A few preparatory comments:

• My examples here will be silly toy examples for the purpose of keeping things simple and focused on the parallelization approaches.

• For those not running jobs on the SCF (or EML) network, the details of job submission and the prescriptions below for controlling the number of cores used by your jobs do not apply, but the material should still serve as an introduction to the basics of parallel programming. Analogous code should work on other clusters and in the cloud, such as Amazon’s EC2.

• For those of you in the EML, the instructions for submitting jobs to the EML cluster are essentially identical to those given here.

1 Overview of parallel processing computers

There are two basic flavors of parallel processing (leaving aside GPUs): distributed memory and shared memory. With shared memory, multiple processors (which I’ll call cores) share the same memory. With distributed memory, you have multiple nodes, each with their own memory. You can think of each node as a separate computer connected by a fast network.

For both shared and distributed memory you can make use of the SCF cluster or of individual stand-alone servers, such as gandalf, arwen, and beren. For those of you working on research grants with Statistics faculty, we have a new high-priority cluster with faster nodes.
1.1 Some useful terminology:

- *cores*: We’ll use this term to mean the different processing units available on a single node.

- *nodes*: We’ll use this term to mean the different computers, each with their own distinct memory, that make up a cluster or supercomputer.

- *processes*: computational tasks executing on a machine; multiple processes may be executing at once. A given program may start up multiple processes at once. Ideally we have no more processes than cores on a node.

- *threads*: multiple paths of execution within a single process; the OS sees the threads as a single process, but one can think of them as 'lightweight' processes. Ideally when considering the processes and their threads, we would have no more processes and threads combined than cores on a node.

- *forking*: child processes are spawned that are identical to the parent, but with different process id’s and their own memory.

- *sockets*: some of R’s parallel functionality involves creating new R processes (e.g., starting processes via *Rscript*) and communicating with them via a communication technology called sockets.

1.2 Shared memory

For shared memory parallelism, each core is accessing the same memory so there is no need to pass information (in the form of messages) between different machines. But in some programming contexts one needs to be careful that activity on different cores doesn’t mistakenly overwrite places in memory that are used by other cores.

Some of the shared memory parallelism approaches that we’ll cover are:

1. threaded linear algebra (from R, C, and Matlab)

2. multicore functionality (in R, Matlab and Python)

3. general purpose threaded C programs using openMP

**Threading**  
Threads are multiple paths of execution within a single process. Using *top* to monitor a job that is executing threaded code, you’ll see the process using more than 100% of CPU. When this occurs, the process is using multiple cores, although it appears as a single process rather than as multiple processes. In general, threaded code will detect the number of cores available on a
machine and make use of them. However, you can also explicitly control the number of threads available to a process.

For most threaded code (that based on the openMP protocol), the number of threads can be set by setting the OMP_NUM_THREADS environment variable (VECLIB_MAXIMUM_THREADS on a Mac). E.g., to set it for four threads in bash:

```
export OMP_NUM_THREADS=4
```

Matlab is an exception to this. Threading in Matlab can be controlled in two ways. From within your Matlab code you can set the number of threads, e.g., to four in this case:

```
feature('numThreads', 4)
```

To use only a single thread, you can use 1 instead of 4 above, or you can start Matlab with the `singleCompThread` flag:

```
m matlab -singleCompThread ...
```

### 1.3 Distributed memory

Parallel programming for distributed memory parallelism requires passing messages between the different nodes. The standard protocol for doing this is MPI, of which there are various versions, including `openMPI`. The R package `Rmpi` implements MPI in R.

Some of the distributed memory approaches that we’ll cover are:

1. embarrassingly parallel computations in R using `foreach`

2. explicit use of MPI (in R and C)

### 1.4 Other type of parallel processing

We won’t cover either of these in this material.

#### 1.4.1 GPUs

GPUs (Graphics Processing Units) are processing units originally designed for rendering graphics on a computer quickly. This is done by having a large number of simple processing units for massively parallel calculation. The idea of general purpose GPU (GPGPU) computing is to exploit this capability for general computation.

In spring 2014, I gave a workshop on using GPUs. The SCF has a GPU on its newest cluster, available to those working on research grants working with Statistics faculty members. It’s also possible to use a GPU on an Amazon EC2 virtual machine.
1.4.2 Spark and Hadoop

Spark and Hadoop are systems for implementing computations in a distributed memory environment, using the MapReduce approach. I’ll be giving a workshop on this later in the fall semester, 2014, with materials to be available here.

2 Running jobs on the department cluster and the queueing system

The information in this section is a shortened, but less terse version of that available on the SCF cluster page and EML cluster page.

Both the SCF (Statistics) and EML (Economics) clusters have 8 nodes, with each node having 32 cores. Most clusters and supercomputers have many more nodes but often fewer cores per node.

To submit a job to the cluster, you need to create a simple shell script file containing the commands you want to run. E.g., the script, named job.sh, say, might contain one of the lines that follow, depending on whether you are using R, Matlab, or Python:

- R CMD BATCH --no-save sim.R sim.Rout
- matlab -nodisplay -nodesktop -singleCompThread < sim.m > sim.out
- python sim.py > sim.out

To submit your job:

$ qsub job.sh

Or to submit to the high priority queue (a user can only use 12 cores at any given time in the high priority queue, but jobs take priority over jobs in the low priority queue).

$ qsub -q high.q job.sh

I/O intensive jobs For jobs that read or write a lot of data to disk, it’s best to read/write from the local hard disk rather than to your home directory on the SCF fileserver.

Here’s an example script that illustrates staging your input data to the local hard disk on the cluster node:

#!/bin/bash
if [ ! -d /tmp/$USER/ ]; then mkdir /tmp/$USER; fi
# position input file on local hard disk of node job is on:
cp ~/projectdir/inputData /tmp/$USER
# your R code should find inputData in /tmp/$USER and write output to /tmp/$USER
R CMD BATCH --no-save sim.R sim.out

# your R code should have written output /tmp/$USER/outputData
cp /tmp/$USER/outputData ~/projectdir/.

Note that you are NOT required to stage your data from the local hard disk. By default files will be read from and written to the directory from which you submitted the job, which will presumably be somewhere in your SCF home directory. If you are reading or writing big files, you may want to stage data to the local disk to increase performance (though if I/O is a small part of your job, it may not matter). And if I/O is a big part of your job (i.e., gigabytes and particularly tens of Gb of I/O), we DO ask that you do do this.

**Long-running jobs**

For jobs you expect to run more than three days, you need to flag this for the system. (This allows us to better optimize sharing of resources on the cluster.) For jobs in the low queue, you would do

```
qsub -l h_rt=672:00:00 job.sh
```

and for jobs in the high queue:

```
qsub -q high.q -l h_rt=168:00:00 job.sh
```

These set the maximum run time to 28 days (low queue) and 7 days (high queue) respectively, whereas the default is 5 days.

### 2.1 Submitting jobs: To ’pe’ or not to ’pe’

**Non-parallel jobs** If you have not written any explicit parallel code, you can submit your job as above.

However, we also need to make sure your job does not use more than one core by using multiple threads. By default the system limits any threaded jobs, including calls to the BLAS, to one core by having OMP_NUM_THREADS set by default to one.

However, Matlab is a bit of a renegade and its threading needs to be controlled explicitly by the user. If you are running in Matlab and you submit a job in the manner described above, you are **REQUIRED** to start Matlab with the singleCompThread flag:

```
matlab -singleCompThread ...
```

Alternatively, from within your Matlab code you can do

```
feature(’numThreads’, 1)
```
Parallel jobs  If you want your job to use more than one core (either by using threaded code or by explicitly parallelizing your code), you need to do the following. First, submit your job with the “-pe smp X” flag to qsub, where X is the number of cores (between 2 and 32) that you want. E.g.,

$ qsub -pe smp 4 job.sh

Next in your code, make sure that the total number of processes multiplied by the number of threads per process does not exceed the number of cores you request, by following the guidelines below. Note that the NSLOTS environment variable is set to the number of cores you have requested via -pe smp, so you can make use of NSLOTS in your code.

