

Mila & DRAC

cheat sheet

by IDT team

getting help

- search in `docs.mila.quebec`
- search for specific strings (or error messages) on the Mila slack
- visit #mila-cluster and #compute-canada (for DRAC)
- visit specific tool channels such as #pytorch and #jax
- go to the IDT office hours (Tuesday 3PM-5PM) by just walking into the IDT lab (room A.17) and saying hi
- open an IT support ticket by emailing it-support@mila.quebec
- contact DRAC support at support@tech.alliancecan.ca

milatools

Quick way to setup SSH to Mila cluster

```
pip install -U milatools; mila init
```

Open VSCode connected to an interactive session on compute node

```
mila code /path/work --alloc [salloc arguments]
```

Milatools can also connect directly to a DRAC cluster (e.g. cedar)

```
mila code /path/work --cluster=cedar --alloc  
--account=def-bengioy [other salloc arguments]
```

Inside VSCode you can also open a remote SSH to mila-cpu to automatically create an interactive session to a CPU node (configured by mila init). You can also ssh mila-cpu from a terminal.

Never run a program that takes more than a few seconds on a login node. Do not edit files remotely with VSCode directly on login nodes.

modules

(NB: Many of those modules are outdated.)

<code>module avail</code>	Displays all the available modules
<code>module load <module></code>	Loads <module>
<code>module spider <module></code>	Shows details about <module>
<code>module load python/3.10</code>	Load python 3.10 to use it
<code>module load httpproxy</code>	Allows Wandb and Comet on DRAC

Slurm commands

```
salloc --gres=gpu:1 -c 2 --mem=12000  
Get an interactive job with one GPU, 2 CPUs and 12000 MB RAM  
sbatch  
Start a batch job (same options as salloc)  
sattach --pty <jobid>.0  
Re-attach a dropped interactive job  
sinfo  
Status of all nodes  
sinfo -O gres:27,nodelist,features -tidle,mix,alloc  
List GPU type and FEATURES that you can request  
savail  
List available gpu (Mila only)  
partition-stats [-v]  
Similar functionality to savail (DRAC only)  
scancel <jobid>  
Cancel a job  
squeue -u $USER  
Summary status of all YOUR active jobs  
squeue -j <jobid>  
Summary status of a specific job  
squeue -O jobid,name,username,partition,state,timeused,nodelist,gres,tres  
Status of all jobs including requested resources  
(see the SLURM squeue doc for all output options)  
scontrol show job <jobid>  
Detailed status of a running job  
sacct -j <job_id> -o NodeList  
Get the node where a finished job ran  
sacct -u $USER -S <start_time> -E <stop_time>  
Find info about old jobs  
sacct -oJobID,JobName,User,Partition,Node,State  
List of current and recent jobs
```

Remember that every map is a simplification of reality.
This is a cheat sheet for Mila students using Slurm,
not a full tutorial, and also not a Linux/Git/PyTorch guide.
See docs.mila.quebec/Cheatsheet.html for pdf,
along with errata. Anticipate one update per year.
The complete up-to-date documentation at docs.mila.quebec.

PREPARED ON

2024-04-01

BEST BEFORE

2025-04-01

sbatch / salloc commands

<code>-n, --ntasks=<number></code>	Number of task in your script, usually =1
<code>-c, --cpus-per-task=<ncpus></code>	Number of cores for each task
<code>-t, --time=<time></code>	Time requested for your job
<code>--mem=<size[units]></code>	Memory requested for all your tasks
<code>--gres=<list></code>	Select generic resources such as GPUs: --gres=gpu:GPU_MODEL
<code>-p, --partition=<name></code>	Partition for resource sharing (Mila cluster only)
<code>--account=<name></code>	DRAC allocation for resources (DRAC only)
<code>-x, --exclude=<nodelist></code>	Exclude certain nodes from job submission

sbatch script example

```
#!/bin/bash  
#SBATCH --ntasks=1 # Default 1 task, optional  
#SBATCH --partition=unkillable # Ask for unkillable job  
#SBATCH --cpus-per-task=2 # Ask for 2 CPUs  
#SBATCH --gres=gpu:1 # Ask for 1 GPU  
#SBATCH --mem=10G # Ask for 10 GB of RAM  
#SBATCH --time=3:00:00 # The job will run for 3 hours  
#SBATCH -o /network/scratch/<u>/<username>/slurm-%j.out  
  
# Load the required modules  
module --quiet load anaconda/3  
# Load your environment  
conda activate "<env_name>"  
# Copy your dataset on the compute node  
cp /network/datasets/<dataset> $SLURM_TMPDIR  
# Launch your job, tell it to save the model in $SLURM_TMPDIR  
# and look for the dataset into $SLURM_TMPDIR  
python main.py --path $SLURM_TMPDIR --data_path $SLURM_TMPDIR  
# Copy whatever you want to save on $SCRATCH  
cp $SLURM_TMPDIR/<to_save> /network/scratch/<u>/<username>/
```

multi-GPU, multi-node

1 node with 1 GPU

See docs.mila.quebec/examples/distributed/index.html for minimalist standalone code.

```
#SBATCH --gpus-per-task=rtx8000:1  
#SBATCH --cpus-per-task=4  
#SBATCH --ntasks-per-node=1  
#SBATCH --mem=16G  
#SBATCH --time=00:15:00
```

