

UPPMAX Introduction

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SciLifeLab







Enabler for Life Sciences



UPPMAX

- Uppsala Multidisciplinary Center for Advanced Computational Science
- http://www.uppmax.uu.se
- 2 clusters
 - Tintin, 160 computer à 16 cores (64GB RAM)
 - Milou, 208 computers à 16 cores (128GB RAM)

• 17 with 256, 17 with 512

- 1 high memory machine à 64 cores, 2TB RAM
- ~7 PB fast parallel storage





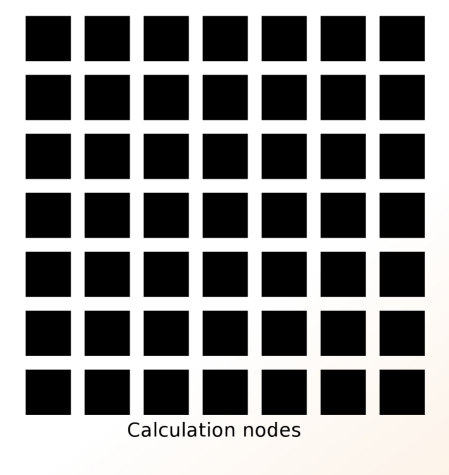
The basic structure of supercomputer







The basic structure of supercomputer

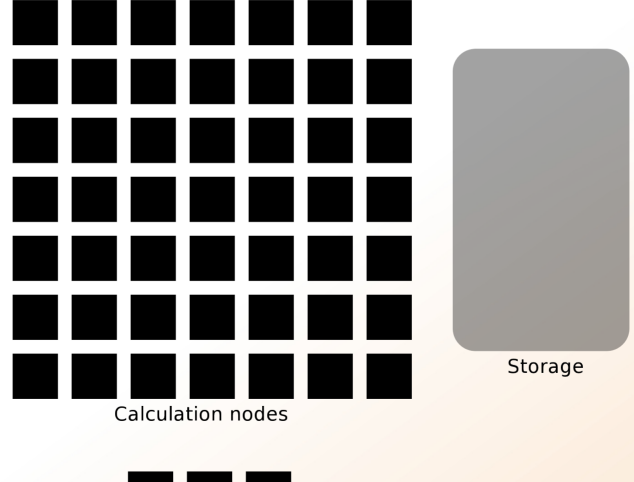








The basic structure of supercomputer



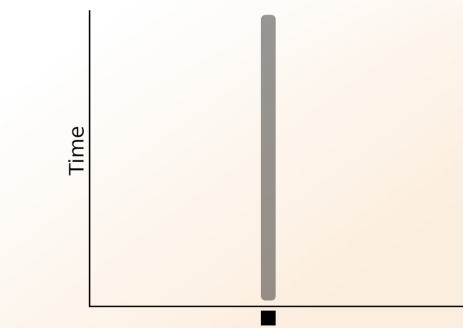






The basic structure of a supercomputer

