Introduction to supercomputing for AI

High performance computing (HPC) resources can be used to accelerate AI workflows. The EuroHPC Joint Undertaking (JU) offers free access to such resources to SMEs as well as larger companies. In this hands-on, you will learn:

- What is an HPC resource and how it is different from a cloud environment;
- What are the available HPC resources through the EuroHPC JU;
- How to connect to a cluster and explore resources:
- How to run a demo Al workflow based on Singularity.

Prerequisites

You will need to have credentials to access the PDC cluster. A working SSH client is needed: it is included on macOS and most Linux flavours; it is also available on Windows in the Powershell or under the Windows Subsystem for Linux (WSL).

Who is the course for?

This course is intended for data scientists that want to take advantage of higher computing power to perform their workflows. Some degree of familiarity with a command-line shell is recommended, but no expertise is required. No previous knowledge of supercomputing environments is required.

About the course

We will train a Unet model to be able to recognise water in satellite pictures. The source code can be found at this repo. The example is based on Tensorflow and will be run using Singularity. The structure of the example is the following:

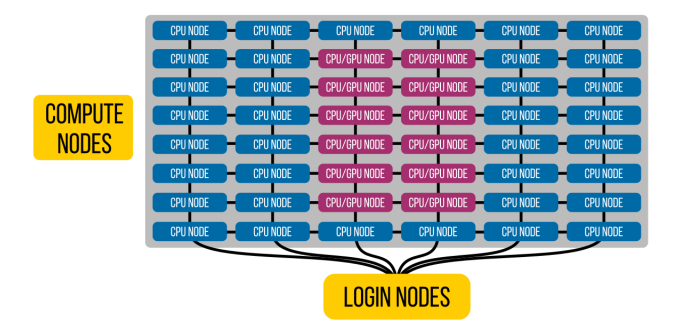
```
./supercomputing4ai_demo
 build_singularity.def
  - images
   - models
   ├─ serving
     └─ main.py
   └─ unet
       ├─ data
          └─ water
              ├─ Images
             └─ Masks
         - main.py
         - result
          — models
          └─ training
  - README.md
```

The models subfolder contains the model to be trained (unet) and the inference code (serving). Under data, the training dataset can be found, with the Images being some satellite images and Masks being the water-covered areas in those images. Upon running main.py in the unet folder, a Unet will be trained, producing a set of weights in the models/ subfolder and training statistics (binary cross-entropy loss and accuracy). Inference is then performed with the models/serving/main.py script, which takes as an input an image and generates a mask of the water parts.

20 min	What is a supercomputer and how is it different from cloud?
20 min	Connecting to a HPC resource
20 min	Move between folders, Is, transferring to/from local storage
20 min	Available software and modules
20 min	Queueing jobs and running a Singularity container

What is a supercomputer and how is it different from cloud?

A supercomputer (or *cluster*) is a high-performance computing infrastructure. The general idea is to have many (thousands) of machines, called *nodes*, working in parallel to increase performance. Each node can have one or several (usually two) multi-core CPUs, local RAM, and possibly accelerators such as GPUs. What distinguishes a cluster from a simple collection of computers is the high-speed connection between all the nodes, called *interconnect*. The speed and density of connections between nodes makes it possible for the nodes to seamlessly communicate and solve a problem in parallel. The general architecture of a cluster looks like below:



Schematics of a typical cluster.

The user accesses the cluster through the *login nodes*, which act as a gateway between the external world and the compute parts. They should not be used to run calculations. Usually a parallel file system is provided (such as Lustre), with independent servers (not shown in the figure) that manage distributed access from all the nodes. Moreover, each node usually has some temporary storage called *scratch* for fast access during computation. On Dardel specifically, each CPU node has two AMD EPYC CPUs with 64 physical cores each and RAM ranging from 256 GB to 2 TB. Each GPU node has one AMD EPYC and 4 AMD Instinct MI250X GPUs, each containing two GCDs (compute chips in AMD lingo); thus, each node has 8 GPUs.

How is HPC different from cloud?