1 node with 4 GPUs

```
#SBATCH --gpus-per-task=rtx8000:1  
#SBATCH --cpus-per-task=4  
#SBATCH --ntasks-per-node=4  
#SBATCH --mem=16G  
#SBATCH --time=00:15:00
```

2 nodes with 4 GPUs each

```
#SBATCH --gpus-per-task=rtx8000:1  
#SBATCH --cpus-per-task=4  
#SBATCH --ntasks-per-node=4  
#SBATCH --nodes=2  
#SBATCH --mem=16G  
#SBATCH --time=00:15:00
```

If you have N parallel jobs that each require 1 GPU, don't try to schedule them in a multi-GPU way. Submit many separate jobs, maybe use job arrays, or consider packing many experiments in a single job with 1GPU.

checkpointing, profiling, scaling

Powerful GPUs cost approximately \$1/h when amortized over their expected life.
If you use only one GPU for active development, it's acceptable to be inefficient.
Consider using a less powerful GPU or a "MIG" instance if possible.

Things change when you run large-scale experiments. You need to

- profile your code to make sure you properly use the GPUs allocated (i.e. "GPU Utilization"),
- use checkpoints properly to resume your experiments when they crash or get preempted.

Easy ways to measure "GPU Utilization" include Wandb, `nvidia-smi` and the DRAC "portal".
See also docs.mila.quebec/examples/good_practices/checkpointing/index.html for an example of proper checkpointing.

Research involves exploring and testing ideas that don't necessarily work out in the end.
This is a good use of the cluster when done properly. Mila is a research institute.

Don't be the researcher who runs 200 jobs each running for 24h and using only 2% of a GPU.
They've just wasted \$5000. Lack of proper checkpointing can lead to same levels of waste.
Avoid grid search for hyperparameter optimization. Better tools reduce unnecessary computation.

Mila

docs.mila.quebec official docs
dashboard.server.mila.quebec node and GPU monitoring
datasets.server.mila.quebec datasets already shared on cluster
clockwork.mila.quebec dashboard for jobs (currently in beta)

The Mila cluster is available for **all students supervised by a Mila core prof** and for Mila employees.
Not MsPro students, nor students of non-core Mila profs. Exceptions exist.

Node Name	Qty	N	GPU Model	Mem (GB)	CPU Cores	Mem (GB)	Tmp (TB)	SLURM features	optimal ratios GPU:CPU:RAM
GPU compute nodes									
cn-a[001-011]	11	8x	RTX8000	48	40	384	3.6	turing,48gb	1 : 5 : 48GB
cn-b[001-005]	5	8x	V100	32	40	384	3.6	volta,nvlink,32gb	1 : 5 : 48GB
cn-c[001-040]	40	8x	RTX8000	48	64	384	3	turing,48gb	1 : 8 : 48GB
cn-g[001-029]	29	4x	A100	80	64	1024	7	ampere,nvlink,80gb	1 : 16 : 256GB
cn-i001	1	4x	A100	80	64	1024	3.6	ampere,80gb	1 : 16 : 256GB
cn-j001	1	8x	A6000	48	64	1024	3.6	ampere,48gb	1 : 8 : 128GB
cn-k[001-004]	1	2x	V100	16	16	256	3.6	volta,16gb	1 : 8 : 128GB

DGX Systems									
cn-d[001-002]	2	8x	A100	40	128	1024	14	ampere,nvlink,dgx,40gb	1 : 16 : 32GB
cn-d[003-004]	2	8x	A100	80	128	2048	28	ampere,nvlink,dgx,80gb	1 : 16 : 64GB
cn-e[002-003]	2	8x	V100	32	40	512	7	volta,nvlink,dgx,32gb	1 : 5 : 16GB

CPU compute nodes									
cn-f[001-004]	4	-	-	-	32	256	10	rome	0 : 1 : 8GB
cn-h[001-004]	4	-	-	-	64	768	7	milan	0 : 1 : 12GB

MIG (a fractional part of a powerful GPU)

a100l.2g.20gb a100l.2	A100 (20GB, 2/7 of compute) sbatch --gres=gpu:a100l.2	48 available -c=4 --mem=32G ...
a100l.3g.40gb a100l.3	A100 (40GB, 3/7 of compute) sbatch --gres=gpu:a100l.3	48 available -c=8 --mem=64G ...
a100l.4g.40gb a100l.4	A100 (40GB, 4/7 of compute) sbatch --gres=gpu:a100l.4	24 available -c=8 --mem=64G ...

partition name	max resource usage	max time	note
unkillable	6 CPUs, mem=32G, 1 GPU	2 days	
unkillable-cpu	2 CPUs, mem=16G	2 days	CPU-only jobs
short-unkillable	24 CPUs, mem=128G, 4 GPUs	3 hours (!)	
main	8 CPUs, mem=48G, 2 GPUs	5 days	
main-cpu	8 CPUs, mem=64G	5 days	CPU-only jobs
long	no limit of resources	7 days	
long-cpu	no limit of resources	7 days	CPU-only jobs

Jobs on the Mila cluster are all *preemptible*, except for those in the *unkillable* partitions. This means that they can be terminated and requested automatically to allow higher-priority jobs to run (based on partition preemption order *unkillable* > *main* > *long*). There is a very limited number of jobs that can run in *unkillable* partitions.

