Running code on the Wesleyan Cluster

ABSTRACT

This document is intended as a quick reference guide for running code on the Wesleyan computing cluster. It is not an introduction to high-performance cluster computing (I would not be qualified to write such a document) but it does cover the details of computing power available, and how to get started using it.

1. CLUSTER BASICS

The Wesleyan computing center (located on the 5th floor of the Excley Science Tower) contains a range of computing capabilities, all available through the Wesleyan network. These capabilities include highly parallelized computing, GPU computing, high memory (~100 GB) computing, etc. A full list of the hardware setup is available online¹, along with a list of available software². A full list of all information about the cluster is also available online³. The cluster is managed by Henk Meij (hmeij@wesleyan.edu); I have always found Henk to be very responsive and helpful when setting up my code, and running through any sort of trouble shooting that I can't handle on my own. The first step in running code on the cluster is to contact Henk, and to request an account on the cluster.

The first point of contact with the computing cluster is through one of the login nodes. The primary login node is cottontail, although secondary nodes cottontail2 and swallowtail are also available. These login nodes can be accessed through ssh

ssh -XY kflaherty@cottontail

You will be deposited into your home directory (/home/kflaherty) on the computing cluster, where you will store your code and data. Note that this home directory is not the same as the home directory on you local machine and your data and code must be copied over from your local machine to the computing clusters home directory

¹ https://dokuwiki.wesleyan.edu/doku.php?id=cluster:126

² https://dokuwiki.wesleyan.edu/doku.php?id=cluster:73

³ https://dokuwiki.wesleyan.edu/doku.php?id=cluster:0

scp mydata.fits kflaherty@cottontail:/home/kflaherty/mycode/ scp mycode.py kflaherty@cottontail:/home/kflaherty/mycode/

There is a total of 10 TB of space allocated for home directories; if more space is needed for your data, contact Henk to make arrangements.

After connecting to the login node, you can access the computing nodes through the use of the OpenLava Scheduler. A job, which is one instance of running your code, is submitted to the computing cluster using the scheduler, which in turn distributes the computational work according to the requested configuration (e.g. the requested computing queue, the requested number of cores, etc.) onto any available cores. The computing cluster is set up with a number of different queues, which provide access to a collection of nodes, which are many up of multiple *chips*. The number of nodes, and the number of *chips* per node depends on the chosen queue. The exact queue that works best for you will depend on the computing capabilities needed for your code. The nodes are specified by n followed by a number (e.g. n33 n14 n27). When multiple cores from a single node are used by a piece of code, you will see the node name repeated multiple times (e.g. n33 n33 n33 n33 n33 for 5 cores from node 33)

The main queues available on the Wesleyan computing cluster are:

- 1. hp12: The default queue. Consists of 32 nodes, with dual quad core chips with 12 GB of memory each
- 2. mw256fd: Contains 8 nodes with dual eight core chips, with 256 GB of memory each.
- 3. mwgpu: Contains four GPUs, 2,500 cores/GPU with GPU memory footprint of 5 GB.

The queue hp12 is designed for highly parallelized, but small memory, jobs (tinymem is a queue that also handles small memory jobs). The queue mw256fd, the primary queue that I use, is designed for large memory jobs, while mwgpu is designed for GPU computations. All queues have access to the same software base, and are all accessible from any login node. As described below your choice of queue is specified when submitting a job to the scheduler.

2. SCRIPTS FOR RUNNING JOBS

The submit a job to the desired queue, a submission script must be created. This script contains a series of commands that setup the computing job, and then call you code to start the computation. Outlined below are the basic commands used in the submission script for either a serial or parallel job. Examples of these pieces of code are available in Henk's home directory (/home/hmeij/jobs within the serial and parallel) directories. Henk has also implemented checkpointing, which allows for a job to be restarted from a saved checkpoint if the cluster crashes. The checkpointing

scripts include the same basic setup, with additional code (that doesn't need to be modified by the user) that handles the checkpointing functionality.

2.1. Basic serial script

The simplest type of job is one that requires only serial computations. Here you simply need to specify the requested queue, provide a name for your job, setup up the software environment, copy the data and code to the scratch directory, and run your code. This is all done within a shell script, with the associated commands for shell scripting available within this submission script.

The first line setups up the bash script

#!/bin/bash

Note that the cluster uses the bash shell, rather than csh or tcsh. Any shell scripts called by you code must be setup in bash.

