

UPPMAX Introduction

2017-05-15

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Enabler for Life Science











Uppsala Multidisciplinary Center for Advanced Computational Science

http://www.uppmax.uu.se

3 clusters

Rackham, 334 computer à 20 cores (128GB RAM)
Milou, 208 computers à 16 cores (128GB RAM)
17 with 256, 17 with 512
Bianca, 200 nodes à 16 cores (128GB RAM)

~11 PB fast parallel storage

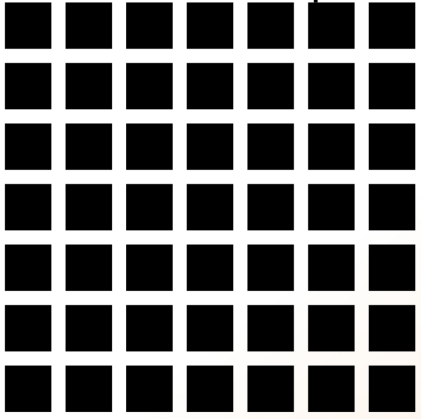


The basic structure of supercomputer





The basic structure of supercomputer

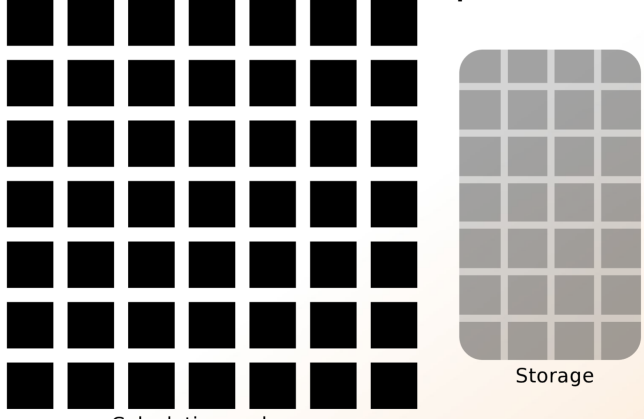


Calculation nodes





The basic structure of supercomputer



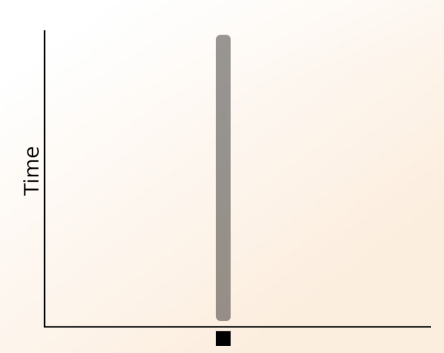






The basic structure of a supercomputer

Parallel computing is key Not one super fast





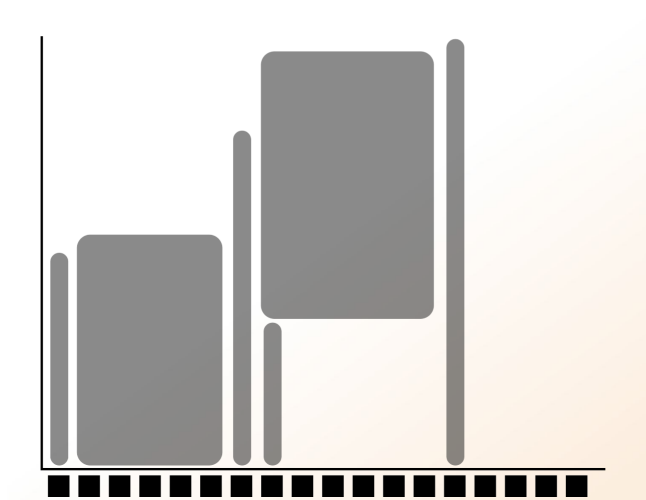
The basic structure of a supercomputer

Parallel computing is key Not one super fast

