Applied HPC with R

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2023-04-09

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Preface

This is a Quarto book.

To learn more about Quarto books visit https://quarto.org/docs/books.

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[1] 2

1 Introduction

1.1 High-Performance Computing: An overview

Loosely, from R's perspective, we can think of HPC in terms of two, maybe three things:

- 1. Big data: How to work with data that doesn't fit your computer
- 2. Parallel computing: How to take advantage of multiple core systems
- 3. Compiled code: Write your own low-level code (if R doesn't has it yet...)

(Checkout CRAN Task View on HPC)

1.2 Big Data

- Buy a bigger computer/RAM memory (not the best solution!)
- Use out-of-memory storage, i.e., don't load all your data in the RAM. e.g. The bigmemory, data.table, HadoopStreaming R packages
- Efficient algorithms for big data, e.g.: biglm, biglasso
- Store it more efficiently, e.g.: Sparse Matrices (take a look at the dgCMatrix objects from the Matrix R package)

1.3 Parallel computing

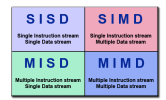


Figure 1.1: Flynn's Classical Taxonomy (Blaise Barney, Introduction to Parallel Computing, Lawrence Livermore National Laboratory)

We will be focusing on the ${\bf S}{\rm ingle}$ Instruction stream Multiple ${\bf D}{\rm ata}$ stream

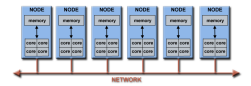
1.4 Parallel computing

1.4.1 Serial vs Parallel

source: Blaise Barney, Introduction to Parallel Computing, Lawrence Livermore National Laboratory

1.5 Parallel computing

knitr::include_graphics("fig/nodesNetwork.png")



source: Blaise Barney, Introduction to Parallel Computing, Lawrence Livermore National Laboratory

1.6 Some vocabulary for HPC

In raw terms

- Supercomputer: A single big machine with thousands of cores/gpus.
- High Performance Computing (HPC): Multiple machines within a single network.
- High Throughput Computing (HTC): Multiple machines across multiple networks.

You may not have access to a supercomputer, but certainly HPC/HTC clusters are more accessible these days, e.g. AWS provides a service to create HPC clusters at a low cost (allegedly, since nobody understands how pricing works)

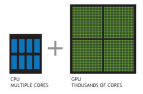


Figure 1.2: NVIDIA Blog

1.7 GPU vs CPU

• Why use OpenMP if GPU is *suited to compute-intensive operations*? Well, mostly because OpenMP is **VERY** easy to implement (easier than CUDA, which is the easiest way to use GPU).

1.8 When is it a good idea?

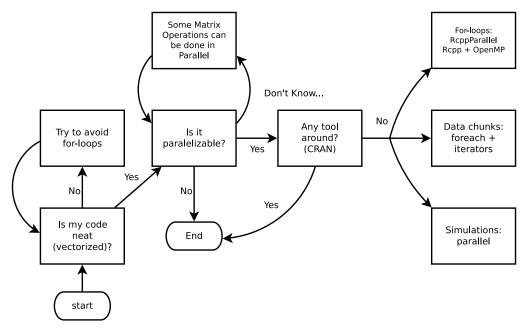


Figure 1.3: Ask yourself these questions before jumping into HPC!

1.9 Parallel computing in R

While there are several alternatives (just take a look at the High-Performance Computing Task View), we'll focus on the following R-packages for **explicit parallelism**:

- parallel: R package that provides '[s]upport for parallel computation, including random-number generation'.
- **future**: '[A] lightweight and unified Future API for sequential and parallel processing of R expression via futures.'
- Rcpp + OpenMP: Rcpp is an R package for integrating R with C++, and OpenMP is a library for high-level parallelism for C/C++ and FORTRAN.

Others but not used here

- foreach for iterating through lists in parallel.
- Rmpi for creating MPI clusters.

And tools for implicit parallelism (out-of-the-box tools that allow the programmer not to worry about parallelization):

- gpuR for Matrix manipulation using GPU
- tensorflow an R interface to TensorFlow.

A ton of other type of resources, notably the tools for working with batch schedulers such as Slurm, and HTCondor.

2 The parallel R package

2.1 Parallel workflow

(Usually) We do the following:

- Create a PSOCK/FORK (or other) cluster using makePSOCKCluster/makeForkCluster (or makeCluster)
- 2. Copy/prepare each R session (if you are using a PSOCK cluster):
 - a. Copy objects with clusterExport
 - b. Pass expressions with clusterEvalQ
 - c. Set a seed
- 3. Do your call: parApply, parLapply, etc.
- 4. Stop the cluster with clusterStop

2.2 Types of clusters: PSOCK

- Can be created with makePSOCKCluster
- Creates brand new R Sessions (so nothing is inherited from the master), e.g.

```
# This creates a cluster with 4 R sessions
cl <- makePSOCKCluster(4)</pre>
```

- Child sessions are connected to the master session via Socket connections
- Can be created outside of the current computer, i.e. across multiple computers!

2.3 Types of clusters: Fork

- Fork Cluster makeForkCluster:
- Uses OS Forking,
- Copies the current R session locally (so everything is inherited from the master up to that point).
- Data is only duplicated if it is altered (need to double check when this happens!)
- Not available on Windows.