For jobs other than Matlab jobs, please follow these guidelines:

1. To use more than one thread within a single process in R or C code, set OMP_NUM_THREADS to NSLOTS in your script:

   export OMP_NUM_THREADS=$NSLOTS

   This will make available as many threads as cores that you have requested. For R jobs, this needs to be set outside of R, in the shell, before running/starting R. An alternative is to use the function `omp_set_num_threads()` in the RhpcBLASctl package at the beginning of your R code:

   ```r
   require(RhpcBLASctl)
   omp_set_num_threads(Sys.getenv("NSLOTS"))
   ```

2. To run multiple processes via explicit parallelization in your code, but with a single thread per process, you need to create only as many processes as you have requested. We’ll see various ways for doing this below when using R, Python or C, and you can make use of NSLOTS in your code. Since OMP_NUM_THREADS defaults to one, only a single thread per process will be used.

3. Finally, to run multiple processes with more than one thread per process (in R or C code), you need to make sure that the total number of threads across all processes is no more than the number of cores you have requested. Thus if you want to have H threads per process and your code starts P processes, you should request H \times P cores via -pe smp and you should then set OMP_NUM_THREADS to H.

For Matlab jobs,
1. To use **more than one thread within a single process**, you should include the following Matlab code in your script:

   ```matlab
   feature('numThreads', str2num(getenv('NSLOTS')))
   ```

   This will make available as many threads as cores that you have requested.

2. To **run multiple processes** in a `parfor` job or when using `parfeval()`, you should set the number of workers to `$NSLOTS` in `parpool()` (see the next section for template code). By default it appears that Matlab only uses one thread per worker so your job will use no more cores than you have requested. When using `parfeval()` you can use multiple threads by setting the number of threads within the function being called, but you need to ensure that the number of threads multiplied by number of jobs does not exceed the number of cores requested.

   Note: if you’re interested in combining threading with `parfor`, email consult@stat.berkeley.edu and we can look into whether there is a way to do this.

   Matlab limits you to at most 12 cores per job with `parpool()`, so there is no point in requesting any more than that. (Note that on the EML you can use Matlab DCS to make use of up to 64 cores - see the information on the EML cluster page.)

**Reserving multiple cores**

If the queue is heavily loaded and you request many slots via the `-pe smp` syntax, that many slots may not come open all at once for a long time, while jobs requesting fewer cores may slip in front of your job. You can request that the queue reserve (i.e., accumulate) cores for your job, preventing smaller jobs from preempting your job. To do so, include “-R y” in your `qsub` command, e.g.,

   ```bash
   $ qsub -pe smp 20 -R y job.sh
   ```

### 2.2 Monitoring jobs

You can get information about jobs using `qstat`.

- `qstat # shows your jobs`
- `qstat -u "*" # shows everyone's jobs`
- `qstat -j 7094 # shows more info about a single job with process ID 7094`
- `qdel 7094 # deletes a single job, in this case that with process ID 7094`
- `qdel -u <username> # deletes all your jobs`
2.3 Interactive jobs

In addition to submitting batch jobs, you can also run a job from the command line on a node of the cluster.

```
qrsh -q interactive.q
```

You can then use the node as you would any other Linux compute server, but please remember to exit the session as soon as you are done so you don’t unnecessarily tie up resources. The primary use of this functionality is likely to be prototyping code and debugging. However, note that cluster nodes are set up very similarly to the other Linux compute servers, so most code should run the same on the cluster as on any other SCF Linux machine.

In some cases, you may want to specify a particular node, such as for data transfer to a particular node or to monitor a specific batch job. In this case (here for node `scf-sm01`), do:

```
qrsh -q interactive.q -l hostname=scf-sm01
```

Once on the node you can transfer files via `scp`, use `top` to monitor jobs you have running, etc.

Note that the NSLOTS environment variable is not set in interactive jobs. Please do not use more threads or cores than you requested (which is of course one by default unless you do `qrsh -q interactive.q -pe smp X`).

3 Threaded linear algebra and the BLAS

The BLAS is the library of basic linear algebra operations (written in Fortran or C). A fast BLAS can greatly speed up linear algebra relative to the default BLAS on a machine. Some fast BLAS libraries are Intel’s MKL, AMD’s ACML, and the open source (and free) openBLAS (formerly GotoBLAS). The default BLAS on the SCF Linux compute servers is openBLAS and on the cluster nodes is ACML. All of these BLAS libraries are now threaded - if your computer has multiple cores and there are free resources, your linear algebra will use multiple cores, provided your program is linked against the specific BLAS and provided OMP_NUM_THREADS is not set to one. (Macs make use of VECLIB_MAXIMUM_THREADS rather than OMP_NUM_THREADS.)

3.1 Fixing the number of threads (cores used)

In general, if you want to limit the number of threads used, you can set the OMP_NUM_THREADS UNIX environment variable. This can be used in the context of R or C code that uses BLAS or your own threaded C code, but this does not work with Matlab. In the UNIX bash shell, you’d do this as follows (e.g. to limit to 3 cores) (do this before starting R):

```
export OMP_NUM_THREADS=3 # or “setenv OMP_NUM_THREADS 1” if using csh/tcsh
```
Alternatively, you can set OMP_NUM_THREADS as you invoke R:

```
OMP_NUM_THREADS=3 R CMD BATCH --no-save job.R job.out
```

OMP_NUM_THREADS is set by default to 1 for jobs submitted to the cluster.

### 3.2 Using threading in R, Matlab and C/C++

#### 3.2.1 Using threading in R

Threading in R is limited to linear algebra, for which R calls external BLAS and LAPACK libraries.

Here’s some code that when run on an SCF Linux server illustrates the speed of using a threaded BLAS:

```r
require(RhpcBLASctl)

## Loading required package: RhpcBLASctl

x <- matrix(rnorm(5000^2), 5000)

blas_set_num_threads(4)

system.time({
  x <- crossprod(x)
  U <- chol(x)
})

## user system elapsed
## 20.885 0.120 6.417

blas_set_num_threads(1)

system.time({
  x <- crossprod(x)
  U <- chol(x)
})

## user system elapsed
## 16.341 0.096 16.512
```

On the SCF, R is linked against openBLAS (for the compute servers) and ACML (for the cluster). On the compute servers, linear algebra will be threaded by default, while on the cluster nodes, OMP_NUM_THREADS is set to one by default, so if you want to enable threading, you need to
follow the instructions in the previous section of this document. If you’re on another system, the R installation manual gives information on how link R to a fast BLAS and you can consult with the system administrator. And if you’d like a fast, threaded BLAS enabled on your own Mac, please email consult@stat.berkeley.edu.

3.2.2 Matlab

Many Matlab functions are automatically threaded (not just linear algebra), so you don’t need to do anything special in your code to take advantage of this. So if you’re running Matlab and monitoring top, you may see a process using more than 100% of CPU. However worker tasks within a parfor() use only a single thread.

On the cluster, make sure to follow the instructions in the previous section about setting the number of threads based on the number of cores you have requested.

