

High-performance computing

High-performance computing (HPC) typically refers to using parallel data processing across multiple computers in order to analyze large data sets in a timely manner. HPC works well for many bioinformatics applications, because many of the analysis steps can proceed independently (e.g. at the same time on different computers).

Running an interactive job

Due to the possibility of being disconnected from the cluster, you may want to use `tmux` or `screen` when running interactive jobs. These programs allow you to disconnect from a terminal session and reconnect later. For example, before requesting the interactive job create a `tmux` session:

```
$ tmux new -s myjob
```

This will create a new `tmux` session named `myjob`. You can then request the interactive job and run your commands. If you are disconnected (or if you detach using `Ctrl + b` then `d`) you can reattach to the `tmux` session:

```
$ tmux attach -t myjob
```

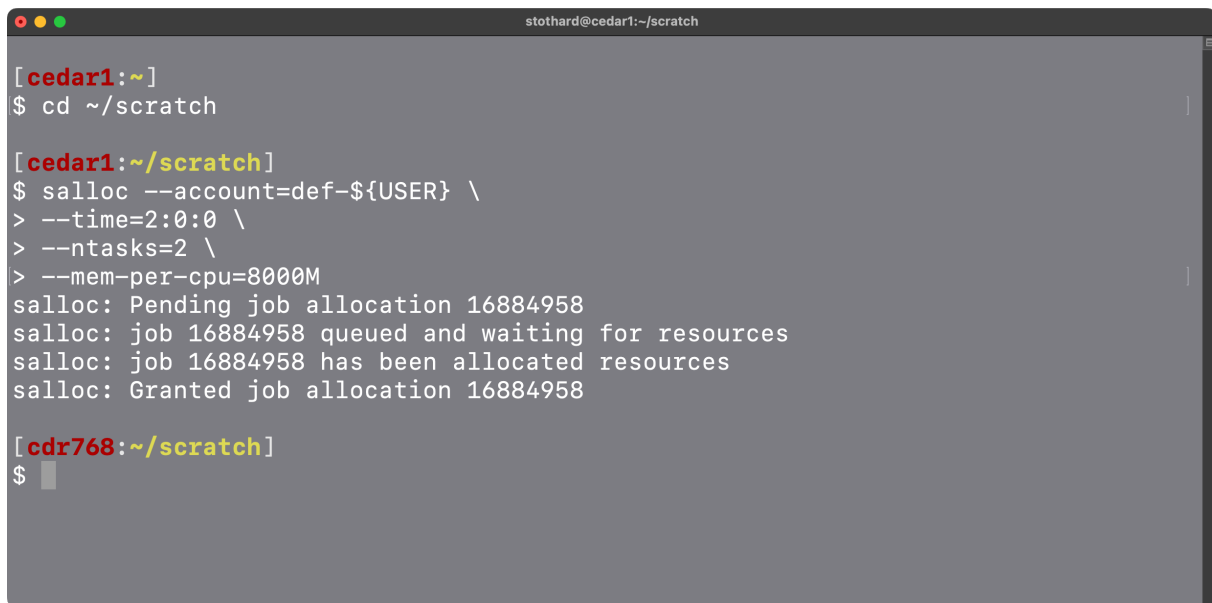
Note that there may be multiple login nodes, and you may be connected to a different login node when you reconnect. If you are unable to reconnect to the `tmux` session, try connecting to a different login node. For example, if you were on the `cedar1` login node when you created the `tmux` session, but are now on the `cedar5` login node, you can quickly switch to the `cedar1` login node using `ssh cedar1`.

If you find that you are frequently disconnected from the cluster, try adding the following to the `~/.ssh/config` file **on your computer** before connecting to the cluster:

```
Host *
  ServerAliveInterval 300
  ServerAliveCountMax 5
  TCPKeepAlive yes
```

On the login node request an interactive job:

```
$ cd ~/scratch
$ salloc --account=def-${USER} \
--time=2:0:0 \
--ntasks=2 \
--mem-per-cpu=8000M
```

A terminal window titled 'stothard@cedar1:~/scratch' showing a user at the 'cedar1' login node. The user navigates to the '~/scratch' directory and runs the 'salloc' command with options: '--account=def-\${USER}', '--time=2:0:0', '--ntasks=2', and '--mem-per-cpu=8000M'. The terminal output shows the job being queued, allocated, and granted. The prompt then changes to 'cdr768:~/scratch', indicating the user has been moved to a compute node.

```
[cedar1:~]
$ cd ~/scratch

[cedar1:~/scratch]
$ salloc --account=def-${USER} \
> --time=2:0:0 \
> --ntasks=2 \
> --mem-per-cpu=8000M
salloc: Pending job allocation 16884958
salloc: job 16884958 queued and waiting for resources
salloc: job 16884958 has been allocated resources
salloc: Granted job allocation 16884958

[cdr768:~/scratch]
$
```

Figure 1: Requesting an interactive job

The `salloc` options mean the following:

| Option | Meaning |
|-------------------------------------|--|
| <code>--account=def-someuser</code> | Use the default allocation for this user |
| <code>--time=2:0:0</code> | Reserve the compute node for 2 hours |
| <code>--ntasks=2</code> | Reserve 2 CPU cores |
| <code>--mem-per-cpu=8000M</code> | Reserve 8000MB of RAM per CPU |

You can request longer times and more RAM and CPU resources. However, the more you request the more likely your `salloc` command will leave you waiting for the resources to become available.

Once the allocation is available you will be logged in to a compute node where you are free to run your commands. You will have access to the same files and folders as on the login node, and any files you create on a compute node will be accessible afterwards from the login node.

Typically you work within the `~/scratch` folder when using compute nodes, hence the `cd ~/scratch` prior to requesting the allocation. When you exit the compute node you can continue to access the contents of this folder from the login node. Note that the contents of `~/scratch` are regularly removed by administrators (with warning) so don't use it for long-term storage. Place files that you want to keep in your `project` folder. Each research group is given a default allocation of 1 TB of "project" storage.

Tutorial: High-performance computing

To use a module, e.g. `emboss`:

```
$ module load emboss
```

Once loaded you can use the usual commands for that package, e.g.:

```
$ stretcher -version
```

A terminal window titled 'stothard@cedar1:~/scratch' showing a sequence of commands and their outputs. The prompt is '[cdr768:~/scratch]'. The first command is '\$ module load emboss'. The second command is '\$ stretcher -version', which outputs 'EMBOSS:6.6.0.0'. The third command is '\$', which shows a cursor.

```
stothard@cedar1:~/scratch  
[cdr768:~/scratch]  
$ module load emboss  
  
[cdr768:~/scratch]  
$ stretcher -version  
EMBOSS:6.6.0.0  
  
[cdr768:~/scratch]  
$
```

Figure 2: Loading a module