- Parallel computing is key
 - Not one super fast

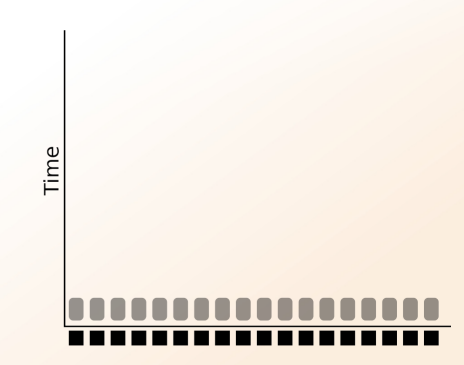






The basic structure of a supercomputer

- Parallel computing is key
 - Not one super fast



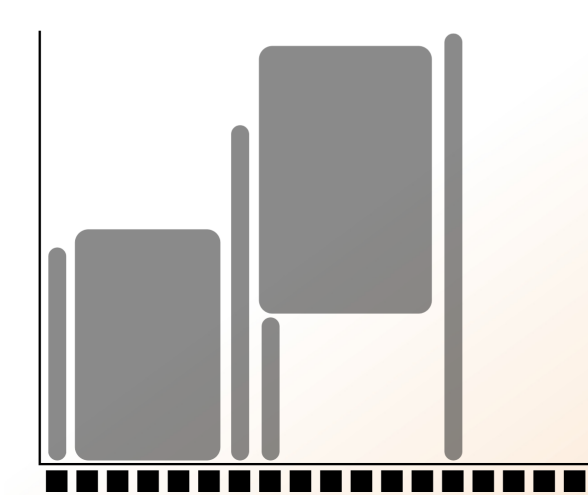


- More users than nodes
 - Need for a queue



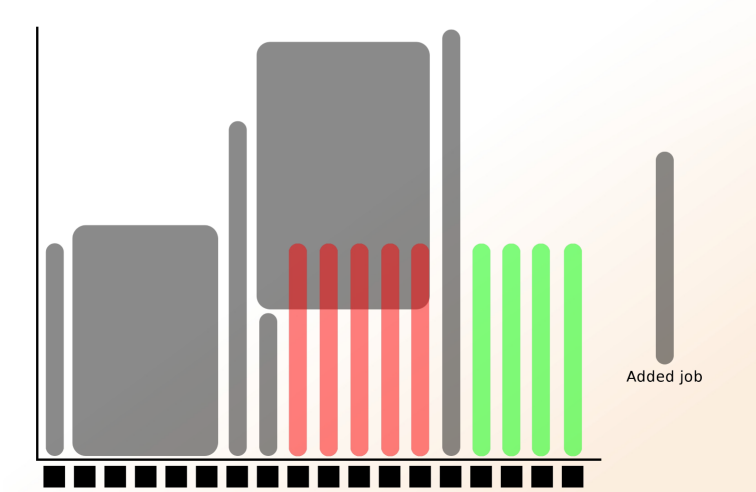


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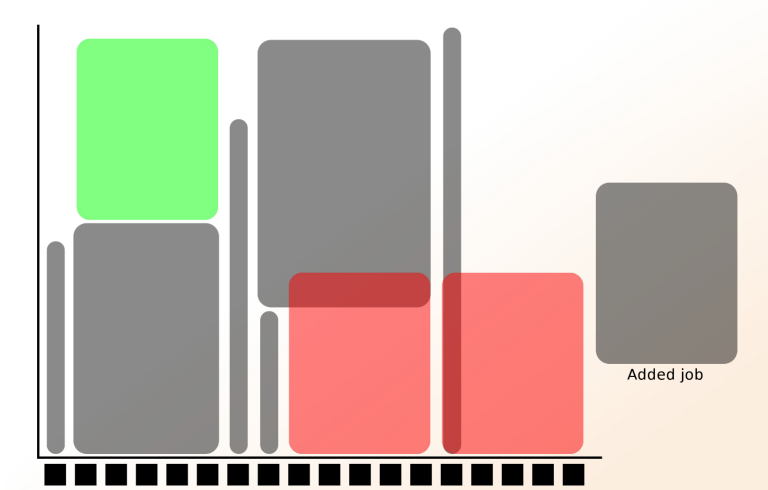


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 - Need for a queue





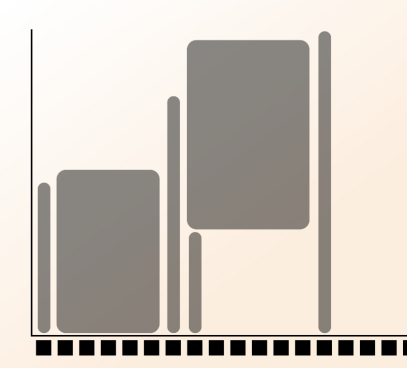
- 1 mandatory setting for jobs:
 - Who pays for it? (-A)
- 3 settings you really should set:

(-n))

(-t)

(default values NOT good)

- Where should it run? (-p)
- (How wide is it?)
- How long is it?