3.2.3 C/C++

To use threaded BLAS calls in a function you compile from C/C++, just make your usual BLAS or Lapack calls in your code. Then link against BLAS and Lapack (which on our system will automatically link against openBLAS or ACML). Here’s an example C++ program (testLinAlg.cpp) and compilation goes like this (the R and Rmath links are because I use R’s rnorm() function):

```bash
    g++ -o testLinAlg testLinAlg.cpp -I/usr/share/R/include -llapack -lblas -lRmath -lR -O3 -Wall
```

    If you’d like to link against ACML’s BLAS and LAPACK (not necessary on the cluster, where ACML is the default):

```bash
    g++ -o testLinAlg testLinAlg.cpp -I/usr/share/R/include -L/opt/acml5.2.0/gfortran64_fma4_mp/lib -lacml_mp -lgfortran -lgomp -lRt -ldl -lm -lRmath -lR -O3 -Wall
```

    That program is rather old-fashioned and doesn’t take advantage of C++. One can also use the Eigen C++ template library for linear algebra that allows you to avoid the nitty-gritty of calling Lapack Fortran routines from C++. Eigen overloads the standard operators so you can write your code in a more natural way. An example is in testLinAlgEigen.cpp. Note that Eigen apparently doesn’t take much advantage of multiple threads even when compiled with openMP, but in this basic test it was competitive with openBLAS/ACML when openBLAS/ACML was restricted to one core.

```bash
    g++ -o testLinAlgEigen testLinAlgEigen.cpp -I/usr/include/eigen3 -I/usr/share/R/include -lRmath -lR -O3 -Wall -fopenmp
```

    Many more details are available from my workshop on C++ and R packages.
3.3 Important warnings about use of threaded BLAS

3.3.1 Conflict between openBLAS and some parallel functionality in R

There are conflicts between forking in R and threaded BLAS that in some cases affect *foreach* (when using the *multicore* and *parallel* backends), *mclapply()*, and (only if *cluster()* is set up with forking (not the default)) *par(L,S,apply()*). The result is that if linear algebra is used within your parallel code, R hangs. This affects (under somewhat different circumstances) both ACML and openBLAS.

To address this, before running an R job that does linear algebra, you can set OMP_NUM_THREADS to 1 to prevent the BLAS from doing threaded calculations. Alternatively, you can use MPI as the parallel backend (via *doMPI* in place of *doMC* or *doParallel*). You may also be able to convert your code to use *par(L,S,apply()* [with the default PSOCK type] and avoid *foreach* entirely.

3.3.2 Conflict between threaded BLAS and R profiling

There is also a conflict between threaded BLAS and R profiling, so if you are using *Rprof()* you may need to set OMP_NUM_THREADS to one. This has definitely occurred with openBLAS; I’m not sure about other threaded BLAS libraries.

3.3.3 Speed and threaded BLAS

In many cases, using multiple threads for linear algebra operations will outperform using a single thread, but there is no guarantee that this will be the case, in particular for operations with small matrices and vectors. Testing with openBLAS suggests that sometimes a job may take more time when using multiple threads; this seems to be less likely with ACML. This presumably occurs because openBLAS is not doing a good job in detecting when the overhead of threading outweighs the gains from distributing the computations. You can compare speeds by setting OMP_NUM_THREADS to different values. In cases where threaded linear algebra is slower than unthreaded, you would want to set OMP_NUM_THREADS to 1.

More generally, if you have an embarrassingly parallel job, it is likely to be more effective to use the fixed number of multiple cores you have access to so as to split along the embarrassingly parallel dimension without taking advantage of the threaded BLAS (i.e., restricting each process to a single thread).

Therefore I recommend that you test any large jobs to compare performance with a single thread vs. multiple threads. Only if you see a substantive improvement with multiple threads does it make sense to have OMP_NUM_THREADS be greater than one.
4  Basic shared memory parallel programming in R, Python, and Matlab

4.1  Basic shared memory parallel programming in R

4.1.1  foreach

A simple way to exploit parallelism in R when you have an embarrassingly parallel problem (one where you can split the problem up into independent chunks) is to use the \texttt{foreach} package to do a for loop in parallel. For example, bootstrapping, random forests, simulation studies, cross-validation and many other statistical methods can be handled in this way. You would not want to use \texttt{foreach} if the iterations were not independent of each other.

The \texttt{foreach} package provides a \texttt{foreach} command that allows you to do this easily. \texttt{foreach} can use a variety of parallel “back-ends”. It can use \texttt{Rmpi} to access cores in a distributed memory setting as discussed further later in this document or the \texttt{parallel} or \texttt{multicore} packages to use shared memory cores. When using \texttt{parallel} or \texttt{multicore} as the back-end, you should see multiple processes (as many as you registered; ideally each at 100\%) when you look at \texttt{top}. The multiple processes are created by forking or using sockets; this is discussed a bit more later in this document.

\begin{verbatim}
require(parallel)  # one of the core R packages
require(doParallel)
# require(multicore); require(doMC)  # alternative to
# parallel/doParallel require(Rmpi); require(doMPI)  # to
# use Rmpi as the back-end
library(foreach)
library(iterators)

taskFun <- function() {
    mn <- mean(rnorm(1e+07))
    return(mn)
}
nCores <- 8  # manually for non-cluster machines
# nCores <- as.numeric(Sys.getenv('NSLOTS'))  # for use
# on cluster
registerDoParallel(nCores)
# registerDoMC(nCores)  # alternative to
# registerDoParallel cl <- startMPIcluster(nCores);
\end{verbatim}
The result of `foreach` will generally be a list, unless `foreach` is able to put it into a simpler R object. Note that `foreach` also provides some additional functionality for collecting and managing the results that mean that you don’t have to do some of the bookkeeping you would need to do if writing your own for loop.

You can debug by running serially using `%do%` rather than `%dopar%`. Note that you may need to load packages within the `foreach` construct to ensure a package is available to all of the calculations.

**Caution:** Note that I didn’t pay any attention to possible danger in generating random numbers in separate processes. More on this issue in the section on RNG.

### 4.1.2 parallel apply and parallel vectorization (parallel package)

The `parallel` package has the ability to parallelize the various `apply()` functions (apply, lapply, sapply, etc.) and parallelize vectorized functions. The `multicore` package also has this ability and `parallel` is built upon `multicore`. `Parallel` is a core R package so we’ll explore the functionality in that setting. It’s a bit hard to find the vignette for the parallel package because parallel is not listed as one of the contributed packages on CRAN.

First let’s consider parallel apply.

```r
require(parallel)
```

```r
cat("Starting ", i, "th job.\n", sep = "")
outSub <- taskFun()
cat("Finishing ", i, "th job.\n", sep = "")
outSub  # this will become part of the out object
```

```r
nCores <- 8  # manually for non-cluster machines
# nCores <- as.numeric(Sys.getenv('NSLOTS')) # for use on cluster
```

```
# using sockets
```

```
# ?clusterApply
```
cl <- \texttt{makeCluster}(n\texttt{Cores}) \# by default this uses the PSOCK
\# mechanism as in the SNOW package - starting new jobs via Rscript
\# and communicating via sockets

nSims <- 60
input <- \texttt{seq_len}(nSims) \# same as 1:nSims but more robust
testFun <- \texttt{function}(i){
    mn <- \texttt{mean}(\texttt{rnorm}(100000))
    \texttt{return}(mn)
}

# \texttt{clusterExport}(cl, c('x', 'y')) \# if the processes need objects
# \ (x and y, here) from the master's workspace
\texttt{system.time}(
    res <- \texttt{parSapply}(cl, input, testFun)
)

\texttt{system.time}(
    res2 <- \texttt{sapply}(input, testFun)
)

res <- \texttt{parLapply}(cl, input, testFun)


Now let's consider parallel evaluation of a vectorized function. This will often only be worthwhile on very long vectors and for computationally intensive calculations. (The \texttt{Matern()} call here is more time-consuming than \texttt{exp()}).

\texttt{require(parallel)}
nCores <- 8
# nCores <- \texttt{as.numeric(Sys.getenv('NSLOTS'))} \# for use
# on cluster
cl <- \texttt{makeCluster}(nCores)
library(fields)
ds <- runif(6e+06, 0.1, 10)
ds_exp <- pvec(ds, exp, mc.cores = nCores)
# here's a more computationally intensive function
system.time(corVals <- pvec(ds, Matern, 0.1, 2, mc.cores = nCores))
system.time(corVals <- Matern(ds, 0.1, 2))

Note that some R packages can directly interact with the parallelization packages to work with multiple cores. E.g., the boot package can make use of the multicore package directly. If you use such a package on the cluster, you should submit your job with -pe smp and set the number of cores/CPUs used to be $NSLOTS (syntax will vary depending which package is being used).

