#### Introduction to scientific and parallel programming in economics

#### Parallel computing infrastructure

Simon Scheidegger simon.scheidegger@unil.com August 30<sup>th</sup>, 2018

Cowles Foundation – Yale

#### **Outline**

- Make first steps on a Linux Cluster
  Login via ssh, remotely, short overview of basic unix commands like cd, pwd, cp, scp,...
- Submit jobs to the queue
- Get lecture notes
  Clone a git repository

#### https://research.computing.yale.edu

For this course, we use Yale's HPC compute cluster.

Yale Center for Research Computing (YCRC)

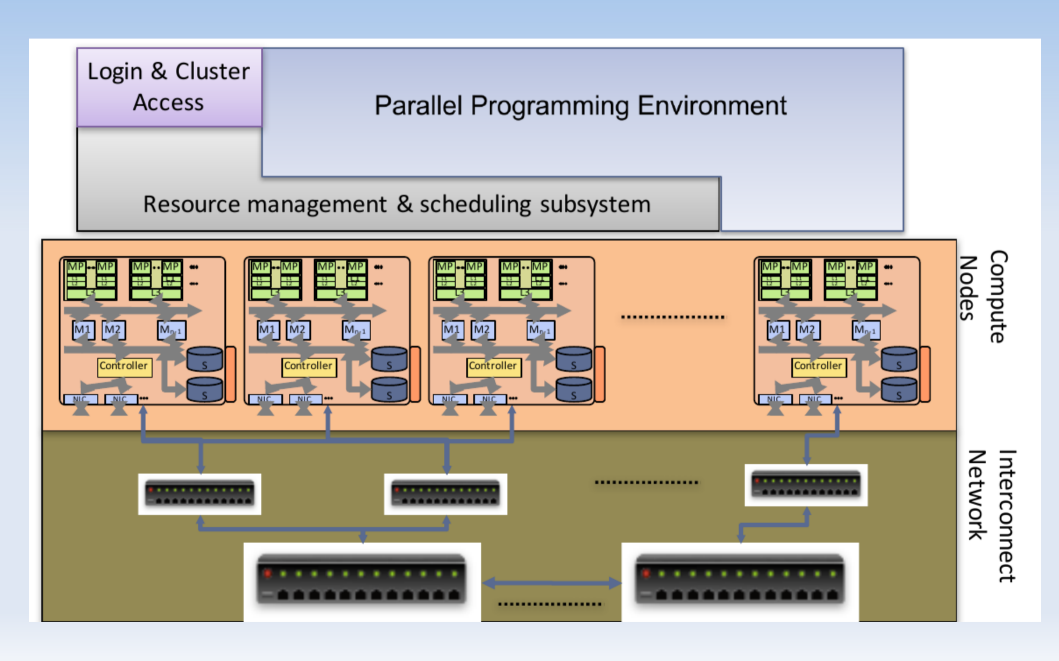
→ Its setup is very similar to any other top system

For the Manual see the documentation site at

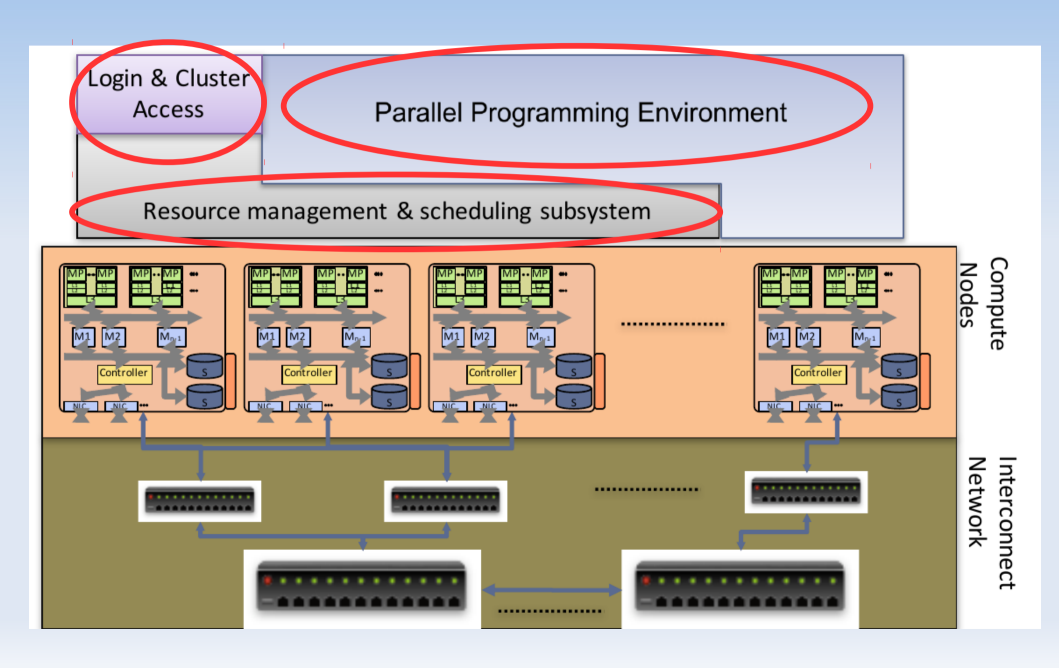
https://research.computing.yale.edu/support/hpc/getting-started