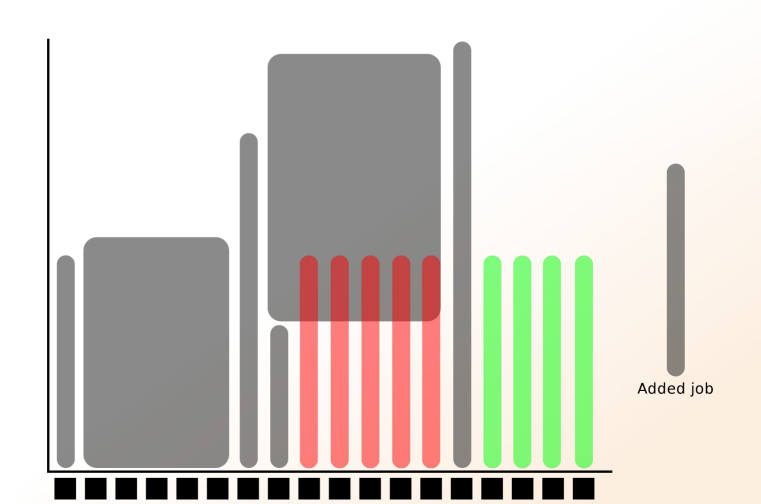




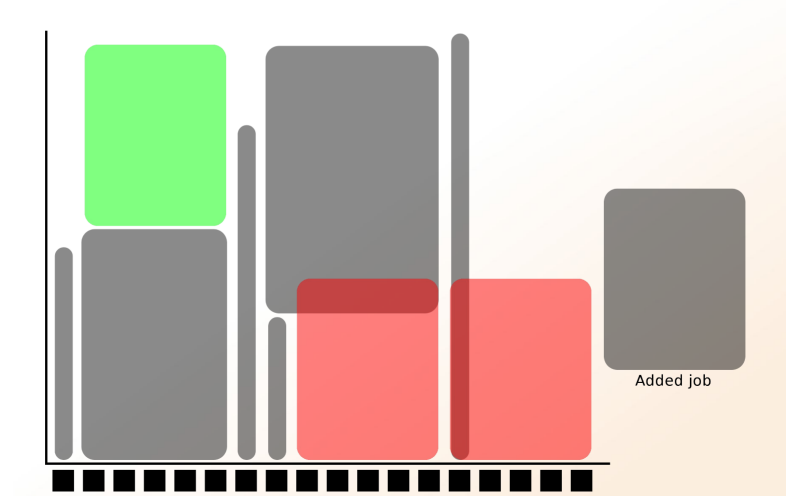














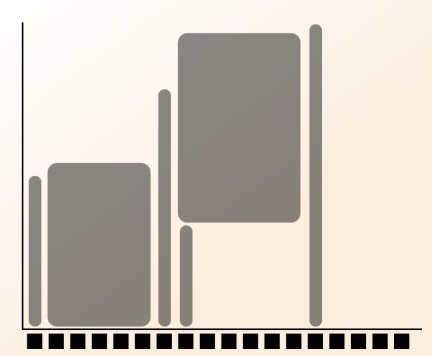
1 mandatory setting for jobs: Who pays for it? (-A)

3 settings you really should set: (default values NOT good)

Where should it run? (-p)

(How wide is it? (-n))

How long is it? (-t)





Who pays for it? (-A)
Only projects can be charged
You have to be a member

This course's project ID: g2017014

-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

1 hour walltime = 16 core hours = expensive

Waste of resources

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 coresDefault value: 1Usually used together with -p core

-N = number of nodes

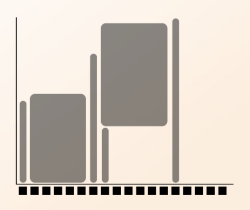


How long is it? (-t)
Always overestimate ~50%
Jobs killed when timelimit reached
Only charged for time used

-t = time (hh:mm:ss)