4.1.3 Using mcparallel() to manually parallelize individual tasks

One can use mcparallel() in the parallel package to send different chunks of code to different processes. Here we would need to manage the number of tasks so that we don’t have more tasks than available cores.

library(parallel)
n <- 1e+07
system.time({
  p <- mcparallel(mean(rnorm(n)))
  q <- mcparallel(mean(rgamma(n, shape = 1)))
  res <- mccollect(list(p, q))
})

system.time({
  p <- mean(rnorm(n))
  q <- mean(rgamma(n, shape = 1))
})

Note that mcparallel() also allows the use of the mc.set.seed argument as with mclapply().

Note that on the cluster, one should create only as many parallel blocks of code as were requested when submitting the job.
4.2  Basic shared memory parallel programming in Matlab

4.2.1  Parallel for loops

To run a loop in parallel in Matlab, you can use the `parfor` construction. Note that once again, this only makes sense if the iterations operate independently of one another. Before running the `parfor` you need to start up a set of workers using `parpool()`. If you’re doing this on the cluster, you must set the `parpool` size to the number of slots requested via the `-pe smp` flag to your `qsub` submission. It appears that Matlab only uses one thread per worker, so you can set the pool size to the number of slots requested. Here is some demo code, also available in `demoParfor.m`.

```matlab
NSLOTS = str2num(getenv('NSLOTS')); # if running on the cluster
% NSLOTS = 8; # otherwise choose how many cores you want to use
pool = parpool(NSLOTS);
n = 3000
nIts = 500
c = zeros(n, nIts);
parfor i = 1:nIts
    c(:,i) = eig(rand(n));
end
delete(pool);
```

4.2.2  Manually parallelizing individual tasks

You can also explicitly program parallelization, managing the individual parallelized tasks. Here is some template code for doing this. We’ll submit our jobs to a pool of workers so that we have control over how many jobs are running at once. Note that here I submit 6 jobs that call the same function, but the different jobs could call different functions and have varying inputs and outputs. Matlab will run as many jobs as available workers in the pool and will queue the remainder, starting them as workers in the pool become available. (So my example is a bit silly in that I have 8 workers but only 6 jobs.) Here is some demo code, also available in `demoBatch.m`.

```matlab
feature('numThreads', 1);
NSLOTS = str2num(getenv('NSLOTS')); % if running on the cluster
% NSLOTS = 8; % otherwise choose how many cores you want to use
pool = parpool(NSLOTS);
% assume you have test.m with a function, test, taking two inputs
% (n and seed) and returning 1 output
n = 10000000;
```
job = cell(1, 6);
job{1} = parfeval(pool, @test, 1, n, 1);
job{2} = parfeval(pool, @test, 1, n, 2);
job{3} = parfeval(pool, @test, 1, n, 3);
job{4} = parfeval(pool, @test, 1, n, 4);
job{5} = parfeval(pool, @test, 1, n, 5);
job{6} = parfeval(pool, @test, 1, n, 6);

% wait for outputs, in order
output = cell(1, 6);
for idx = 1:6
    output{idx} = fetchOutputs(job{idx});
end

% alternative way to loop over jobs:
for idx = 1:6
    jobs(idx) = parfeval(pool, @test, 1, n, idx);
end

% wait for outputs as they finish
output = cell(1, 6);
for idx = 1:6
    [completedIdx, value] = fetchNext(jobs);
    output{completedIdx} = value;
end

delete(pool);

And if you want to run threaded code in a given job, you can do that by setting the number of
threads within the function called by parfeval(). See the testThread.m file in the demo code files
for the testThread() function.

NSLOTS = str2num(getenv('NSLOTS')); % if running on the cluster
% NSLOTS = 8; % otherwise choose how many cores you want to use
pool = parpool(NSLOTS); % matlabpool(NSLOTS) in R2013a and earlier
n = 5000;
nJobs = 2;
pool = parpool(nJobs);
% pass number of threads as number of slots divided by number of jobs
% testThread() function should then do:
% feature('numThreads', nThreads);
% where nThreads is the name of the relevant argument to testThread()
jobt1 = parfeval(pool, @testThread, 1, n, 1, NSLOTS/nJobs);
jobt2 = parfeval(pool, @testThread, 1, n, 2, NSLOTS/nJobs);
jobt3 = parfeval(pool, @testThread, 1, n, 3, NSLOTS/nJobs);
jobt4 = parfeval(pool, @testThread, 1, n, 4, NSLOTS/nJobs);
output1 = fetchOutputs(jobt1);
output2 = fetchOutputs(jobt2);
output3 = fetchOutputs(jobt3);
output4 = fetchOutputs(jobt4);
delete(pool);

Note that functions such as batch(), createJob(), createTask() and submit() appear to be designed for jobs that you submit to a queueing system from within Matlab, which is not how our cluster is set up, so don’t use these on the cluster without emailing consult@stat.berkeley.edu.

4.3 Basic shared memory parallel programming in Python

There are a number of approaches to parallelization in Python. We’ll cover two of the basic approaches here. Note that Python has something called the Global Interpreter Lock that interferes with threading in Python. The approaches below make use of multiple processes.

4.3.1 Multiprocessing package

We’ll first see how to use the multiprocessing package to do multi-core calculations. First we’ll use the Pool.map() method to iterate in a parallelized fashion, as the Python analog to foreach or parfor. Pool.map() only supports having a single argument to the function being used, so we’ll use list of tuples, and pass each tuple as the argument.

```python
import multiprocessing as mp
import numpy as np
```
import os
nCores = int(os.environ['NSLOTS'])
# nCores = 8 # otherwise choose how many cores you want to use

nSmp = 10000000
m = 40
def f(input):
    np.random.seed(input[0])
    return np.mean(np.random.normal(0, 1, input[1]))

# create list of tuples to iterate over, since
# Pool.map() does not support multiple arguments
inputs = [(i, nSmp) for i in xrange(m)]
inputs[0:2]

pool = mp.Pool(processes = nCores)
results = pool.map(f, inputs)
print(results)

We can also manually dispatch the jobs as follows. However, this method will not manage the
processes such that only as many jobs are being done as there are cores, so you would need to
manually manage that.

# set up a shared object to store results
result_queue = mp.Queue()
def f(i, n, out):
    np.random.seed(i)
    # return both index and result as tuple to show how to do that
    out.put(((i, np.mean(np.random.normal(0, 1, n))))

jobs = [] # list of processes
nProc = nCores # don't have more processes than cores available
for i in range(nCores):
    p = mp.Process(target = f, args = (i, nSmp, result_queue))
    jobs.append(p)
p.start()

# wait for results...
for p in jobs:
    p.join()

results = [result_queue.get() for i in range(nCores)]

print(results)

## 4.3.2 pp package

Here we create a server object and submit jobs to the server object, which manages the farming
out of the tasks. Note that this will run interactively in iPython or as a script from UNIX, but will
not run interactively in the base Python interpreter (for reasons that are unclear to me). Also note
that while we are illustrating this as basically another parallelized for loop, the individual jobs can
be whatever calculations you want, so the \( f() \) function could change from job to job.

import numpy
import pp

# on cluster, when using qsub:
import os
nCores = int(os.environ['NSLOTS'])

job_server = pp.Server(ncpus = nCores, secret = 'mysecretphrase')
# set 'secret' to some passphrase (you need to set it but
# what it is should not be crucial)
job_server.get_ncpus()

nSmp = 10000000
m = 40
def f(i, n):
    numpy.random.seed(i)
    return (i, numpy.mean(numpy.random.normal(0, 1, n)))
# create list of tuples to iterate over
inputs = [(i, nSmp) for i in xrange(m)]
# submit and run jobs
jobs = [job_server.submit(f, invalue, modules = ('numpy',)) for invalue in inputs]
# collect results (will have to wait for longer tasks to finish)
results = [job() for job in jobs]
print(results)
job_server.destroy()

5 RNG

The key thing when thinking about random numbers in a parallel context is that you want to avoid having the same 'random' numbers occur on multiple processes. On a computer, random numbers are not actually random but are generated as a sequence of pseudo-random numbers designed to mimic true random numbers. The sequence is finite (but very long) and eventually repeats itself. When one sets a seed, one is choosing a position in that sequence to start from. Subsequent random numbers are based on that subsequence. All random numbers can be generated from one or more random uniform numbers, so we can just think about a sequence of values between 0 and 1.