- Who pays for it? (-A)
 - Only projects can be charged
 - You have to be a member
 - This course's project ID: g2015006

- -A = account (the account you charge)
 - No default value, mandatory



- Where should it run? (-p)
 - Use a whole node or just part of it?
 - 1 node = 16 cores
 - I hour walltime = 16 core hours = expensive
 - Waste of resouces
 - Default project size: 2000 core hours
- -p = partition (node or core)
 - Default value: core

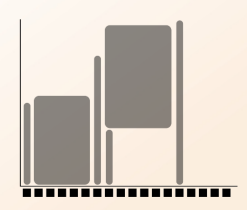


- How wide is it? (-n)
 - How much of the node should be booked?
 - 1 node = 16 cores
 - Any number of cores
 - 1, 2, 5, 13, 15 etc

- -n = number of cores
 - Default value: 1
 - Usually used together with -p core



- How long is it? (-t)
 - Always overestimate ~50%
 - Jobs killed when timelimit reached
 - Only charged for time used
- -t = time (hh:mm:ss)
 - 78:00:00 or 3-6:00:00
 - Default value: 7-00:00:00

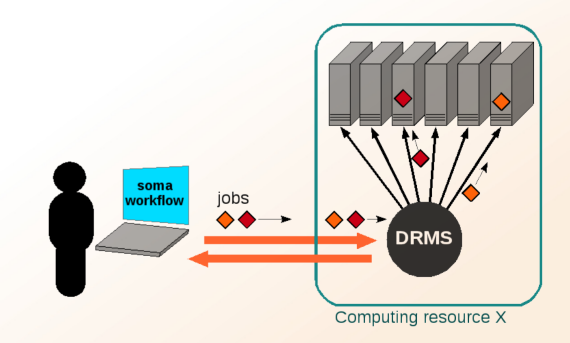






Job = what happen during the booked time

- Script file
 - Start programs
 - Move files
 - and more





- How to submit a job
 - Write a script (bash)
 - Queue options
 - Rest of the script

```
#! /bin/bash -l
#SBATCH -A g2012157
#SBATCH -p core
#SBATCH -J Template_script
#SBATCH -t 08:00:00
# go to some directory
cd ~/glob
# do something
echo Hello world!
```



- How to submit a job
 - Write a script (bash)
 - Queue options
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```
#! /bin/bash -l
#SBATCH -A g2012157
#SBATCH -p node
#SBATCH -J Template_script
#SBATCH -t 08:00:00
```

go to the correct directory
cd /home/dahlo/glob/work/uppmaxScripts/misc

run tophat on the data, using 8 cores
tophat -p 8 /bubo/proj/g2012157/indexes/bowtie/hg19 tophat/input/ad12.fq



How to submit a job

Script written, now what?

```
[dahlo@kalkyl1 temp]$ ls -l
total 16
-rw-r--r-- 1 dahlo uppmax 169 Jan 28 15:45 test.sbatch
[dahlo@kalkyl1 temp]$ cat test.sbatch
#! /bin/bash -l
#SBATCH -A q2012157
#SBATCH -p core
#SBATCH -J Template script
#SBATCH -t 08:00:00
# go to some directory
cd ~/glob
# do something
echo "Hello world!"
[dahlo@kalkyl1 temp]$ sbatch test.sbatch
Submitted batch job 1745244
[dahlo@kalkyl1 temp]$
```



SLURM Output

- Prints to a file instead of terminal
 - slurm-<job id>.out

```
[dahlo@biologin glob]$ ll
total 16
-rw-r--r-- 1 dahlo uppmax 1025 Sep 22 19:46 my_script.sb
[dahlo@biologin glob]$
```



SLURM Output

- Prints to a file instead of terminal
 - slurm-<job id>.out

```
[dahlo@biologin glob]$ ll
total 16
-rw-r--r-- 1 dahlo uppmax 1025 Sep 22 19:46 my_script.sb
[dahlo@biologin glob]$
[dahlo@biologin glob]$ sbatch my_script.sb
Submitted batch job 2226951
[dahlo@biologin glob]$
[dahlo@biologin glob]$ ll
total 32
-rw-r--r-- 1 dahlo uppmax 1025 Sep 22 19:46 my_script.sb
-rw-r--r-- 1 dahlo uppmax 87 Sep 22 19:56 slurm-2226951.out
[dahlo@biologin glob]$
```



SLURM Output

Prints to a file instead of terminal

slurm-<job id>.out

