

Parallel architectures and production environment

Introduction to Parallel Computing with MPI and OpenMP

P. Dagna

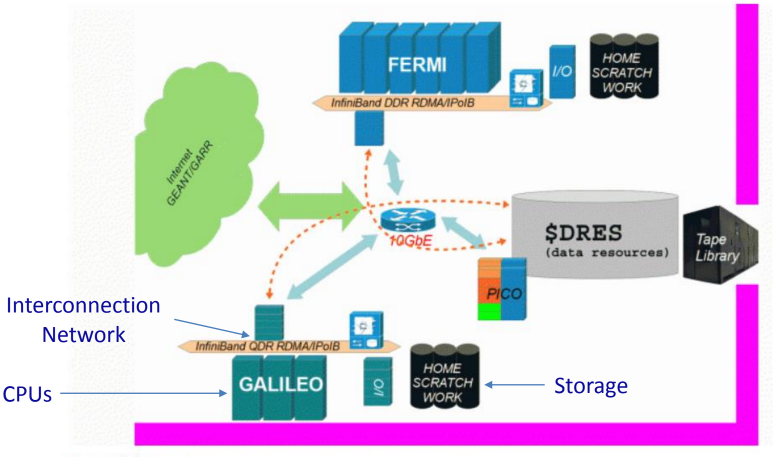
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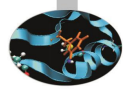




The HPC infrastructure

Cineca HPC infrastructure





The HPC infrastructure



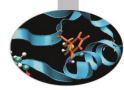
The HPC infrastructure

GALILEO CHARACTERISTICS

Model: IBM NeXtScale Architecture: Linux Infiniband Cluster Processors Type: 8-cores Intel Haswell 2.40 GHz (2 per node) Number of nodes: 516 Compute Number of cores: 8256 Accelerators: 2 Intel Phi 7120p per node on 384 nodes (768 in total) 4 nVIDIA Tesla K40 on 40 nodes (160 in total) RAM: 128 GB/node, 8 GB/core

OS: RedHat CentOS release 7.0, 64 bit









The HPC infrastructure

MARCONI – CHARACTERISTICS

- Model: Lenovo NeXtScale
- Architecture: Intel OmniPath Cluster
- Internal Network: Intel OmniPath
- Disk Space: 17PB (raw) of local storage

A1 Section

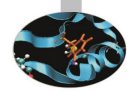
- Nodes: 1.512
- Processors: 2 x 18-cores Intel Xeon E5-2697 v4 (Broadwell) at 2.3 GHz
- Cores: 36 cores/node, 54.432 cores in total
- RAM: 128 GB/node
- Peak Performance: 2 PFlop/s

A2 Section

- Nodes: 3.600
- Processors: 1 x 64-cores Intel Xeon Phi7250
- (Intel Knights Landing) at 1.4 GHz
- Cores: 64 cores/node, 244.800 cores in total
- RAM: 96 GB/node
- Peak Performance: 11 PFlop/s

A3 Section

- Nodes: 1.512
- Processors: 2 x 24-cores Intel Xeon 8160 (Intel SkyLake) at 2.1 GHz
- Cores: 48 cores/node, 72.576 cores in total
- RAM: 192 GB/node
- Peak Performance: 5 PFlop/s

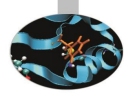




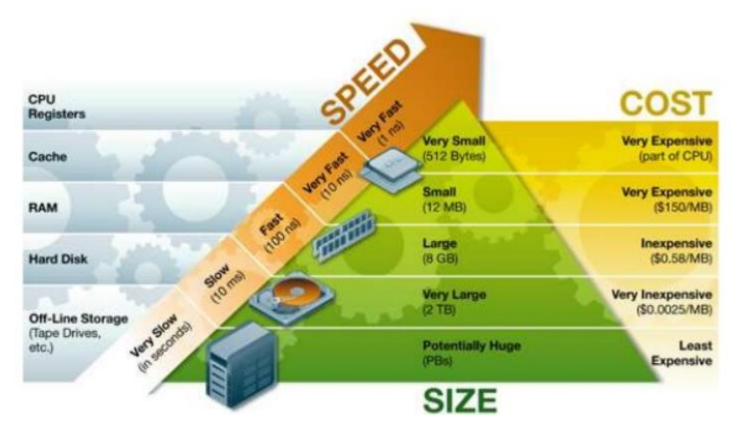




Memory hierarchy



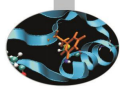
Storage organization and speed access can be thought of as a pyramid



In a shared working environment storage access changes depending to the load of the cluster



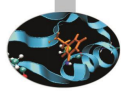




Production Environment







Login and working areas

How to login

- Establish a ssh connection : ssh <username>@login.galileo.cineca.it
- Remarks:
 - ssh available on all linux distros
 - Putty (free) or Tectia ssh on Windows
 - secure shell plugin for Google Chrome!