The worst thing that could happen is that one sets things up in such a way that every process is using the same sequence of random numbers. This could happen if you mistakenly set the same seed in each process, e.g., using `set.seed(mySeed)` in R on every process.

The naive approach is to use a different seed for each process. E.g., if your processes are numbered \( id = 1, \ldots, p \), with \( id \) unique to a process, using `set.seed(id)` on each process. This is likely not to cause problems, but raises the danger that two (or more sequences) might overlap. For an algorithm with dependence on the full sequence, such as an MCMC, this probably won’t cause big problems (though you likely wouldn’t know if it did), but for something like simple simulation studies, some of your 'independent' samples could be exact replicates of a sample on another process. Given the period length of the default generators in R, Matlab and Python, this is actually quite unlikely, but it is a bit sloppy.

One approach to avoid the problem is to do all your RNG on one process and distribute the random deviates, but this can be infeasible with many random numbers.

More generally to avoid this problem, the key is to use an algorithm that ensures sequences that do not overlap.
5.1 Ensuring separate sequences in R

In R, there are two packages that deal with this, `rlecuyer` and `rsprng`. We’ll go over `rlecuyer`, as I’ve heard that `rsprng` is deprecated (though there is no evidence of this on CRAN) and `rsprng` is (at the moment) not available for the Mac.

The L’Ecuyer algorithm has a period of $2^{191}$, which it divides into subsequences of length $2^{127}$.

5.1.1 With the parallel package

Here’s how you initialize independent sequences on different processes when using the `parallel` package’s parallel apply functionality (illustrated here with `parSapply()`).

```r
require(parallel)
require(rlecuyer)
nSims <- 250
testFun <- function(i) {
  val <- runif(1)
  return(val)
}
nSlots <- 4
RNGkind()
cl <- makeCluster(nSlots)
iseed <- 0
# ?clusterSetRNGStream
clusterSetRNGStream(cl = cl, iseed = iseed)
RNGkind()  # clusterSetRNGStream sets RNGkind as L'Ecuyer-CMRG
# but it doesn't show up here on the master
res <- parSapply(cl, 1:nSims, testFun)
clusterSetRNGStream(cl = cl, iseed = iseed)
res2 <- parSapply(cl, 1:nSims, testFun)
identical(res, res2)
stopCluster(cl)
```

If you want to explicitly move from stream to stream, you can use `nextRNGStream()`. For example:
RNGkind("L'Ecuyer-CMRG")
seed <- 0
set.seed(seed)  ## now start M workers
s <- .Random.seed
for (i in 1:M) {
  s <- nextRNGStream(s)
  # send s to worker i as .Random.seed
}

When using `mclapply()`, you can use the `mc.set.seed` argument as follows (note that `mc.set.seed` is TRUE by default, so you should get different seeds for the different processes by default), but one needs to invoke `RNGkind("L'Ecuyer-CMRG")` to get independent streams via the L'Ecuyer algorithm.

```r
require(parallel)
require(rlecuyer)
RNGkind("L'Ecuyer-CMRG")
res <- mclapply(seq_len(nSims), testFun, mc.cores = nSlots,
               mc.set.seed = TRUE)
# this also seems to reset the seed when it is run
res2 <- mclapply(seq_len(nSims), testFun, mc.cores = nSlots,
                 mc.set.seed = TRUE)
identical(res, res2)
```

The documentation for `mcparallel()` gives more information about reproducibility based on `mc.set.seed`.

### 5.1.2 With foreach

**Getting independent streams** One question is whether `foreach` deals with RNG correctly. This is not documented, but the developers (Revolution Analytics) are well aware of RNG issues. Digging into the underlying code reveals that the `doMC` and `doParallel` backends both invoke `mclapply()` and set `mc.set.seed` to TRUE by default. This suggests that the discussion above r.e. `mclapply()` holds for `foreach` as well, so you should do `RNGkind("L'Ecuyer-CMRG")` before your `foreach` call. For `doMPI`, as of version 0.2, you can do something like this, which uses L'Ecuyer behind the scenes:

```r
cl <- makeCluster(nSlots)
setRngDoMPI(cl, seed=0)
```
Ensuring reproducibility  While using foreach as just described should ensure that the streams on each worker are distinct, it does not ensure reproducibility because task chunks may be assigned to workers differently in different runs and the substreams are specific to workers, not to tasks.

For doMPI, you can specify a different RNG substream for each task chunk in a way that ensures reproducibility. Basically you provide a list called .options.mpi as an argument to foreach, with seed as an element of the list:

```r
nslaves <- 4
library(doMPI, quietly = TRUE)
c1 <- startMPIcluster(nslaves)
## 4 slaves are spawned successfully. 0 failed.
registerDoMPI(c1)
result <- foreach(i = 1:20, .options.mpi = list(seed = 0)) %dopar%
        { out <- mean(rnorm(1000)) }
result2 <- foreach(i = 1:20, .options.mpi = list(seed = 0)) %dopar%
          { out <- mean(rnorm(1000)) }
identical(result, result2)
## [1] TRUE
```

That single seed then initializes the RNG for the first task, and subsequent tasks get separate substreams, using the L’Ecuyer algorithm, based on nextRNGStream(). Note that the doMPI developers also suggest using the chunkSize option (also specified as an element of .options.mpi) when using seed. See ?“doMPI-package” for more details.

For other backends, such as doParallel, there is a package called doRNG that ensures that foreach loops are reproducible. Here’s how you do it:

```r
rm(result, result2)
nCores <- 4
library(doRNG, quietly = TRUE)
```
### Loading required package: methods

### Loading required package: pkgmaker

### Loading required package: registry

```r
library(doParallel)

library(doParallel)(nCores)
result <- foreach(i = 1:20, .options.RNG = 0) %dorng% {
    out <- mean(rnorm(1000))
}
result2 <- foreach(i = 1:20, .options.RNG = 0) %dorng% {
    out <- mean(rnorm(1000))
}
identical(result, result2)
```

## [1] TRUE

Alternatively, you can do:

```r
rm(result, result2)
library(doRNG, quietly = TRUE)
library(doParallel)
registerDoParallel(nCores)
registerDoRNG(seed = 0)
result <- foreach(i = 1:20) %dopar% {
    out <- mean(rnorm(1000))
}
```

```
## Warning: error 'Interrupted system call' in select
## Warning: error 'Interrupted system call' in select
```

```r
registerDoRNG(seed = 0)
result2 <- foreach(i = 1:20) %dopar% {
    out <- mean(rnorm(1000))
}
```

```
## Warning: error 'Interrupted system call' in select
## Warning: error 'Interrupted system call' in select
```
identical(result, result2)

## [1] TRUE

### 5.2 Python

Python uses the Mersenne-Twister generator. If you’re using the RNG in *numpy/scipy*, you can set the seed using *(numpy,scipy).random.seed()* . The advice I’m seeing online in various Python forums is to just set separate seeds, so it appears the Python community is not very sophisticated about this issue. There is a function *random.jumpahead()* that allows you to move the seed ahead as if a given number of random numbers had been generated, but this function will not be in Python 3.x, so I won’t suggest using it.

### 5.3 Matlab

Matlab also uses the Mersenne-Twister. We can set the seed as: *rng(seed)* , with seed being a non-negative integer.