While they both provide access to high-performance computational resources, a few important differences exist between cloud and HPC systems:

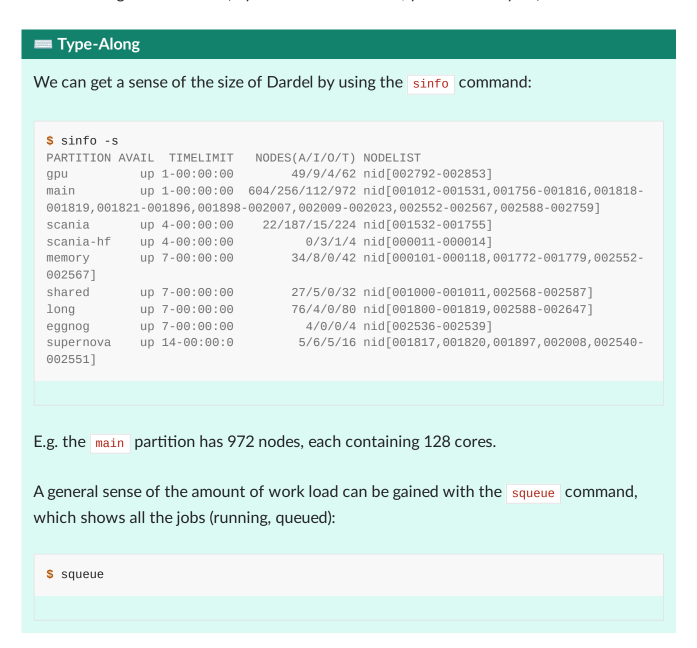
- An HPC resource is shared among several users; usually users do not have root access, unlike cloud where each user has their own sandboxed environment;
- Each workload pushed by a user (called *job*) contends for the available resources; usually, a queueing system is managed by a *job manager* (more on that later) which decides which jobs are going to run to maximise cluster usage. This means that a job may have to wait for resources before it is allowed to run.
- (Through EuroHPC) the cost of HPC does not scale with amount of resources used; with cloud it does.
- Engineering/scientific applications usually expect an HPC-like environment which is harder to set up in the cloud;
- Access to bare metal vs virtual machines.

Connecting to a HPC resource

Once SSH keys are created and uploaded on the PDC interface, entering the cluster is as simple as:

```
$ ssh -Y <username>@dardel.pdc.kth.se
```

Which should get you into the PDC supercomputer. The _-y flag is used to be able to open graphical windows on the supercomputer, e.g. to visualise images. This will work only if you have a running local X server (if you are on Linux/WSL, you most likely do).



Move between folders, ls, transferring to/from local storage

Upon logging in, you should be in your "home" folder, as reported by the prompt:

```
fiusco@login1:~>
```

where login is the name of the host (in this case, the login node) and represents the home folder. The full path of this directory can be printed using the pwd command (print working directory):

```
$ pwd
/cfs/klemming/home/f/fiusco
```

The contents of a directory can be listed with the ls command:

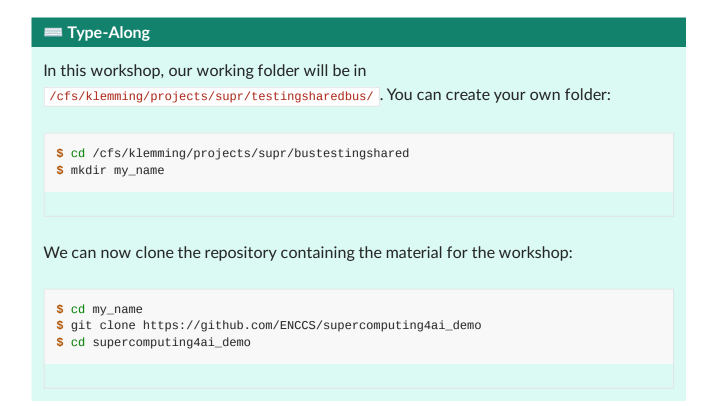
```
$ ls
Private Public spack-user
```

The cd (change directory) command can be used to navigate the filesystem.

Moving files/folders from/to the cluster can be achieved via the scp command to be run locally (i.e. not on the cluster):

```
$ scp [-r] user@dardel.pdc.kth.se:/path/to/source /path/to/local/destination
```

The optional _-r flag is used to indicate recursive copying of whole folders and their contents.