Next are a series of parameters for the scheduler:

```
#BSUB -q mw256fd
#BSUB -J test
#BSUB -o test.stdout
#BSUB -e test.stderr
#BSUB -N
```

The first line specifies the queue to which the code is sent (e.g. hp12, mwd26fd). The second line provides the name of the job, while the third and fourth lines specify the names for the output and error files. These two files are created once the job is complete and they contain outputs from you code, as well an error messages generated while running your code. The test.stdout file will also contain basic information on the name of the job, the home directory, the working directory, the job id, a snippet of the code, plus some other information about the run. The last line requests that the scheduler send you an email when the code has finished running.

Next you need to set up the software environment for your code. This can require defining path variables for various packages (in this case Python and MIRIAD).

```
export PYTHONHOME=/home/apps/python/2.6.1
export PYTHONPATH=/home/apps/python/2.6.1/site-packages
export PATH=$PYTHONHOME/bin:$PATH
. /home/apps/miriad/MIRRC.sh
export PATH=$MIRBIN:$PATH
```

The first three lines set up the python paths, specifically referencing python 2.6.1. Note that multiple versions of python exist on the cluster, but version 2.6.1 contains numpy, scipy, astropy and emcee which are all used in my own code. Version 2.7.10 also contains these packages, along with PyTransit. There is also a miniconda version of python that has the Galario package, along with astropy and emcee. The last two lines set up the MIRIAD installation. First it calls the MIRIAD initialization script, which sets up the variable \$MIRBIN, which is then added to the path. Note that the MIRIAD installation requires a fortran library that is only available on the mw256fd queue.

The next step is to copy the data and code over to the working directory. Upon the start of a job a temporary scratch directory is created, with a name specified by the ID number associated with job. This directory is removed once the job is complete, but is a useful place for running the code in order to reduce the load on the home directory server. By default this happens in the /sanscratch directory, but if large amounts of data are being generated, then the /localscratch5tb scratch directory should be used⁴.

```
cp -r mydata.fits /sanscratch/$LSB_JOBID
cp -r mycode.py /sanscratch/$LSB_JOBID
cd /sanscratch/$LSB_JOBID
```

The system variable \$LSB_JOBID contains the job ID number that is generated when a job is sent to the Scheduler. This sequence of commands not only copies the data (mydata.fits) and code (mycode.py) into the scratch directory, but it also moves into that directory. This is important so that any read/write commands happen within the scratch directory. Otherwise these commands will be sent to your home directory, which will put unnecessary load onto the server that runs the home directory, slowing down operations on the home directory.

Once the setup is complete, you can call your actual code (test.py), making sure to copy over any results (myresults.txt) from the scratch directory back into the home directory.

```
python test.py
cp myresults.txt /home/kflaherty/mycode/
```

2.2. Basic Parallel script

Running a parallel code that distributes jobs over multiple cores (which may or may not be on the same node) follows the same basic flow as the serial script, with additional parameters for the scheduler, and additional PATH variables to provide

⁴ The /localscratch5tb directory is only available to the mw256fd queue

access to the software that does the parallelization. Listed below are the commands that are included in addition to those already specified within the basic serial script.

The first addition is in the parameters for the scheduler. Since a parallel job requires resources from multiple cores you need to specify the necessary computing resources.

```
#BSUB -n 8
#BSUB -R ''span[hosts=1]''
```

The first command specifies the number of cores requested for this job, in this case 8. The second command tells the scheduler that all of the cores must come from one node. This, in turn, limits the maximum number of cores that can be requested in the first line; in the hp12 queue there are 8 cores per node, while for the mw256fd queue there are 16 cores per node. Conversely, you can remove the second command and replace it with

```
#BSUB -R ''span[ptile=1]''
```

which tells the scheduler to use one core per node (e.g. use 8 nodes, instead of 8 cores within a single node). Here the limit is the number of nodes within a queue (32 for hp12, and 8 for mw256fd). I have always found that openmpi only works when all of the cores are within a single node, rather than being spread out over multiple nodes.