Happily, like R, we can set up independent streams, using either of the Combined Multiple Recursion (’mrg32k3a’) and the Multiplicative Lagged Fibonacci (’mlfg6331_64’) generators. Here’s an example, where we create the second of the 5 streams, as if we were using this code in the second of our parallel processes. The ’Seed’, 0 part is not actually needed as that is the default.

```plaintext
thisStream = 2;
totalNumStreams = 5;
seed = 0;
cmrg1 = RandStream.create('mrg32k3a', 'NumStreams', totalNumStreams, 'StreamIndices', thisStream, 'Seed', seed);
RandStream.setGlobalStream(cmrg1);
randn(5, 1)
```

### 6 Forking in R

The *fork* package and *fork()* function in R provide an implementation of the UNIX *fork* system call for forking a process.
library(fork)
{
  # this set of braces is REQUIRED, unless you pass a
  # function to the slave argument of fork()
  pid <- fork(slave = NULL)
  if (pid == 0) {
    cat("Starting child process execution.\n")
    tmpChild <- mean(rnorm(1e+07))
    cat("Result is ", tmpChild, "\n", sep = "")
    save(tmpChild, file = "child.RData")
    cat("Finishing child process execution.\n")
    exit()
  } else {
    cat("Starting parent process execution.\n")
    tmpParent <- mean(rnorm(1e+07))
    cat("Finishing parent process execution.\n")
    wait(pid) # wait til child is finished so can read in
    # updated child.RData below
  }
}
load("child.RData")
print(c(tmpParent, tmpChild))

Note that if we were really running the above code, we’d want to be careful about the RNG. As it stands, it will use the same random numbers in both child and parent processes.

7 OpenMP for C

7.1 Compilation and parallel for

It’s straightforward to write threaded code in C and C++ (as well as Fortran). The basic approach is to use the openMP protocol. Here’s how one would parallelize a loop in C/C++ using an openMP compiler directive. As with foreach in R, you only want to do this if the iterations do not depend on each other.

// see testOpenMP.cpp
#include <iostream>
using namespace std;

// compile with:  g++ -fopenmp -L/usr/local/lib
// testOpenMP.cpp -o testOpenMP

int main(){
    int nReps = 20;
    double x[nReps];
    #pragma omp parallel for
    for (int i=0; i<nReps; i++){
        x[i] = 0.0;
        for ( int j=0; j<1000000000; j++){
            x[i] = x[i] + 1.0;
        }
        cout << x[i] << endl;
    }
    return 0;
}

We would compile this program as follows
$ g++ -L/usr/local/lib -fopenmp testOpenMP.cpp -o testOpenMP

The main thing to be aware of in using openMP is not having different threads overwrite variables used by other threads. In the example above, variables declared within the #pragma directive will be recognized as variables that are private to each thread. In fact, you could declare 'int i' before the compiler directive and things would be fine because openMP is smart enough to deal properly with the primary looping variable. But big problems would ensue if you had instead written the following code:

int main(){
    int nReps = 20;
    int j; // DON'T DO THIS !!!!!!!!!!!!!!!
    double x[nReps];
    #pragma omp parallel for
    for (int i=0; i<nReps; i++){
        x[i] = 0.0;
        for (j=0; j<1000000000; j++){

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\[
x[i] = x[i] + 1.0;
\]

} cout << x[i] << endl;
}
return 0;
}

Note that we do want \( x \) declared before the compiler directive because we want all the threads to write to a common \( x \) (but, importantly, to different components of \( x \)). That’s the point!

We can also be explicit about what is shared and what is private to each thread:

```cpp
int main(){
    int nReps = 20;
    int i, j;
    double x[nReps];
    #pragma omp parallel for private(i,j) shared(x, nReps)
    for (i=0; i<nReps; i++){
        x[i] = 0.0;
        for (j=0; j<1000000000; j++){
            x[i] = x[i] + 1.0;
        }
        cout << x[i] << endl;
    }
    return 0;
}
```

As discussed, when running on the cluster, you are required to use the `-pe smp` parallel environment flag and to set OMP_NUM_THREADS to NSLOTS.

### 7.2 OpenMP for C/C++: more advanced topics

The goal here is just to give you a sense of what is possible with openMP.

The OpenMP API provides three components: compiler directives that parallelize your code (such as `#pragma omp parallel for`), library functions (such as `omp_get_thread_num()`), and environment variables (such as OMP_NUM_THREADS)

*OpenMP* constructs apply to structured blocks of code.

Here’s a basic “Hello, world” example that illustrates how it works:
void myFun(double *in, int id){
    // this is the function that would presumably do the heavy lifting
}

int main()
{
    int nthreads, myID;
    double* input;
    /* make the values of nthreads and myid private to each thread */
    #pragma omp parallel private (nthreads, myID)
    { // beginning of block
        myID = omp_get_thread_num();
        printf("Hello, I am thread \%d\n", myID);
        myFun(input, myID); // do some computation on each thread
        /* only master node print the number of threads */
        if (myid == 0)
        {
            nthreads = omp_get_num_threads();
            printf("Number of threads = \%d\n", nthreads);
        }
    } // end of block
    return 0;
}

The parallel directive starts a team of threads, including the master, which is a member of the
team and has thread number 0. The number of threads is determined in the following ways - here
the first two options specify four threads:

1. #pragma omp parallel NUM_THREADS (4) // set 4 threads for this parallel block
2. omp_set_num_threads(4) // set four threads in general
3. the value of the OMP_NUM_THREADS environment variable
4. a default - usually the number of cores on the compute node

Note that in \texttt{#pragma omp parallel for}, there are actually two instructions, 'parallel' starts a team of threads, and 'for' farms out the iterations to the team. In our \textit{parallel for} invocation, we have done it more explicitly as:

\begin{verbatim}
#pragma omp parallel
#pragma omp for
We can also explicitly distribute different chunks of code amongst different threads:

// see sectionsOpenMP.cpp
#pragma omp parallel // starts a new team of threads
{
    Work0(); // this function would be run by all threads.
    #pragma omp sections // divides the team into sections
    {
        // everything herein is run only once.
        #pragma omp section
        { Work1(); }  
        #pragma omp section
        {
            Work2();
            Work3();
        }
    #pragma omp section
    { Work4(); }  
}
} // implied barrier
\end{verbatim}

Here Work1, \{Work2 + Work3\} and Work4 are done in parallel, but Work2 and Work3 are done in sequence (on a single thread).

If one wants to make sure that all of a parallized calculation is complete before any further code is executed you can insert

\begin{verbatim}
#pragma omp barrier
Note that a \#pragma for statement includes an implicit barrier as does the end of any block specified with \#pragma omp parallel
You can use 'nowait' if you explicitly want to prevent threads from waiting at an implicit barrier: e.g., \#pragma omp parallel sections nowait or \#pragma omp parallel for nowait
\end{verbatim}
One should be careful about multiple threads writing to the same variable at the same time (this is an example of a *race condition*). In the example below, if one doesn’t have the `#pragma omp critical` directive two threads could read the current value of `’sum’` at the same time and then sequentially write to `sum` after incrementing their local copy, which would result in one of the increments being lost. A way to avoid this is with the `critical` directive (for single lines of code you can also use `atomic` instead of `critical`):

```cpp
// see criticalOpenMP.cpp
double sum = 0.0;
double tmp;
#pragma omp parallel for private (tmp, i) shared (sum)
for (int i=0; i<n; i++){
    tmp = myFun(i);
    #pragma omp critical
    sum += tmp;
}
```

You should also be able to use syntax like the following for the parallel for declaration (in which case you shouldn’t need the `#pragma omp critical`):

```cpp
#pragma omp parallel for reduction(+:result)
```

## 8 Distributed memory

### 8.1 MPI basics

There are multiple MPI implementations, of which *openMPI* and *mpich* are very common.

In MPI programming, the same code runs on all the machines, but each machine is assigned a unique identifying number, so the code can include logic so that different code paths are followed on different machines. Since MPI operates in a distributed fashion, any transfer of information between machines must be done explicitly via send and receive calls (*MPI_Send*, *MPI_Recv*, *MPI_Isend*, and *MPI_Irecv*).

The latter two of these functions (*MPI_Isend* and *MPI_Irecv*) are so-called non-blocking calls. One important concept to understand is the difference between blocking and non-blocking calls. Blocking calls wait until the call finishes, while non-blocking calls return and allow the code to continue. Non-blocking calls can be more efficient, but can lead to problems with synchronization between processes.
Debugging MPI/Rmpi code can be tricky because communication can hang, error messages from the workers may not be seen or readily accessible and it can be difficult to assess the state of the worker processes.