https://research.computing.yale.edu/support/hpc/user-guide

#### An abstract compute cluster

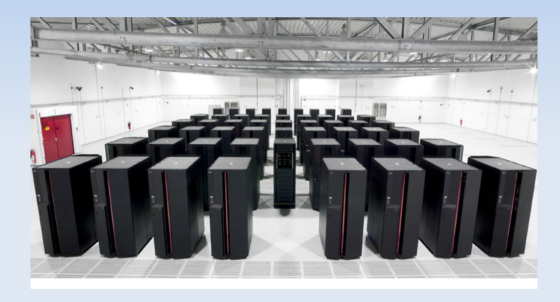


### An abstract compute cluster



# A real compute cluster





### Login for participants

- If you don't have an account on Yale's HPC infrastructure, request one (infrastructure for this course)

  https://research.computing.yale.edu/support/hpc/account-request
- For MS-Windows users: Download and install Putty
  - → http://the.earth.li/~sgtatham/putty/latest/x86/putty.exe
  - → Download and install Winscp
  - → http://winscp.net/download/winscp576setup.exe

# **Basic Linux commands (1)**

Command	Description		
pwd	Print name of current/working directory		
cd [Directory]	Change directory (no directory $\rightarrow$ change to home)		
ls [Directory]	List directory contents (no directory $ o$ list current)		
cat FILE	Concatenate files and print on the standard output		
mkdir DIRECTORY	Make directories		
mkdir -p DIRECTORY	Make directories, make parent directories as needed		
cp SOURCEDIRECTORY	Copy files and directories		
cp -r SOURCEDIRECTORY	Copy files and directories, copy directories recursively		
mv SOURCEDIRECTORY	Move (rename) files		
man COMMAND	An interface to the on-line reference manuals		

# Basic Linux commands (2)

Command	Description	
ssh -X foo@host.com	OpenSSH SSH client (remote login program), access to host.com with user foo	
scp foo@host.com:/home/bar ./	Secure copy (remote file copy program), copy file bar from /home on host.com to directory	
scp bar foo@host.com:/home/	Secure copy (remote file copy program), copy file bar from the local host to /home on host.com	
git clone git@github.com:whatever folder-name	The stupid content tracker, Clone a repository (whatever) into a new directory (folder-name).	
git checkout	Checkout a branch or paths to the working tree.	

#### Other clusters — Step-by-Step

 First login, change password and get lecture notes (MS-Windows: Putty, Linux/MacOS: Terminal)

```
> ssh -X ssh YALE_ID@grace.hpc.yale.edu
> passwd #Change password for USERNAME.
  (current) UNIX password:
  Enter new UNIX password:
  Retype new UNIX password:
  Password changed
> git clone ***lecture-folder*** #clone lecture
> cd ***lecture_folder*** #go into folder
> ls # list content of folder
```

#### Step-by-Step (2)

#### → Perform some basic operations on the cluster

```
> ssh -X ss3835@grace.hpc.yale.edu
> pwd
/home/USERNAME
> mkdir -p firstFolder/secondFolder
> 1s
FirstFolder
> ls firstFolder
secondFolder
> cd firstFolder
> pwd
/home/USERNAME/firstFolder
> 1s
secondFolder
> exit
```