The next step is to set up paths to the openmpi code

```
export PATH=/share/apps/openmpi/1.2+intel-10/bin:$PATH
export LD_LIBRARY_PATH=/share/apps/openmpi/1.2+intel-10/lib:$LD_LIBRARY_PATH
```

The miniconda python environment uses mpich2 instead of openmpi. This can be accessed with:

```
export PATH=/share/apps/CENTOS6/miniconda2/bin:$PATH
export LD_LIBRARY_PATH=/share/apps/CENTOS6/miniconda2/lib:$LD_LIBRARY_PATH
```

From here your code can be called using the line:

```
./lava.openmpi.wrapper python mpi_run_models.py
```

Calling your code is slightly different here that in the serial case. The call is done via a wrapper (lava.openmpi.wrapper) that sets up variables needed for parallel jobs. This wrapper takes the place of a direct call to e.g. mpirun that you would use on your local machine. The wrapper is available from Henk's direction.

tory (/home/hmeij/jobs/parallel/). This particular wrapper works with openmpi (used by python 2.6.1 and python 2.7.10), but a different wrapper is needed for mpich2 for miniconda2. The wrapper is called mpich-mpirun and is available in /opt/openlava-3.1/bin/.

2.3. Checkpointing

Recently Henk has set up the cluster to take advantage of checkpointing within the code, based on the Berkeley Laboratory Checkpoint/Restart tool. What this means is that if the cluster crashes, you can restart from the last checkpoint, rather than having to restart from the very beginning. This is very useful if you e.g. are running a week long MCMC chain and the cluster crashes on day four. The details of how this works are complicated, but Henk has hidden most of this within new execution files, having worked closely with me and Jesse Tarnes ('16, within Seth's group) to make sure our code works with checkpointing. Information on both serial⁵ and parallel⁶ jobs can be found online. Note that checkpointing only works with the versions of openmpi available within python 2.7.10, and not with mpich2 within the miniconda2 python environment (e.g. if you need to use Galario then you cannot do any checkpointing).

Once you have copied over Henk's checkpointing scripts from his directory (/home/hmeij/jobs/blcr/blcr_wrapper.serial), you will notice some small differences from the original sample scripts. The first thing the script does is set up the code to operate from the scratch directory

```
export MYSANSCRATCH=/sanscratch/$LSB_JOBID

cd $MYSANSCRATCH

pre_cmd=''scp $HOME/kflaherty/test.py .''
post_cmd=''scp $MYSANSCRATCH/result.txt $HOME/kflaherty/''
```

These commands first set up the \$MYSANSCRATCH system variable, move to the temporary directory and set up commands that will copy over your code at the start of the run, and copy back any results (result.txt) of the run. The operation of copying your code and data into the scratch directory is not executed right away, this just sets up the commands to be called later. Moving everything to the temporary directory, including your code, is important when checkpointing because the BLCR code has to recover everything that was used by the code in order to properly restart it. This includes every file created by the code while it is running. In the example above, I only have the file test.py that needs to be copied over at the beginning, but this can be expanded to include any pieces of code, data sets, support files, etc. that are needed. In general it is good practice to move your code and any data into

⁵ https://dokuwiki.wesleyan.edu/doku.php?id=cluster:147

⁶ https://dokuwiki.wesleyan.edu/doku.php?id=cluster:148

the scratch, and to execute your code from their in order to reduce the load on the home directories.

Next you need to tell the wrapper that you are starting a new job:

```
mode=start
queue=mwd256fd
cmd=''python test.py''
```

where queue specifies the queue to which the code is sent (the same as specified with BSUB -q), while the second line specifies command used to call the code. If you are restarting a job then the code block above is replaced by:

```
mode=restart
queue=mw256fd
orgjobid=250
```

The queue must be the same as the queue used for the original job. The orgjobid parameter is the job ID number for the original job.

Finally, the time interval over which the checkpointing occurs is specified:

```
cpti=15m
```

This requests that a complete copy of the state of system is saved every 15 minutes. This copy includes all of the files that have been generated, all of the code used to generate them, and the state of the memory. These states are saved within /sanscratch/checkpoints/\$JOBID where \$JOBID is the unique ID number for the job. For any full run of your code a checkpoint every 15 minutes is overkill; more reasonable values are 12h (12 hours) or 2d (2 days).

You can check on the status of the script using two files within the checkpoints folder titled cr_checkpoint.err and cr_mpirun.err. The first of these keeps a log of errors associated with the checkpointing process. The second keeps tracks of any errors/warning output by your code during the course of its operation. This later file can be useful for checking on the progress of your code (e.g. if your code crashes without stopping, you will likely see an error message here).