Partitions are specified with the `--partition` flag (obsolete in late 2024).

path	storage/inodes	speed	backup?	mounted
\$HOME	100GB / 1M	low	yes	all nodes
\$SCRATCH	20TB / infity	high	no	all nodes
\$SLURM_TMPDIR	-	highest	no	cn-*
/network/projects	varies	medium	no	all nodes
/network/datasets	read-only	high	no	all nodes
/network/weights	read-only	high	no	all nodes
\$ARCHIVE	500GB	low	no	login-*

Use `disk-quota` to see your current usage of storage (\$HOME and \$SCRATCH).

Use `savail` to list the GPUs available. Slurmatively, go to dashboard.server.mila.quebec.

MIG instances should never be used for multi-GPU training. It is slow and absurd compared to using a single full GPU. For up-to-date tips to avoid hitting a MIG node in your multi-GPU training, refer to either docs.mila.quebec or the [#mila-cluster](#) Slack channel. MIG instances have no graphics APIs (OpenGL/Vulkan).

DRAC

docs.alliancecan.ca/wiki
portail.narval.calculquebec.ca/secure/jobstats/<username>/<jobid>
portail.beluga.calculquebec.ca/secure/jobstats/<username>/<jobid>

DRAC (Digital Research Alliance of Canada, formerly known as Compute Canada) offers access to compute clusters to all researchers in Canada. Any prof with an account on DRAC can add ("sponsor") whoever they want on their "default" allocation, but they will usually add their own students. Additionally, **all students supervised by a Mila core prof** and all Mila employees can be added to a "mega allocation" under Yoshua Bengio's name. See https://docs.mila.quebec/Extra_compute.html#account-creation.

DRAC uses a concept of "Reference GPU Unit" (RGU) to measure allocated GPUs in a way that attributes a different costs to GPUs based on their performance and memory. In the first column below, we provide values for RGU-years as well as their GPU-years equivalents as estimates (in brackets). Ex: P100-12GB is 1 RGU. A100-40GB is 4.0 RGU.

	(shared mega-allocation)		(your supervisor's default allocation)		GPU types	unrestricted internet?	Wandb?	Comet?
	rrg-bengio-ad_gpu	rrg-bengio-ad_cpu	def-yourprof-gpu	def-yourprof-cpu				
narval	440 (~110)	580	3 GPUs	195 cores	A100	no	httpproxy (limited)	httpproxy
beluga	250 (~111)	125	4 GPUs	63 cores	V100	no	httpproxy (limited)	httpproxy
cedar	305 (~118)	125	3 GPUs	71 cores	P100, V100	yes	yes	yes

RGUs (~GPUs)

CPU cores

Columns with *rrg-bengio* values are guaranteed resources for the year. However, the *def-yourprof* values are estimates of the best effort to share excess resources available across Canada, and they could be off by a factor of 2 in reality.

1. Contrary to the Mila cluster, DRAC allocations have a single *fair share* value for all users under the account. Jobs that are the easiest to run will run first, no matter to whom they belong. This might not feel fair. IDT does not control this. DRAC does.

Don't be a bad actor by submitting 1k small jobs because you will end up monopolizing the clusters to the detriment of everyone. This is the price for DRAC clusters having preemption disabled.

If your jobs are queued and never run, try to make them more appealing to the scheduler by asking for optimal resources.

2. You have access to "default accounts" that starts with a "def-". These are shared between members of your research group instead of the whole Mila. This resources are underused. Free CPUs/GPUs!

3. The shared storage on DRAC is particular because we run out of inodes (i.e. number of files) faster than the actual storage space.

4. IDT does not admin rights on DRAC clusters.

DRAC compute nodes have similar roles as the Mila cluster for \$HOME, \$SCRATCH and \$SLURM_TMPDIR. See also `$HOME/projects/<account>/<your_username>`. Use `diskusage_report` to see usage.

jobs that the scheduler likes

time bins <=3h <=12h <=24h <=72h <=168h

GPU:CPU:RAM ratios	Cedar	1 : 8 : 46G x4 (V100 32GB)
Beluga	favor10 : 46G x4	1 : 6 : 31G x4 (P100 12GB)
Narval	1 : 12 : 123G x4	1 : 6 : 62G x4 (P100 16GB)

DRAC clusters use a different method to queue jobs. You cannot specify a `--partition`.

Jobs fall in "bins" based on time and resources requested. Ask for things that are easy to schedule, the scheduler will be much nicer to you. If you break your 12h job into 4 chunks of 3h (with checkpointing), you will get resources more easily. If you ask for 13h, you will be put into the <=24h bin, which is not advantageous to you.

If you ask for certain ratios of GPU:CPU:RAM when submitting jobs, the scheduler will also favor you.