#### Step-by-Step (3)

- How to copy folders and files to your PC?
- MS-Windows, start WinSCP
  - → Host-Name: grace.hpc.yale.edu
  - → User: YALE\_ID
- Linux/MacOS, replace /YOUR-LOCAL-PATH/
  - → with /home/LOCAL-LOGIN-NAME/ for linux
  - → with /Users/LOCAL-LOGIN-NAME/ for MacOS

#### Step-by-Step (4)

- Copy folders and files from your notebook create a file named firstFile in firstFolder
  - → MS-Windows: use WinSCP to copy the directory back
  - → Linux/MacOS

```
>scp -r /YOUR-LOCAL_PATH/firstFolder/FILENAME YALE_ID@grace.hpc.yale.edu:
```

Check that file is there by

```
>ssh -X YALE_ID@grace.hpc.yale.edu:
> ls
FILENAME
>cat FILENAME #shows content of file
```

#### **Environment setup**

Supporting diverse user community requires supporting diverse tool sets (different vendors, versions of compilers, debuggers, libraries, apps, etc)

User environments are customized via modules system (or softenv)

- > module avail #shows list of available modules
- > module list #shows list of modules loaded by user
- > module load module\_name #load a module e.g. compiler
- > module unload module\_name #unload a module

### Example – environment setup

```
> vi ~/.bashrc  #here you can setup/store your profile
module load python  #always load this lib upon login
```

#### Using an editor on a cluster

Compute clusters like Grace's infrastructure have a variety of simple text editors available.

#### $\rightarrow$ vi, vim

```
>vi helloworld.cpp
#include <iostream>
int main()
{
  std::cout << "Hello World!" << std::endl;
  return 0;
}</pre>
```

#### More low bandwidth editors

Depending on network and preference, you may want to use an editor without a graphical user interface; common options:

- vi/vim

- emacs

- nano

emacs: Two modes – insertion and command mode

Insertion mode begins upon an insertion

Undo: C- [ESC] returns to command mode

Find/create file: C-x C-f Command mode options:

:w save

Save file: C-x C-s :wq save and exit

:q exit as long as there are no changes

Exit Emacs: C-x C-c :q! exit without saving

Quit: C-g Insertion:

i (insert before cursor)

Deletion: x

a (append)

j (down) I (right)

k (up)

Motion: h (left)

# Compiling & running code interactively

→ go to Root/day1/code/examples → cd Root/day1/codes/examples

If your program is only in one file (a hello-world program, or any simple code that doesn't require external libraries), the compilation is straightforward:

> gfortran helloworld.f90 -o helloworld.exe #Fortran

```
> g++ helloworld.cpp -o helloworld.exe #C++
```

Once you produced the executable, you can run it (serial code) by

```
> ./helloworld.exe
```

> hello

Example: ...

#### Compiling Code with a makefile

In case your program consists of many routines (files), compiling by hand gets very cumbersome

> g++ -o abc abc.cpp a.cpp b.cpp c.cpp

- ightarrow A makefile is just a set of rules to determine which pieces of a large program need to be recompiled, and issues commands to recompile them
- → For large programs, it's usually convenient to keep each program unit in a separate file. Keeping all program units in a single file is impractical because a change to a single subroutine requires recompilation of the entire program, which can be time consuming.
- → When changes are made to some of the source files, only the updated files need to be recompiled, although all relevant files must be linked to create the new executable.

### Compiling Code with a makefile (2)

Basic makefile structure: a list of rules with the following format:

```
target ... : prerequisites ... <TAB> construction-commands
```

A "target" is usually the name of a file that is generated by the program (e.g, executable or object files). It can also be the name of an action to carry out, like "clean".