Also, the blcr wrapper will pipe any text that would normally be written on the screen into a file in your home directory. It is located in /home/kflaherty/.lsbatch and the name will have a format (random number).\$JOBID.out. If your code regularly prints text to the screen, then this is a good place to look for the text as the code is running.

Once the wrapper has been set up it can be sent to the scheduler in the same manner as any other script. As soon as a file titled chk.PID shows up, the code has completed its first checkpoint. From here on the code can be resumed from this point after it crashes.

3. SUBMITTING A JOB

Regardless of whether a job is serial or parallel, or includes checkpointing, submitting the job to the scheduler is handled in the same way. This is done with the bsub command (Figure 1):

bsub < myscript

where myscript is the script containing the commands listed above. This pipes the myscript file into the bsub command, including all of the bsub parameters specified within the script.

```
[kflaherty@greentail code]$ bsub < turbulence_run.parallel
Job <66285> is submitted to queue <mw256fd>.
```

Figure 1. Example of the bsub command. A script is submitted to the mw256fd queue and is assigned a job id number.

Before submitting a job, you can check the status of the queues to e.g. determine how many other jobs are still pending on your desired queue. This is done with bqueues (Figure 2).

[kflaherty@gre	entail	~]\$ bqueues								
QUEUE_NAME		STATUS	MAX	JL/U	JL/P	JL/H	NJOBS	PEND	RUN	SUSP
test _	90	Open:Active	16	-	-	8	0	0	0	0
hp12	50	Open:Active	256	-	-	8	402	146	256	0
mw256fd	50	Open:Active	256	-	-	32	329	90	235	0
mw256	50	Open:Active	140	-	-	28	1176	1037	139	0
mwgpu	50	Open:Active	20	-	-	4	5	0	5	0
matlab	50	Open:Active	16	-	-	8	0	0	0	0
mathematica	50	Open:Active	64	-	-	8	0	0	0	0
stata	50	Open:Active	6	-	-	8	0	0	0	0
bss24	50	Open:Active	104	-	-	2	0	0	0	0

Figure 2. Example of the bqueues command. This lists all of the available queues, the maximum number of jobs, the allowable jobs per host, the number of submitted jobs, the number of pending jobs, the number of running jobs and the number of suspended jobs. Most of the queues are currently overloaded, although this tends to fluctuate with time.

This lists the name of all available queues, the status of the queue, the number of jobs per host, the number of total jobs submitted to the queue, the number of pending jobs, and the number of running jobs.

You can check on any submitted jobs using the bjobs command (Figure 3).

This command will list any jobs that are running, or have been recently submitted. It lists the unique job ID number, the user that submitted the job, the status of the

```
[kflaherty@greentail ~]$ bjobs
JOBID USER STAT QUEUE
                     FROM HOST
                             EXEC HOST
                                     JOB NAME
                                            SUBMIT TIME
               mw256fd
    kflaherty RUN
                      greentail
                              28712
turb_nw200_n300_all_w32_vcs Jun 10 08:36
54341
    kflaherty RUN
               mw256fd
                      greentail
                              turb_nw200_n600_13C021_vcs Jun 13 15:37
                      greentail
    kflaherty RUN
               mw256fd
                              turb nw200 n600 CO21 vcs Jun 13 15:42
```

Figure 3. Example of a call to bjobs. I have submitted three jobs from greentail, all of which are running. Each job is taking up eight cores on different nodes.

job (e.g. running vs pending), the queue that the job was sent to, the login node from which the job was submitted, the cores that are executing the job, the name of the job, and the time that the job that was submitted. A useful variant of bjobs is:

```
bjobs -u all -q mw256fd
```

This lists the jobs from all users that have been submitted to a particular queue. This can be useful for looking in detail at the load on a particular queue. The bjobs command can also be called with the -m node flag, where node is the node name (e.g. n44); bjobs will then show the jobs operating on a particular node.