```
[dahlo@biologin glob]$ ll
total 16
-rw-r--r-- 1 dahlo uppmax 1025 Sep 22 19:46 my script.sb
[dahlo@biologin glob]$
[dahlo@biologin glob]$ sbatch my script.sb
Submitted batch job 2226951
[dahlo@biologin glob]$
[dahlo@biologin glob]$ ll
total 32
-rw-r--r-- 1 dahlo uppmax 1025 Sep 22 19:46 my script.sb
-rw-r--r-- 1 dahlo uppmax 87 Sep 22 19:56 slurm-2226951.out
[dahlo@biologin glob]$
[dahlo@biologin glob]$ cat slurm-2226951.out
Example of error with line number and message
slurm script: 40: An error has occurred.
[dahlo@biologin glob]$
```





Shows information about your jobs

squeue -u <user>

[dahlo@kalkyl4 work]\$ squeue -u dahlo JOBID PARTITION NAME USER ST 2215978 core My_job dahlo PD [dahlo@kalkyl4 work]\$

TIME	NODES	NODELIST(REASON)
0:00	1	(Resources)





Shows information about your jobs

squeue -u <user>

[dahlo@kalkyl4 work]\$ squeue -u dahlo JOBID PARTITION NAME USER ST 2215978 core My job dahlo R [dahlo@kalkyl4 work]\$

TIME	NODES	NODELIST(REASON)
0:13	1	q148





Shows information about your jobs

squeue -u <user>

[dahlo@kalkyl4 work]\$ squeue -u dahlo JOBID PARTITION NAME USER ST 2215978 core My job dahlo R [dahlo@kalkyl4 work]\$

TIME	NODES	NODELIST(REASON)
0:13	1	q148

jobinfo -u <user>



SSH

Connects to a node

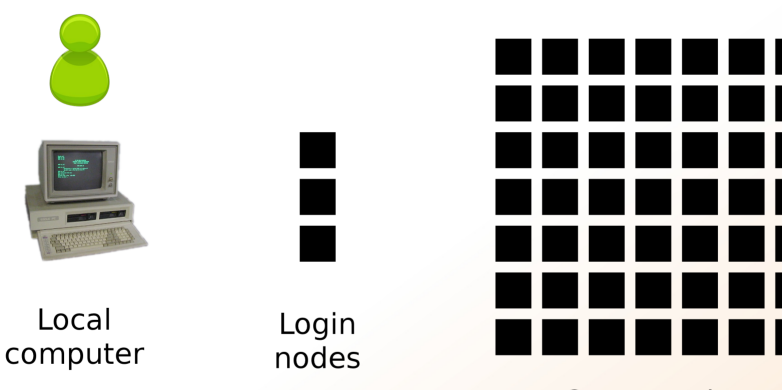
ssh -X <node name>

[dahlo@kalkyl4 work]\$ squeue -u dahlo JOBID PARTITION NAME USER ST 2215978 core My_job dahlo R [dahlo@kalkyl4 work]\$ [dahlo@kalkyl4 work]\$ ssh -X q148 [dahlo@q148 work]\$

TIME	NODES	NODELIST(REASON)
0:13	1	q148



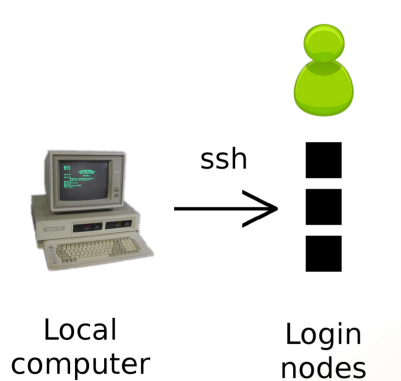
SSH

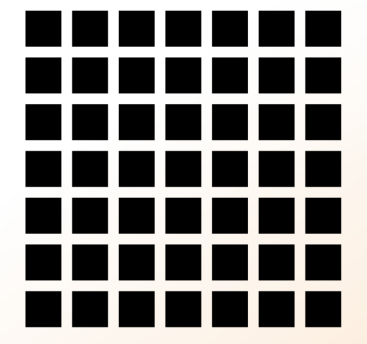


Computation nodes



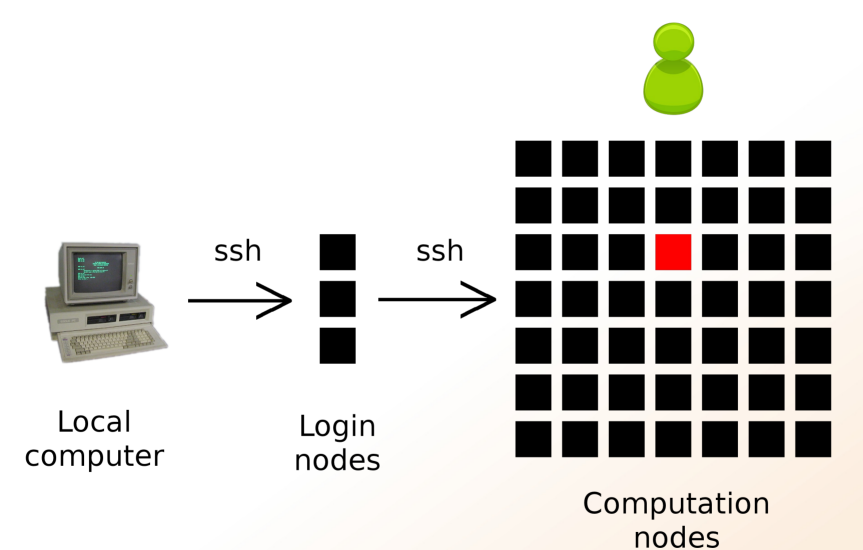
SSH





Computation nodes







Books a node and connects you to it

interactive -A <proj id> -p <core or node> -t <time>

[dahlo@kalkyl3 work]\$ interactive -A g2012205 -t 00:15:00 -p core



Books a node and connects you to it

interactive -A <proj id> -p <core or node> -t <time>

[dahlo@kalkyl3 work]\$ interactive -A g2012205 -t 00:15:00 -p core Your job is assigned a high interactive priority.