Available software and modules

A variety of software, libraries and compiler toolchains is usually available on most HPC resources. Usually a module system is used to access software provided by the cluster. A list of all available modules can be printed with the following syntax

```
$ module avail
```

The presence of a particular piece of software/library can be gueried with:

```
$ ml spider <name>
```

This command will return all available versions of a module (if present), as well as specific instructions to load it if needed. The module can then be loaded with module_name(s). For example, the Julia runtime can be loaded with:

```
$ ml PDC julia
```

Type-Along

We will use a Singularity container to get a Tensorflow environment. For that, we need a few modules:

```
$ ml PDC Singularity
```

We can also inspect the produced images directly on the cluster using the display command, available in the ImageMagick toolkit:

\$ ml ImageMagick

Queueing jobs and running a Singularity container

Most supercomputers use a system called SLURM to manage jobs and maximise cluster utilisation. There are two main workflows:

- Interactive mode, in which the user asks for resources, logs into the compute nodes and runs all they need to run
- Batch mode, in which the user prepares a submit script (usually Bash) that contains all the instructions to be executed. The script is then placed in the queue and runs without any further inputs.

Interactive resources can be requested with the salloc command in the following fashion:

```
$ salloc -n <n_cores> -t HH:MM:SS -A <allocation_number> -p <partition>
```

Most parameters are self-explanatory. -t is the wall time, -A is an allocation dependent on the project. -p is required by some clusters that have different types of nodes, e.g. Dardel has a set of nodes reserved for interactive use on a partition called shared and a set of nodes with GPUs (gpu). There are many other options that are explained here. Once the requested resources are granted, a MPI job can be executed with srun -n <n_cores>my_command; alternatively, the user can also ssh directly into the allocated node.

Type-Along

Let's book a node with a GPU to run our script:

```
$ salloc -n 1 -t 00:50:00 -A pdc-bus-2024-8 -p gpu --gpus=1
```

Once we get our node, we can train our model:

```
$ srun -n 1 singularity exec --rocm -B ./models:/models
/cfs/klemming/projects/supr/bustestingshared/ENCCS/rocm_tensorflow/ python
models/unet/main.py
```

The _B flag is used to bind a directory on the filesystem to a directory in the container and the _-rocm flag is used to expose the GPU to the container. The _rocm_tensorflow folder contains a pre-built container with the necessary Python packages. If the ImageMagick module was loaded, we can inspect a the accuracy and losses in the _results/training folder with the _display command. Once the network is trained, we can perform inference:

```
$ srun -n 1 singularity --exec --rocm -B ./models:/models,./images:/images/cfs/klemming/projects/supr/bustestingshared/ENCCS/rocm_tensorflow python models/serving/main.py -f water_body_17.jpg
```

The image and generated mask can be shown with:

```
$ display images/water_body_17.jpg
$ display images/generated-images/water_body_17.jpg
```

Alternatively, you can open another local terminal (i.e. on your computer) and copy the images back for local visualisation:

```
scp -r
<yourusername>@dardel.pdc.kth.se:/cfs/klemming/projects/supr/bustestingshared/<yournam
/local/path</pre>
```

Bonus - Batch mode job

It is often easier to submit a job to the general queue rather than waiting for an interactive resource. As an example, we can book two nodes and print their host name. Let's create the following submission script and call it *submit_test.sh*:

```
#!/bin/bash -1
#SBATCH -J test # Job name
#SBATCH -t 00:02:00 # Wall time
#SBATCH -A pdc-bus-2024-8 # Allocation number
#SBATCH -p main # Partition - in this case the normal (non-GPU) is fine
#SBATCH -nodes 2 # We want two different nodes
#SBATCH -ntasks-per-node=1 # We want to book only one core on each node - We need
just the hostname after all :)
#SBATCH --mail-type=ALL # Get an email when the job starts and when the job ends
# Now for the actual instruction to be executed
srun hostname > out.txt 2>&1 # Black magic to redirect both stdout and stderr to
the same file
```

We can now submit our short job to the queue:

```
$ sbatch submit.sh
```

We can check the status in queue with:

```
$ squeue -u <username>
```

Quick Reference

Instructor's guide

Why we teach this lesson

Intended learning outcomes

Timing

Preparing exercises

e.g. what to do the day before to set up common repositories.

Other practical aspects

Interesting questions you might get

Typical pitfalls See also

Further introductory material can be found on the Introduction to LUMI and HPC carpentry pages.

Credits

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