Similarly, the **bhist** command (Figure 4) lists any submitted jobs, including the amount of time they have spent in various states (e.g. running vs pending).

```
[kflaherty@greentail ~]$ bhist
Summary of time in seconds spent in various states:
                 JOB_NAME
                          PEND
                                    PSUSP
                                             RUN
                                                      USUSP
                                                               SSUSP
                                                                       UNKWN
                                                                                TOTAL
JOBID
        USER
28712
        kflaher *w32 vcs
                           15
                                    0
                                                      4298
                                                              18
                                                                       0
                                             540400
                                                                                544731
        kflaher *021 vcs
                           8
                                    0
                                                                       0
54341
                                             260294
                                                      0
                                                               0
                                                                                260302
54442
        kflaher *021 vcs
                                    0
                                                               0
                                                                       0
                                                                                259968
```

Figure 4. Example of a call to bhist. This shows the three jobs currently running, along with the number of seconds they have spent in various states.

Once a job is running, a number of commands can be used to check its status. The first is lsload (Figure 5), which lists the load on a particular node, in the form of the number of cores that are currently operating, and the amount of memory being used. The number of cores should be at least as large as the number of cores requested by your code (assuming you have requested that all cores be located within a single node). If not, then the code has likely crashed without stopping (see below).

```
[kflaherty@greentail ~]$ lsload n38
HOST NAME
                 status r15s
                                       r15m
                                               ut
                                                     pg
                                                                it
                                                                      tmp
                                                                            SWD
                                                                                   mem
n38
                          40.1
                                40.5
                                       40.1
                                              99%
                                                    0.0
                                                           0 2e+08 9384M
                                                                            32G
                                                                                  237G
```

Figure 5. Example of a call to lsload. The values r15s, r1m, r15m shows the core usage averaged over the previous 15 seconds, 1 miute and 15 minutes.

Another method for checking on the operation of your code is with the top command, which can be called as :

```
ssh <node> top -u <username> -b -n 1
```

where <node> is the node name (e.g. n42) and <username> is your username. This will execute top on all of your commands executing on the specified node (Figure 6). This contains information on the total memory usage (important to keep track of) and the fraction of CPU being devoted to a particular command.

```
[kflaherty@greentail ~]$ ssh n38 top -u kflaherty -b -n 1
top - 15:58:42 up 20 days, 5:32, 0 users, load average: 40.05, 40.44, 40.09
Tasks: 738 total, 41 running, 697 sleeping, 0 stopped, 0 zombie
Cpu(s): 64.1%us, 1.1%sy, 0.0%ni, 33.1%id, 1.6%wa, 0.0%hi, 0.1%si, 0.0%st
Mem: 264498348k total, 75679324k used, 188819024k free, 438936k buffers
Swap: 33554424k total, 0k used, 33554424k free, 68631904k cached
```

PID USER	PR	NI VIRT	RES	SHR S %CPU	%MEM	TIME+	COMMAND
4466 kflahert	20	0 1055m	66m	12m R 97.6	0.0	4312:48	python
4467 kflahert	20	0 1093m	102m	11m R 97.6	0.0	3425:10	
4469 kflahert	20	0 1093m	102m	11m R 97.6	0.0	3414:13	
4475 kflahert	20	0 1093m	102m	11m R 97.6	0.0	3419:58	
4479 kflahert	20	0 1093m	102m	11m R 97.6	0.0	3522:22	python
4481 kflahert	20	0 1093m	102m	11m R 97.6	0.0	3551:21	
4478 kflahert	20	0 1093m	102m	11m R 91.9	0.0	3543:32	python
4477 kflahert	20	0 1093m	102m	11m R 76.9	0.0	3539:46	python
4480 kflahert	20	0 1093m	102m	11m R 73.2	0.0	3546:18	python
4470 kflahert	20	0 1093m	102m	11m R 69.4	0.0	3421:59	python
4471 kflahert	20	0 1090m	100m	11m R 67.5	0.0	3426:43	
4476 kflahert	20	0 1093m	102m	11m R 65.7	0.0	3419:38	
4472 kflahert	20	0 1093m	102m	11m R 63.8	0.0	3422:58	
4473 kflahert	20	0 1093m	102m	11m R 63.8	0.0	3422:43	
4474 kflahert	20	0 1093m	102m	11m R 56.3	0.0	3425:31	

Figure 6. Example of calling top for a particular node. There are lots of python processes running right now.

If you need to terminate a job for any reason, then this can be done with the bkill command:

bkill <jobid>

where <jobid> is the unique ID number for the particular job that you want to kill. This may take a minute or two depending on the complexity of the job that is being killed.