Please note that you must not use more than three GB of memory. Waiting for job 2216477 to start... Starting job now -- you waited for 3 seconds.



Books a node and connects you to it

interactive -A <proj id> -p <core or node> -t <time>

[dahlo@q229 work]\$



Books a node and connects you to it

interactive -A <proj id> -p <core or node> -t <time>

[dahlo@q229 work]\$

 Add -n to get more cores if you need more memory



UPPMAX Software

- 100+ programs installed
- Managed by a 'module system'
 - Installed, but hidden
 - Manually loaded before use

module avail - Lists all available modules

module load <module name> - Loads the module

module unload <module name> - Unloads the module

module list - Lists loaded modules



UPPMAX Software

- Most bioinfo programs hidden under bioinfo-tools
 - Load bioinfo-tools first, then program module

[dahlo@kalkyl3 work]\$ module load cufflinks/1.2.1 ModuleCmd_Load.c(200):ERROR:105: Unable to locate a modulefile for 'cufflinks/1.2.1' [dahlo@kalkyl3 work]\$ module load bioinfo-tools [dahlo@kalkyl3 work]\$ module load cufflinks/1.2.1 [dahlo@kalkyl3 work]\$

or

[dahlo@kalkyl3 work]\$ module load samtools ModuleCmd_Load.c(200):ERROR:105: Unable to locate a modulefile for 'samtools' [dahlo@kalkyl3 work]\$ module load bioinfo-tools samtools [dahlo@kalkyl3 work]\$ [dahlo@kalkyl4 work]\$ module load bioinfo-tools
[dahlo@kalkyl4 work]\$ module avail

	/bubo/sw/r	nf/kalkvl/bioinfo-too	ls/alignment
MUMmer/3.22(default)			maq/0.7.1(default)
anfo/0.97	blast/2.2.2		mosaik-aligner/1.0.1388(default)
anfo/0.98(default)	blast/2.2.2		mosaik-aligner/1.1.0021
blast/2.2.15	blat/34		<pre>mpiblast/1.6.0(default)</pre>
blast/2.2.18	bwa/0.5.8a		splitseek/1.3.2
blast/2.2.23	bwa/0.5.9		splitseek/1.3.4(default)
blast/2.2.23+	hmmer/3.0		
	(hubo (cu/i	mf/kalkul/higinfo_tool	ls (assembly
Ray/0.0.4	abyss/1.2.4	abyss/1.3.0	ls/assemblyvelvet/1.0.03(default)
			velvet/1.1.04
Ray/0.0.7(default) Ray/1.6.1	abyss/1.2.5(derautt)		
abyss/1.2.3	abyss/1.2.7 abyss/1.2.7-maxk96	mira/3.0.0 mira/3.2.0(default)	velvet/1.1.04_KI01
aby 55/1.2.5	aby55/1.2.7-11axx50	mild/5.2.0(delautt)	Ve(Ve()1.1.0/