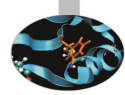
Working Environment

- **\$HOME**:
 - Permanent, backed-up, and local to GALILEO.
 - 50 Gb of quota. For source code or important input files.
- \$CINECA_SCRATCH:
 - Large, parallel filesystem (GPFS).
 - No quota. Run your simulations and calculations here.





Module System



All the optional software on the machine is made available through the "module" system

• provides a way to rationalize software and its environment variables

Modules are divided in profiles

- profile/core (default stable and tested modules)
- profile/extra (more recent versions non completely tested)

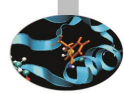
Profiles are divided in 4 categories

- compilers (GNU, intel, openmpi)
- libraries (e.g. LAPACK, BLAS, FFTW, ...)
- tools (e.g. Scalasca, GNU make, VNC, ...)
- applications (software for chemistry, physics, ...)





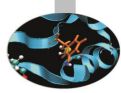
Module System



- CINECA's work environment is organized in modules, a set of installed libraries, tools and applications available for all users.
- "loading" a module means that a series of (useful) shell environment variables will be set
- E.g. after a module is loaded, an environment variable of the form "<MODULENAME>_HOME" is set

/c	developers/devenv/environm developers/devenv/current/	opt/modulefiles/pro				
	levelopers/devenv/current/					
	levelopers/devenv/current/					
	levelopers/devenv/current/					
	levelopers/devenv/current/					
	levelopers/devenv/current/	opt/modulefiles/cor	ce/compil	ers		
intel/cs-xe-201	_3					
[pdagna00@node1	.71 ~]\$ module load intel/	'cs-xe-2013				
	.71 ~]\$ ls \$INTEL_HOME					
		inspector_xe	lib		vtune_amplifier_x	
advisor_xe_2013				server_2017.lic		e_2013 NECA
oin		ipp	man	server.lic		
	include		mkl	tbb		





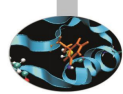
Module Commands

COMMAND	DESCRIPTION		
module av	list all the available modules		
module load <module_name(s)></module_name(s)>	load module <module_name></module_name>		
module list	list currently loaded modules		
module purge	unload all the loaded modules		
module unload <module_name></module_name>	unload module <module_name></module_name>		
module help <module_name></module_name>	print out the help (hints)		
module show <module_name></module_name>	print the env. variables set when loading the module		





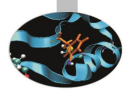
Launching Jobs



- As in every HPC cluster, users are allowed to run their own simulations by submitting "jobs" to the **compute nodes**.
- The job is then taken in consideration by a scheduler, that adds it to a queuing line and allows its execution when the resources required are available.
- The operative scheduler in GALILEO is **PBS**.
- The scheduler has a proprietary scripting language necessary to submit jobs





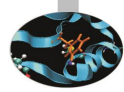


• The scheme of a PBS job script is as follows:

#!/bin/bash
#PBS keywords
variables environment
execution line





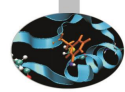


• Example of parallel job:

#!/bin/bash
#PBS -N <jobname>
#PBS -o job.out
#PBS -o job.err
#PBS -e job.err
#PBS -l walltime=1:00:00
#PBS -l select=2:ncpus=16:mpiprocs=16:mem=10GB
#PBS -q <queuename>
cd \$PBS_O_WORKDIR # points to the folder you are actually working
into
module load autoload openmpi
mpirun ./myprogram







• PBS Keyword Analysis:

#PBS -N myname

• Defines the name of your job

#PBS -o job.out

 Specifies the file where the standard output is directed (default=jobname.o<jobID>)

#PBS -e job.err

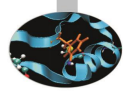
 Specifies the file where the standard error is directed (default=jobname.e<jobID>)

#PBS -I walltime=00:30:00

• Specifies the maximum duration of the job (queue dependency).







PBS Keyword Analysis:

#PBS -l select=2:ncpus=16:mpiprocs=16:mem=10GB

- Specifies the resources needed for the simulation.
 - select number of compute nodes ("chunks")
 - ncpus number of cpus per node (max. 16)
 - mpiprocs number of MPI tasks per node (max=ncpus)
 - mem memory allocated for each node (default=8GB, max.=120 GB)

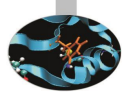
#PBS -q <queuename>

• Specifies the queue requested for the job.





PBS Commands



qsub <job_script>

 Your job will be submitted to the PBS scheduler and executed when there will be nodes available (according to your priority and the queue you requested)

qstat -u <username>

• Shows the list of all your scheduled jobs, along with their status(idle, running, closing, ...) Also, shows you the job id required for other PBS commands.

qstat -f <job_id>

• Provides a long list of information for the job requested. In particular, if your job isn't running yet, you'll be notified about its estimated start time or, if you made an error on the job script, you will learn that the job won't ever start

qdel <job_id>

• Removes the job from the scheduled jobs by killing it