A "prerequisite" is a file that is used as input to create the target.

```
# makefile : makes the ABC program
abc: a.o b.o c.o ### by typing "make", the makefile generates an executable denotes as "abc"
q++ -o abc a.o b.o c.o
a.o: a.cpp
      g++ -c a.cpp
b.o:b.cpp
      g++ -c b.cpp
c.o: c.cpp
      q++ -c c.cpp
clean: ### by typing "make clean", the executable, the *.mod as well as the *.o files are deleted
      rm *.mod *.o abc
```

### Compiling Code with a makefile (3)

- By default, the first target listed in the file (the executable abc) is the one that will be created when the make command is issued.
- Since abc depends on the files a.o, b.o and c.o, all of the .o files must exist and be up-to-date. make will take care of checking for them and recreating them if necessary. Let's give it a try!
- Makefiles can include comments delimited by hash marks (#).
- A backslash (\) can be used at the end of the line to continue a command to the next physical line.
- The make utility compares the modification time of the target file with the modification times of the prerequisite files.
- Any prerequisite file that has a more recent modification time than its target file forces the target file to be recreated.
  - → A lot more can be done with makefiles (beyond the scope of this lecture)

#### Slurm Workload Manager

http://slurm.schedmd.com/

Simple Linux Utility for Resource Management (SLURM).

Open-source workload manager designed for Linux clusters of all sizes.

#### Provides three key functions:

- 1) It allocates exclusive and/or non-exclusive access to resources (computer nodes) to users for some duration of time so they can perform work.
- 2) It provides a framework for starting, executing, and monitoring work (typically a parallel job) on a set of allocated nodes.
- 3) It arbitrates contention for resources by managing a queue of pending work.
  - > sbatch submit\_helloworld.sh (submit job)
  - > squeue -u NAME (status of job)
  - > scancel JOBID (cancel job)

#### Run an executalbe with "slurm"

https://research.computing.yale.edu/support/hpc/user-guide/slurm

```
#!/bin/bash -I

#SBATCH --ntasks=1 ## how many cpus used here

#SBATCH --time=01:00:00 ## walltime requested

#SBATCH --output=slurm_test.out ## output file

#SBATCH --error=slurm_test.err ## error

### executable
./helloworld.cpp.exe
```

- → Try NOW on grace
- > cd Root/day1/code/examples
- > make -f makefile\_helloworld\_cpp
- > sbatch submit\_grace.sh
- →What is the output? Play with it a bit.

#### Nodes available at Yale

#### Compute

We maintain 5 clusters, with roughly 29,000 cores total. Please click on cluster names for more information. To download a Word document that describes our facilities, equipment, and other resources for HPC and research computing, click here.

Cluster Name	Approx. Core Count	Login Address	Monitor Dashboard	Purpose
Grace	11,000	grace.hpc.yale.edu	cluster.ycrc.yale.edu/grace ♂	general
Farnam	6,300	farnam.hpc.yale.edu	cluster.ycrc.yale.edu/farnam ♂	medical/life science
Omega	6,500	omega.hpc.yale.edu	cluster.ycrc.yale.edu/omega 🗷	highly parallel, tightly coupled
Ruddle	3,600	ruddle.hpc.yale.edu	cluster.ycrc.yale.edu/ruddle 🗷	Yale Center for Genome Analysis ☑
Milgram	1,600	milgram.hpc.yale.edu	n/a	Psychology dept. HIPAA cluster

#### Storage

We maintain several high performance storage systems which amount to about 9 PB total. Listed below are the shared filesystems and the clusters where they are available. We distinguish where clusters store their home directories with an asterisk. Using /home will always refer to the home directory of the cluster you are on.

/gpfs/loomis         2.6 PB         Grace*, Omega*, Farnam           /gpfs/ysm         1.5 PB         Farnam*           /gpfs/ycga         2.0 PB         Ruddle*           /ycga-ba         1.1 PB         Ruddle           /gpfs/milgram         1.1 PB         Milgram*	Filesystem	Size	Mounting Clusters
/gpfs/ycga 2.0 PB Ruddle* /ycga-ba 1.1 PB Ruddle	/gpfs/loomis	2.6 PB	Grace*, Omega*, Farnam
/ycga-ba 1.1 PB Ruddle	/gpfs/ysm	1.5 PB	Farnam*
77-3	/gpfs/ycga	2.0 PB	Ruddle*
/gpfs/milgram 11 PB Milgram*	/ycga-ba	1.1 PB	Ruddle
7gPorting-an	/gpfs/milgram	1.1 PB	Milgram*

### Clone a git repository

```
>ssh -X YALE_ID@grace.hpc.yale.edu
```

- > git clone .... # clone the git repository
- > cd .. # go into the repository
- > Is ... # check that all is there