8.2 Using message passing on a single node

First let’s get a field for MPI in the simple context of a single node. Of course it’s a bit silly to do this in reality on a single node on which one can take advantage of shared memory.

MPI (specifically openMPI) is installed on all the Linux compute servers, including the cluster nodes. R users can use Rmpi to interface with MPI or can use the functionality in the parallel package to make a socket cluster, which doesn’t use MPI, but does its own version of message passing. There’s not much reason to use MPI on a single node as there’s a cost to the message passing relative to shared memory, but it is useful to be able to test the code on a single machine without having to worry about networking issues across nodes.

8.2.1 MPI example

There are C (mpicc) and C++ (mpic++, mpicxx, mpiCC are synonyms on the SCF system) compilers for MPI programs.

Here’s a basic hello world example (I’ll use the MPI C++ compiler even though the code is all plain C code).

```c
#include <stdio.h>
#include <math.h>
#include <mpi.h>

int main(int argc, char** argv) {
    int myrank, nprocs, namelen;
    char processor_name[MPI_MAX_PROCESSOR_NAME];
    MPI_Init(&argc, &argv);
    MPI_Comm_size(MPI_COMM_WORLD, &nprocs);
    MPI_Comm_rank(MPI_COMM_WORLD, &myrank);
    MPI_Get_processor_name(processor_name, &namelen);
    printf("Hello from processor %d of %d on %s\n", myrank, nprocs, processor_name);
    MPI_Finalize();
}
```

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To compile and run the code, do:

```bash
mpicxx mpiHello.c -o mpiHello
mpirun -np 4 mpiHello
```

This will run multiple (four in this case) processes on the machine on which it is run. Here’s the output:

Hello from processor 2 of 4 on arwen
Hello from processor 3 of 4 on arwen
Hello from processor 0 of 4 on arwen
Hello from processor 1 of 4 on arwen

Note that `mpirun`, `mpiexec`, and `orterun` are synonyms under `openMPI`.

To actually write real MPI code, you’ll need to go learn some of the MPI syntax. See `quad_mpi.c` and `quad_mpi.cpp`, which are example C and C++ programs (for approximating an integral via quadrature) show some of the basic MPI functions. Compilation and running are as above:

```bash
mpicxx quad_mpi.cpp -o quad_mpi
mpirun -np 4 quad_mpi
```

And here’s the output:

03 January 2013 04:02:47 PM

**QUAD_MPI**

C++/MPI version

Estimate an integral of \( f(x) \) from \( A \) to \( B \).

\[
f(x) = \frac{50}{\pi \times (2500 \times x \times x + 1)}
\]

\( A = 0 \)

\( B = 10 \)

\( N = 999999999 \)

\( \text{EXACT} = 0.4993633810764567 \)

Use MPI to divide the computation among 4 total processes, of which one is the master and does not do core computations.

Process 2 contributed \( \text{MY\_TOTAL} = 0.00095491 \)

Process 1 contributed \( \text{MY\_TOTAL} = 0.49809 \)

Process 3 contributed \( \text{MY\_TOTAL} = 0.000318308 \)
Estimate = 0.4993634591634721
Error = 7.808701535383378e-08
Time = 9.904016971588135

QUAD_MPI:
Normal end of execution.

03 January 2013 04:02:56 PM

Note that `mpirun` just dumbly invokes as many processes as you request without regard to the hardware (i.e., the number of cores) on your machine.

Here’s how we would run the program on the SCF cluster via `qsub` using a single node. One would often put the command passed to `qsub` inside a shell script but here I just do it at the command line.

```bash
np=4; qsub -pe smp $np -b y 'mpirun -np $np quad_mpi > results 2>&1'
```

### 8.2.2 Rmpi example

To use `Rmpi`, you can simply start R as you normally do by invoking a command-line R session or using R CMD BATCH.

Here’s R code for using `Rmpi` as the back-end to `foreach`.

```r

# start R as usual: 'R' or via a batch job
library(Rmpi)
library(doMPI)
nslaves <- 4
c1 <- startMPIcluster(nslaves)
registerDoMPI(c1)
clusterSize(c1)  # just to check

## [1] 4

result <- foreach(i = 1:20) %dopar% {
  out <- mean(rnorm(1e+07))
}

closeCluster(c1)

```

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Here’s some example code that uses actual Rmpi syntax, which is very similar to the MPI C syntax we’ve already seen. This code runs in a master-slave paradigm where the master starts the slaves and invokes commands on them. It may be possible to run Rmpi in a context where each process runs the same code based on invoking with Rmpi, but I haven’t investigated this further.

```r
library(Rmpi)

mpi.spawn.Rslaves(nslaves = 4)

## 4 slaves are spawned successfully. 0 failed.
## master (rank 0, comm 1) of size 5 is running on: smeagol
## slave1 (rank 1, comm 1) of size 5 is running on: smeagol
## slave2 (rank 2, comm 1) of size 5 is running on: smeagol
## slave3 (rank 3, comm 1) of size 5 is running on: smeagol
## slave4 (rank 4, comm 1) of size 5 is running on: smeagol

n <- 5

mpi.bcast.Robj2slave(n)

mpi.bcast.cmd(id <- mpi.comm.rank())

mpi.bcast.cmd(x <- rnorm(id))

mpi.remote.exec(ls(.GlobalEnv), ret = TRUE)

## $slave1
## [1] "id" "n" "x"
##
## $slave2
## [1] "id" "n" "x"
##
## $slave3
```
## 

```r
## [1] "id" "n" "x"
##
## $slave4
## [1] "id" "n" "x"

mpi.bcast.cmd(y <- 2 * x)

mpi.remote.exec(print(y))

## $slave1
## [1] -0.9894
##
## $slave2
## [1] -2.120 1.101
##
## $slave3
## [1] 1.6586 -0.8279 -0.6253
##
## $slave4
## [1] 2.1371 1.5313 0.4866 2.2468

objs <- c("y", "z")

# next command sends value of objs on _master_ as argument to rm

mpi.remote.exec(rm, objs)

## $slave2
## [1] 0
##
## $slave4
## [1] 0

mpi.remote.exec(print(z))

## $slave1
## [1] "Error in print(z) : object 'z' not found\n"
## attr(,"class")
## [1] "try-error"
## attr(,"condition")
```
## <simpleError in print(z): object 'z' not found>

## $slave2
## [1] "Error in print(z) : object 'z' not found\n"
## attr(,"class")
## [1] "try-error"
## attr(,"condition")
## <simpleError in print(z): object 'z' not found>

## $slave3
## [1] "Error in print(z) : object 'z' not found\n"
## attr(,"class")
## [1] "try-error"
## attr(,"condition")
## <simpleError in print(z): object 'z' not found>

## $slave4
## [1] "Error in print(z) : object 'z' not found\n"
## attr(,"class")
## [1] "try-error"
## attr(,"condition")
## <simpleError in print(z): object 'z' not found>

# collect results back via send/recv
mpi.remote.exec(mpi.send.Robj(x, dest = 0, tag = 1))

## $slave1
## used (Mb) gc trigger (Mb) max used (Mb)
## Ncells 93666 5.1 350000 18.7 239437 12.8
## Vcells 173426 1.4 786432 6.0 786056 6.0

## $slave2
## used (Mb) gc trigger (Mb) max used (Mb)
## Ncells 93666 5.1 350000 18.7 239437 12.8
## Vcells 173427 1.4 786432 6.0 786056 6.0

## $slave3
## used (Mb) gc trigger (Mb) max used (Mb)
## Ncells 93666 5.1 350000 18.7 239437 12.8
## Vcells 173429 1.4 786432 6.0 786056 6.0

## $slave4
## used (Mb) gc trigger (Mb) max used (Mb)
## Ncells 93666 5.1 350000 18.7 239437 12.8
## Vcells 173429 1.4 786432 6.0 786056 6.0

results <- list()
for (i in 1:mpi.comm.size() - 1) {
  results[[i]] <- mpi.recv.Robj(source = i, tag = 1)
}

print(results)