78:00:00 or 3-6:00:00

Default value: 7-00:00:00

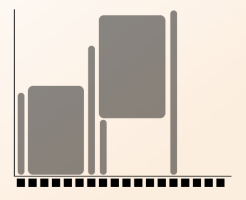




Slurm user guide at

http://www.uppmax.uu.se/support/user-guides/slurm-user-guide/#tocjump_8905574359956505_5

(just google "uppmax slurm user guide")

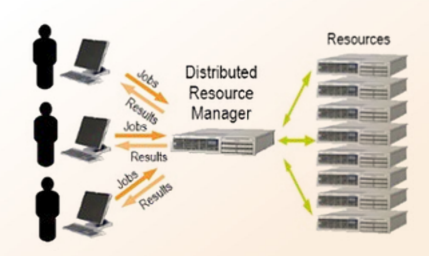




A job?

Job = what happen during the booked time Script file

Start programs
Move files
and more





How to submit a job Write a script (bash)

```
#! /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!
```

Queue options

Rest of the script



How to submit a job Write a script (bash)

```
#! /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]$
```



Squeue

Shows information about your jobs

squeue -u <user>

```
[dahlo@kalkyl4 work]$ squeue -u dahlo

JOBID PARTITION NAME USER ST TIME NODES NODELIST(REASON)

2215978 core My_job dahlo PD 0:00 1 (Resources)

[dahlo@kalkyl4 work]$
```



Squeue

Shows information about your jobs

squeue -u <user>

```
[dahlo@kalkyl4 work]$ squeue -u dahlo

JOBID PARTITION NAME USER ST TIME NODES NODELIST(REASON)

2215978 core My job dahlo R 0:13 1 q148

[dahlo@kalkyl4 work]$
```



Squeue

Shows information about your jobs

squeue -u <user>

```
[dahlo@kalkyl4 work]$ squeue -u dahlo

JOBID PARTITION NAME USER ST TIME NODES NODELIST(REASON)

2215978 core My job dahlo R 0:13 1 q148

[dahlo@kalkyl4 work]$
```

jobinfo -u <user>





Connects to a node

ssh -Y <node name>

```
[dahlo@kalkyl4 work]$ squeue -u dahlo
JOBID PARTITION NAME USER ST TIME NODES NODELIST(REASON)
2215978 core My_job dahlo R 0:13 1 q148
[dahlo@kalkyl4 work]$
[dahlo@kalkyl4 work]$ ssh -Y q148
[dahlo@q148 work]$
```