4. MISC.

4.1. Crashing without Stopping

Most of the time if your code crashes the job will be killed, and error messages will be output to the .stderr and .stdout files. But I have run into some instances where the code will crash, but the job will not kill itself. This can happen if *emcee* runs into a problem with a particular model and isn't able to proceed any

further along its chains. If you have piped your output into test.out (described below), then you can see this happen when test.out stops growing. At this point you have to kill the job and examine the error file to discern what went wrong. Similarly, for checkpointed jobs you can look at ~/.lsbatch/[0-9]*.JOBID.out or /sanscratch/checkpoints/\$JOBID/cr_mpirun.err to see if it has registered any errors.

4.2. MIRIAD on the Cluster

Miriad runs the same as on our local machines, with one small exception having to do with how uvmodel handles polarized data and an unpolarized model. On my local machine, if the data has dimensions $N_{base}xN_{chan}x3x2$, where N_{base} is the number of baselines and N_{chan} is the number of channels and the last dimension splits the data into XX and YY, then when the model is run through uvmodel it results in a visibility file with dimensions $2N_{base}xN_{chan}x3x1$. The last dimension has been reduced while the first dimension has been doubled, with the intensity copied into twice as many baseline positions. On the cluster uvmodel does not do this. It maintains the original dimensions of the data $(N_{base}xN_{chan}x3x2)$ and copies intensity into the XX,YY positions in the array. This requires slightly different handling of the model visibilities when they are loaded into python.

Also note that miriad is very talkative, and likes to tell you all about things that are going on when it is running. This can become burdensome when hundreds of thousands of models are run since all of this information will be dumped into the .stderr and .stdout files created at the end of each run (described below). To avoid keeping this useless information, I usually pipe the command into a file (e.g. python code.py > test.out. The file test.out will slowly grow with time, which a useful way to test that your code has not crashed. If you place this file in the scratch directory then it will deleted once the code is done running. When checkpointing is used but the output is instead placed in ~/.lsbatch/[0-9]*.JOBID.out. When the job ends, this file becomes the .out file.

4.3. Multithreading vs OpenMPI

Two possible methods that are often seen for distributing computing task are multithreading and multi-core (e.g. OpenMPI). Multi-threading refers to a single core executing multiple threads concurrently, while multi-core refers to distributing processes across multiple cores. Both kinds of parallelization are available on the cluster, but it is more designed for multi-core work, and problems can arise when multi-threading is combined with multi-core. In particular, using both multi-threading and multi-core can overtax the cores (one of the few things that will result in a worried email from Henk). This will appear as a >100% CPU usage for a particular function; this can be checked by using top on the particular node that is being used, as discussed above.

Table 1. Cluster Commands

Command	Description				
bsub < <myjob></myjob>	Submit a job (<myjob>) to the scheduler</myjob>				
bqueues	Report on the status of the queues within the cluster. The output includes the number of submitted jobs NJOBS, the number of pending jobs PEND, the number of running jobs RUN, the number of allowed jobs per host JL/H, and the status (e.g. Open vs Closed) for each queue on the cluster.				
bjobs	List your submitted jobs, including their job id, the queue they were submitted to, the name of the job, and, if the job is running, a list of the cores on which the code is running.				
bjobs -u all -q <queue></queue>	List all of the jobs currently running on a specific queue.				
bhist	List a history of time spent on your submitted jobs				
lsload <node></node>	Lists the number of currently running cores, averaged over the past 15 seconds (r15s), 1 minute (r1m), and 15 minutes (r15m), on a particular node. It also lists the memory usage for that particular node.				
ssh <node> top -u <username> -b -n 1</username></node>	Run top for node <node> for all jobs submitted by user <username>.</username></node>				
bkil <jobid></jobid>	End/kill a particular job, specified by its unique ID number.				

Note—A description of common commands for use on Wesleyan's computing cluster

If at all possible limit your code to one thread, while farming the jobs out to multiple cores via e.g. OpenMPI. This can often be done in how the code is called, as with emcee, which has both an option for specifying the number of threads as well as sending it an MPI pool. In this case you would skip the threads keyword and simply send it the MPI pool object. Some pieces of code try to perform multithreading in the background. This is the case for Galario, for which you need to restrict the code to one thread (galario.double.threads(1)). At this level multithreading doesn't always provide a substantial increase in speed, while putting a substantial burden on the CPUs.