## [[1]]
## [1] -0.4947
##
## [[2]]
## [1] -1.0599  0.5507
##
## [[3]]
## [1]  0.8293 -0.4139 -0.3127
##
## [[4]]
## [1]  1.0686  0.7657  0.2433  1.1234

A caution concerning Rmpi/doMPI: when you invoke startMPIcluster(), all the slave R processes become 100% active and stay active until the cluster is closed. In addition, when foreach is actually running, the master process also becomes 100% active. So using this functionality involves some inefficiency in CPU usage. This inefficiency is not seen with a sockets cluster (see next) nor when using other Rmpi functionality - i.e., starting slaves with mpi.spawn.Rslaves() and then issuing commands to the slaves.
8.2.3 Sockets in R example

One can also set up a cluster via sockets. In this case we’ll do it on a remote machine but put all the workers on that same machine.

```r
# example with PSOCK cluster
library(parallel)

machine <- "arwen"
cl <- makeCluster(rep(machine, 4))
# cl = makeCluster(4) # this would do it by default on
# your local machine
n <- 1e+07
clusterExport(cl, c("n"))
fun <- function(i) out <- mean(rnorm(n))

result <- parSapply(cl, 1:20, fun)

stopCluster(cl) # not strictly necessary
```

One reason to use Rmpi on a single node is that for machines using openBLAS (all of the compute servers except for the cluster nodes), there is a conflict between the multicore/parallel’s forking functionality and openBLAS that causes foreach to hang when used with the doParallel or doMC parallel back ends.

8.2.4 Threaded linear algebra

Note that if you use linear algebra in your R calculations (or if you run an openMP program via mpirun), you will end up using more than the number of cores specified by the number of slaves requested. To control this you can set OMP_NUM_THREADS before starting your program. However if you are running on a remote machine, such as the sockets example, this variable would not get propagated to the other machines, unless you had set it (say, temporarily) in your .bashrc file. At the moment I don’t know of any other way to control this.
8.3 Using message passing in a distributed memory environment (multiple nodes)

8.3.1 MPI example

To run on multiple machines, we need to let `mpirun` know the names of those machines. We can do this in two different ways.

First, we can pass the machine names directly, replicating the name if we want multiple processes on a single machine.

```
mpirun --host arwen,arwen,treebeard -np 3 mpiHello
```

Here’s the output:

Hello from processor 0 of 3 on arwen
Hello from processor 2 of 3 on treebeard
Hello from processor 1 of 3 on arwen

```
mpirun --host arwen,arwen,treebeard -np 3 quad_mpi
```

Alternatively, we can create a file with the relevant information. Here we’ll specify two processes on arwen for every one process on treebeard.

```
echo 'arwen slots=2' >> .hosts
echo 'treebeard slots=1' >> .hosts
mpirun -machinefile .hosts -np 3 mpiHello
mpirun -machinefile .hosts -np 12 mpiHello # this seems to just recycle what is in .hosts
```

Here’s the output from the last command that requested 12 processes:

Hello from processor 1 of 12 on arwen
Hello from processor 9 of 12 on arwen
Hello from processor 10 of 12 on arwen
Hello from processor 3 of 12 on arwen
Hello from processor 0 of 12 on arwen
Hello from processor 6 of 12 on arwen
Hello from processor 7 of 12 on arwen
Hello from processor 4 of 12 on arwen
Hello from processor 5 of 12 on treebeard
Hello from processor 8 of 12 on treebeard
Hello from processor 11 of 12 on treebeard
Hello from processor 2 of 12 on treebeard
To limit the number of threads for each process, we can tell `mpirun` to export the value of `OMP_NUM_THREADS` to the processes.

```bash
export OMP_NUM_THREADS=2
mpirun -machinefile .hosts -np 3 -x OMP_NUM_THREADS quad_mpi
```

### 8.3.2 Rmpi example

Here for interactive use we run `mpirun` with `-np 1` but provide the hosts information, so `Rmpi` can make the requisite connections for the worker processes. In my testing it appears that `Rmpi` wants the first host in the machine file to be the machine on which you run `mpirun`.

```bash
mpirun -machinefile .hosts -np 1 R --no-save
```

Our host file specifies three slots so I’ll ask for two slaves in the R code, since there is also the master process and it will (inefficiently) use up the resources on a core. But note that whatever is in .hosts will get recycled if you ask for more processes than the number of slots.

```r
# start R as either: mpirun -hostfile .hosts -np 1 R
# --no-save mpirun -hostfile .hosts -np 1 R CMD BATCH
# --no-save

library(Rmpi)
library(doMPI)
nslaves <- 2
# 2 slaves since my host file specifies 3 slots, and one
# will be used for master
cl <- startMPIcluster(nslaves)
registerDoMPI(cl)
clusterSize(cl)  # just to check

results <- foreach(i = 1:200) %dopar% {
  out <- mean(rnorm(1e+07))
}

if (FALSE) {
  foreach(i = 1:20) %dopar% {
    out <- chol(crossprod(matrix(rnorm(3000^2), 3000)))[1, 2]
  }
}
```

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For a batch job, you can do the same as above, but of course with R CMD BATCH. Alternatively, you can do the following with $np set to a number greater than one:

```r
mpirun -machinefile .hosts -np $np R CMD BATCH --no-save tmp.R tmp.out
```

Then in your R code if you call `startMPIcluster()` with no arguments, it will start up $np-1 slave processes by default, so your R code will be more portable.

If you specified -np with more than one process then as with the C-based MPI job above, you can control the threading via OMP_NUM_THREADS and the -x flag to `mpirun`. Note that this only works when the R processes are directly started by `mpirun`, which they are not if you set -np 1. The `maxcores` argument to `startMPIcluster()` does not seem to function (perhaps it does on other systems).

Also note that if you do this in interactive mode, some of the usual functionality of command line R (tab completion, scrolling for history) is not enabled and errors will cause R to quit. This occurs because passing things through `mpirun` causes R to think it is not running interactively.

### 8.3.3 Running an MPI job on the cluster

To run a job on the cluster that uses MPI, the syntax is similar to the examples given just above.

First, you need to request resources via the queueing system using the MPI parallel environment, in this case asking for 40 cores.

```bash
qsub -pe mpi 40 job.sh
```

Then your submission script should invoke `mpirun` and specify `$TMPDIR/machines` as the hosts file and OMIT the number of processors, e.g., as follows for an R job

```bash
mpirun -machinefile $TMPDIR/machines R CMD BATCH --no-save tmp.R tmp.out
```

Note that for Matlab jobs, our license prevents use of more than 12 cores via parpool for use with `parfor`, so one should not use -pe mpi when using Matlab. (Those in the EML can make use of Matlab across multiple nodes - please see the instructions concerning Matlab DCS on the EML cluster page.)

It is possible to use MPI with Python but I haven’t provided any details here.
8.3.4 Note on supercomputers

Note: in some cases a cluster/supercomputer will be set up so that Rmpi is loaded and the worker processes are already started when you start R. In this case you wouldn’t need to load Rmpi or use mpi.spawn.Rslaves() (though you would still need to run startMPIcluster() if using foreach with doMPI). You can always check mpi.comm.size() to see if the workers are already set up.

8.3.5 Sockets in R example

This is not much different from the single node case. You just need to specify a character vector with the machine names as the input to makeCluster().

```r
# multinode example with PSOCK cluster

library(parallel)

machineVec <- c(rep("arwen", 4), rep("treebeard", 2), rep("beren", 2))
cl <- makeCluster(machineVec)

n <- 1e+07
clusterExport(cl, c("n"))
fun <- function(i) out <- mean(rnorm(n))
result <- parSapply(cl, 1:20, fun)

stopCluster(cl)  # not strictly necessary
```

As before, be aware that if you use linear algebra, your processes will use more cores than the number of slaves specified.