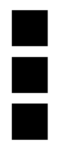


SSH

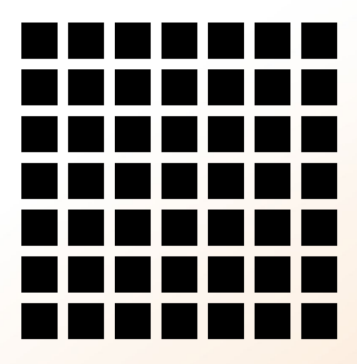




Local computer



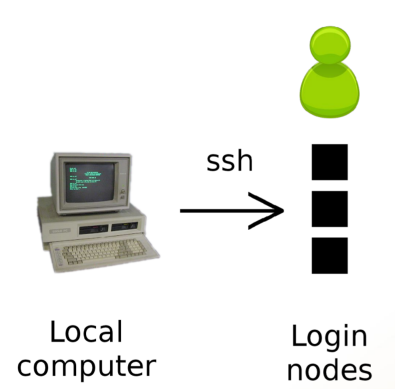
Login nodes

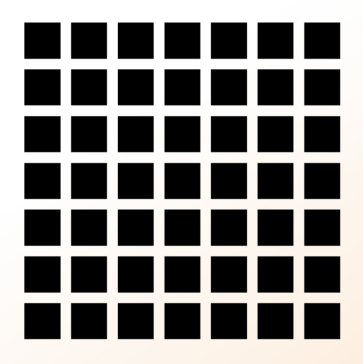


Computation nodes



SSH

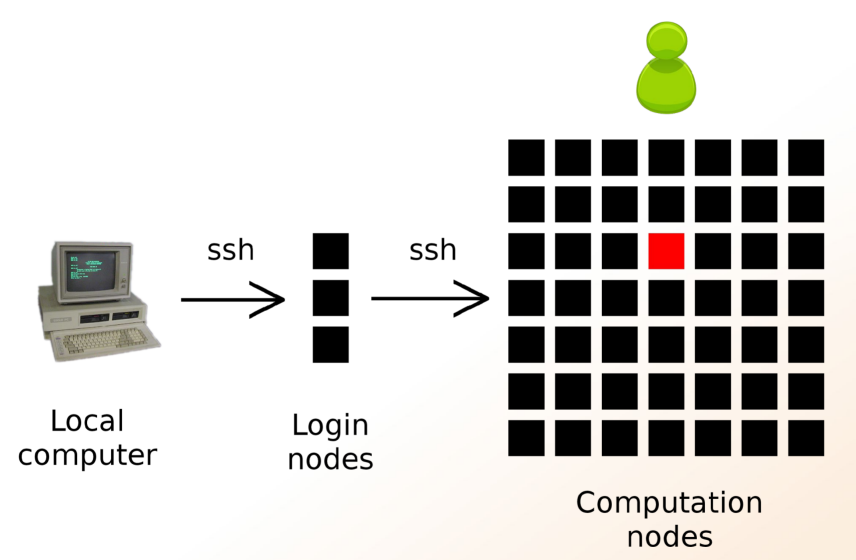




Computation nodes



SSH





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 module spider <word> - Searches all modules after 'word'



