

Access and usage of the IGBMC HPC cluster

First you need to verify that your environment is properly set.

```
>ssh login@surf
```

The sbatch command allow you to submit a script with commands to the cluster

```
>SBATCH -p surf cmd
```

- The command then gives a job execution number.
- `>sccontrol show job <job-number>` allows you to follow the execution of the job
- `>scancel <job-number>` allows you to kill the job

The job will generate output files:

- `slurm-<job-number>.out` in the directory from which the sbatch command was executed
- `slurm-<job-number>.e` in case of errors

In case of necessity the debug line `sbatch -p debug` allows you to use two additional lames (16 procs with 24GB memory)

There are options to:

- specify the memory required `>sbatch -p surf -mem=xxMB`
- specify the number of processors required `>sbatch -p surf -cpus-per-task=24` (if you specify more than 24, the number of available lames for your job will be very limited)

The cmd script must start with

```
#!/bin/bash
```

and should set the necessary environment. All alias can be checked with the command `>alias`

From:

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