	/bubo/sv	w/mf/kalkyl/bioinfo-to	ools/misc
BclConverter/1.7.1			samtools/0.1.12-10(default)
BioPerl/1.6.1	freebayes/		samtools/0.1.16
BioPerl/1.6.1_PERL5.10).1(default) gcta/0.92.0	Ð	samtools/0.1.18
BioPerl/1.6.1_PERL5.12			samtools/0.1.7a
FastQC/0.6.1	2	s/1.3.5(default)	samtools/0.1.8
FastQC/0.7.2(default)			samtools/0.1.9
Fastx/0.0.13(default)	htseq/0.5.	1	snpEff/2.0.3
IGV/1.5.51	matrix2png,		trinity/2011-05-13
biopython/1.56	picard/1.4		trinity/2011-10-29
cellprofiler/20111024		1	
emmax/beta-07Mar2010	plink/1.07		
	/huho/cu//	mf/kalkyl/biainfa_taal	ls (phylogopy
concaterpillar/1.4 garli/2.0 raxml/7.0.4(default) raxml/7.2.8			
	mrbayes/3.1.2-mpi		Taxint/7.2.0
ga. cz, 0.0000 (deradec)	mages, state mpr		
pipelines			
	cufflinks/0.9.2		
<pre>bowtie/0.12.6(default)</pre>	cufflinks/0.9.3	cufflinks/1.2.1	tophat/1.3.3



UPPMAX Commands

uquota

[dahlo@biologin work]\$ uquota Your File Area Usage (GB) Quota Limit (GB) Over Quota Grace	Time
dahlo glob 196 2048	-
dahlo home 4 32	-
/proj/b2010015 229 256	
/proj/b2010015/nobackup 0 512	-
/proj/b2010033 132 6348	
/proj/b2010033/nobackup 27 512	-



projinfo

UPPMAX Commands

[dahlo@kalkyl4 work]\$ projinfo
(Counting the number of core hours used since 2012-08-19/00:00:00 until now.)

Project User	Used[h]	Current allocation [h/month]
b2010015 ameur	1257.20 1257.20	2000
b2010069	0.00	2000
b2010074 dahlo seba	110.98 1.01 109.97	2000
b2012044	0.00	2000
g2012005	0.00	2000
g2012083	0.00	2000
g2012157 dahlo	0.12 0.12	2000

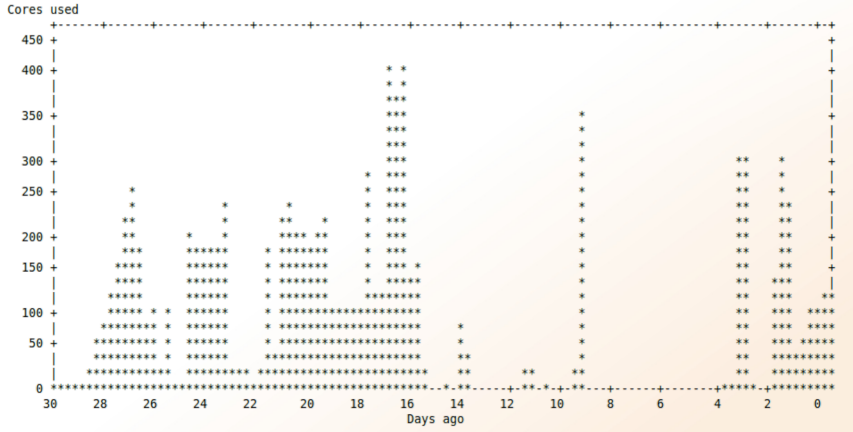
[dahlo@kalkyl4 work]\$



UPPMAX Commands

projplot -A <proj-id> (-h for more options)

Core hour usage during the last 30 days Project: a2009002 Cluster: kalkyl Core hours used in interval: ~29 173 (72.93%) Limit: 40 000



[dahlo@biologin slurm-usage]\$



UPPMAX

Summary

- All jobs are run on nodes through queue system
- A job script usually consists of
 - Job settings (-A, -p, -n, -t)
 - Modules to be loaded
 - Bash code to perform actions
 - Run a program, or multiple programs

More info on UPPMAX homepage

http://www.uppmax.uu.se/milou-user-guide



• Laboratory time! (again)