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@kalkvl4 work]$ module avail
                            -- /bubo/sw/mf/kalkyl/bioinfo-tools/alignment ------
MUMmer/3.22(default)
                             blast/2.2.24(default)
                                                          mag/0.7.1(default)
anfo/0.97
                                                          mosaik-aligner/1.0.1388(default)
                             blast/2.2.24+
anfo/0.98(default)
                             blast/2.2.25
                                                          mosaik-aligner/1.1.0021
                                                          mpiblast/1.6.0(default)
blast/2.2.15
                             blat/34
blast/2.2.18
                             bwa/0.5.8a
                                                          splitseek/1.3.2
                                                          splitseek/1.3.4(default)
blast/2.2.23
                             bwa/0.5.9
blast/2.2.23+
                             hmmer/3.0
-----/bubo/sw/mf/kalkyl/bioinfo-tools/assembly ------
                               abyss/1.3.0
                                                           velvet/1.0.03(default)
Ray/0.0.4 abyss/1.2.4
Ray/0.0.7(default) abyss/1.2.5(default) abyss/1.3.2 velvet/1.1.04
Ray/1.6.1
                    abyss/1.2.7
                                        mira/3.0.0
                                                             velvet/1.1.04 K101
                                        mira/3.2.0(default) velvet/1.1.07
                    abyss/1.2.7-maxk96
abyss/1.2.3
-----/bubo/sw/mf/kalkyl/bioinfo-tools/misc ------------------------
BclConverter/1.7.1
                              freebayes/0.8.9
                                                            samtools/0.1.12-10(default)
                                                            samtools/0.1.16
                              freebayes/0.9.4
BioPerl/1.6.1
BioPerl/1.6.1 PERL5.10.1(default) gcta/0.92.0
                                                            samtools/0.1.18
BioPerl/1.6.1 PERL5.12.3
                              gcta/0.92.6
