	idle	dp-dam[11-16]
dp-esb	up	20:00:00	51	drain*	dp-esb[11,26-75]
dp-esb	up	20:00:00	2	drain	dp-esb[08,23]
dp-esb	up	20:00:00	2	alloc	dp-esb[09-10]
dp-esb	up	20:00:00	20	idle	dp-esb[01-07,12-22,24-25]
dp-sdv-esb	up	20:00:00	2	resv	dp-sdv-esb[01-02]
psgw-cluster	up	20:00:00	1	idle	nfgw01
psgw-booster	up	20:00:00	1	idle	nfgw02
debug	up	20:00:00	1	drain*	dp-dam08
debug	up	20:00:00	4	drain	dp-cn33,dp-dam[03,07],knl01
debug	up	20:00:00	10	resv	dp-cn[09-10,25,49-50],dp-dam[05,09-10],dp-sdv-esb[01-02]
debug	up	20:00:00	2	alloc	dp-dam[01,04]
debug	up	20:00:00	69	idle	deeper-sdv[06-16],dp-cn[01-08,11-24,26-32,34-48],dp-dam[02,06,11-16],knl[04-06

Please refer to the man page for ${\tt sinfo}$ for more information.

Can I join stderr and stdout like it was done with -joe in Torque?

Not directly. In your batch script, redirect stdout and stderr to the same file:

```
...
#SBATCH -o /point/to/the/common/logfile-%j.log
#SBATCH -e /point/to/the/common/logfile-%j.log
...
```

(The %j will place the job id in the output file). N.B. It might be more efficient to redirect the output of your script's commands to a dedicated file.