                                                            samtools/0.1.7a
Fast0C/0.6.1
                              genometools/1.3.5(default)
                                                            samtools/0.1.8
FastQC/0.7.2(default)
                              htseq/0.4.6
                                                            samtools/0.1.9
                                                            snpEff/2.0.3
Fastx/0.0.13(default)
                              htseq/0.5.1
IGV/1.5.51
                              matrix2png/1.2.1
                                                            trinity/2011-05-13
biopython/1.56
                              picard/1.40
                                                            trinity/2011-10-29
cellprofiler/20111024
                              picard/1.41
emmax/beta-07Mar2010
                              plink/1.07
      ------/bubo/sw/mf/kalkyl/bioinfo-tools/phylogeny -------/bubo/sw/mf/kalkyl/bioinfo-tools/phylogeny
concaterpillar/1.4 garli/2.0
                                       raxml/7.0.4(default) raxml/7.2.8
garli/0.96b8(default) mrbayes/3.1.2-mpi raxml/7.2.7
```



uquota

Usage (GB)	Quota Limit (GB)	Over Quota	Grace Time
196	2048		-
4	32		-
229	256		
Θ	512		-
132	6348		
27	512		-
	4 229 0 132	196 2048 4 32 229 256 0 512 132 6348	196 2048 4 32 229 256 0 512 132 6348



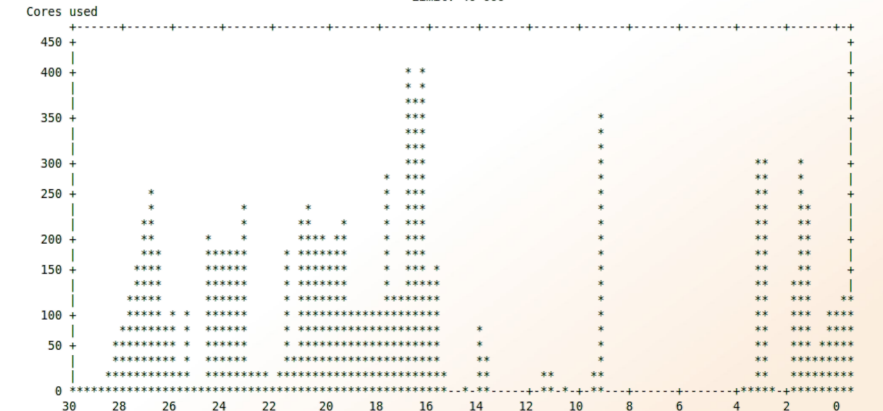
projinfo

1257.20 b2010015 2000 1257.20 ameur b2010069 0.00 2000 b2010074 110.98 2000 dahlo 1.01 seba 109.97 b2012044 0.00 2000 g2012005 0.00 2000 g2012083 0.00 2000 q2012157 0.12 2000 dahlo 0.12

[dahlo@kalkyl4 work]\$



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



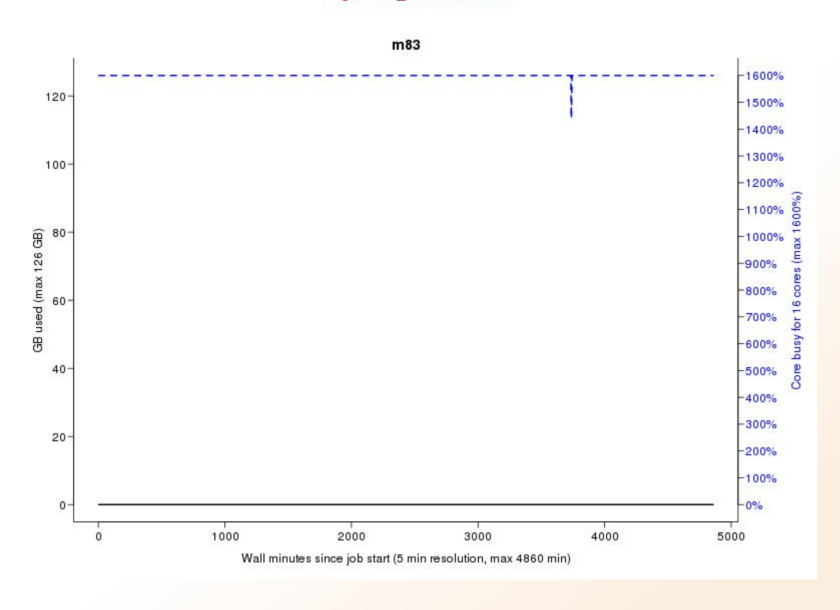
Days ago



Plot efficiency jobstats -p -A projid>

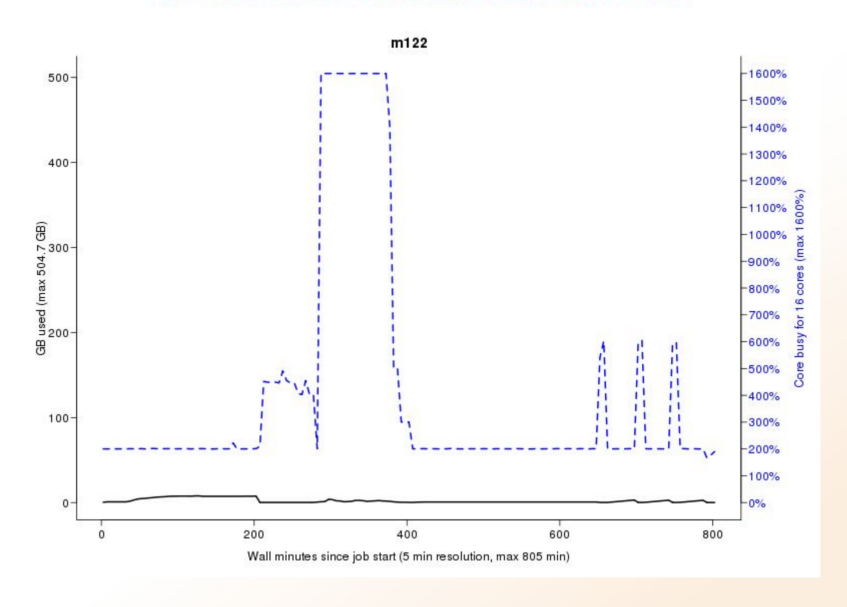
2719328 on 'milou' end: 2014-09-09T08:26:34 runtime: 03-08:59:53

Flags: mem_underused:126:0



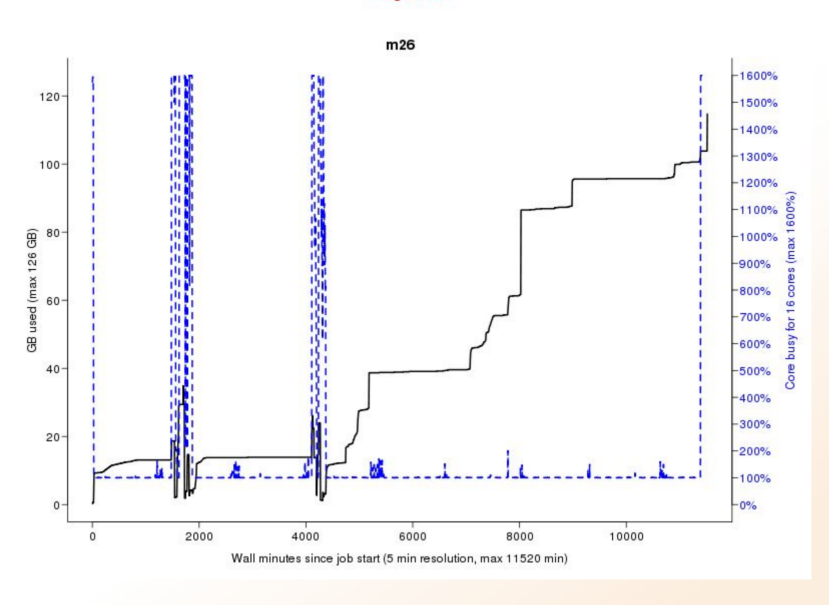
2934193 on 'milou' end: 2014-09-26T01:40:23 runtime: 13:30:23

Flags: mem_underused:504.7:7.9 node_type_misbooked:mem512GB:mem128GB



2799665 on 'milou' end: 2014-09-18T07:36:54 runtime: 07-23:56:23







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)