Other makeCluster: passed to **snow** (Simple Network of Workstations)

2.4 Ex 1: Parallel RNG with makePSOCKCluster

```
# 1. CREATING A CLUSTER
  library(parallel)
  nnodes <- 4L
         <- makePSOCKcluster(nnodes)
  # 2. PREPARING THE CLUSTER
  clusterSetRNGStream(cl, 123) # Equivalent to `set.seed(123)`
  # 3. DO YOUR CALL
  ans <- parSapply(cl, 1:nnodes, function(x) runif(1e3))</pre>
  (ans0 <- var(ans))</pre>
              [,1]
                            [,2]
                                           [,3]
                                                         [,4]
[1,] 0.0861888293 -0.0001633431 5.939143e-04 -3.672845e-04
[2,] -0.0001633431  0.0853841838  2.390790e-03 -1.462154e-04
[3,] 0.0005939143 0.0023907904 8.114219e-02 -4.714618e-06
[4,] -0.0003672845 -0.0001462154 -4.714618e-06 8.467722e-02
```

Making sure is reproducible

```
# I want to get the same!
clusterSetRNGStream(cl, 123)
ans1 <- var(parSapply(cl, 1:nnodes, function(x) runif(1e3)))
# 4. STOP THE CLUSTER</pre>
```

```
stopCluster(cl)
all.equal(ans0, ans1) # All equal!
```

[1] TRUE

2.5 Ex 2: Parallel RNG with makeForkCluster

In the case of makeForkCluster

```
# 1. CREATING A CLUSTER
  library(parallel)
  # The fork cluster will copy the -nsims- object
  nsims <- 1e3
  nnodes <- 4L
        <- makeForkCluster(nnodes)
  # 2. PREPARING THE CLUSTER
  clusterSetRNGStream(cl, 123)
  # 3. DO YOUR CALL
  ans <- do.call(cbind, parLapply(cl, 1:nnodes, function(x) {</pre>
   runif(nsims) # Look! we use the nsims object!
               # This would have fail in makePSOCKCluster
                # if we didn't copy -nsims- first.
   }))
  (ans0 <- var(ans))</pre>
                          [,2]
             [,1]
                                       [,3]
[1,] 0.0861888293 -0.0001633431 5.939143e-04 -3.672845e-04
[3,] 0.0005939143 0.0023907904 8.114219e-02 -4.714618e-06
[4,] -0.0003672845 -0.0001462154 -4.714618e-06 8.467722e-02
```

Again, we want to make sure this is reproducible

```
# Same sequence with same seed
clusterSetRNGStream(cl, 123)
ans1 <- var(do.call(cbind, parLapply(cl, 1:nnodes, function(x) runif(nsims))))
ans0 - ans1 # A matrix of zeros</pre>
```

```
[,1] [,2] [,3] [,4]
[1,] 0 0 0 0
[2,] 0 0 0 0
[3,] 0 0 0 0
[4,] 0 0 0 0
```

```
# 4. STOP THE CLUSTER stopCluster(cl)
```

Well, if you are a Mac-OS/Linux user, there's a simpler way of doing this...

2.6 Ex 3: Parallel RNG with mclapply (Forking on the fly)

In the case of mclapply, the forking (cluster creation) is done on the fly!

```
# 1. CREATING A CLUSTER
  library(parallel)
  # The fork cluster will copy the -nsims- object
  nsims <- 1e3
  nnodes <- 4L
           <- makeForkCluster(nnodes) # mclapply does it on the fly</pre>
  # 2. PREPARING THE CLUSTER
  set.seed(123)
  # 3. DO YOUR CALL
  ans <- do.call(cbind, mclapply(1:nnodes, function(x) runif(nsims)))
  (ans0 <- var(ans))</pre>
             [,1]
                         [,2]
                                       [,3]
                                                    [,4]
[1,] 0.085384184 0.002390790 0.006576204 -0.003998278
     0.002390790 0.081142190
                               0.001846963 0.001476244
[3,] 0.006576204 0.001846963 0.085175347 -0.002807348
[4,] -0.003998278 0.001476244 -0.002807348 0.082425477
```

Once more, we want to make sure this is reproducible

```
# Same sequence with same seed
  set.seed(123)
  ans1 <- var(do.call(cbind, mclapply(1:nnodes, function(x) runif(nsims))))</pre>
  ans0 - ans1 # A matrix of zeros
    [,1] [,2] [,3] [,4]
[1,]
                0
            0
[2,]
            0
                   0
[3,]
       0
          0
              0
                   0
[4,]
    0
         0
                0
                   0
```

```
# 4. STOP THE CLUSTER
```

[#] stopCluster(cl) no need of doing this anymore

3 Fundamentals

4 Hello world

For a quick-n-dirty intro to Slurm, we will start with a simple "Hello world" using Slurm + R. For this, we need to go through the following steps:

- 1. Copy a Slurm script to HPC,
- 2. Logging to HPC, and
- 3. Submit the job using sbatch.

4.1 Step 1: Copy the Slurm script to HPC

We need to copy the following Slurm script to HPC (00-hello-world.slurm):

```
#!/bin/sh
#SBATCH --output=00-hello-world.out
module load usc r
Rscript -e "paste('Hello from node', Sys.getenv('SLURMD_NODENAME'))"
```

Which has four lines:

- 1. #!/bin/sh the shebang (shewhat?)
- 2. #SBATCH --output=00-hello-world.out an option to be passed to sbatch, in this case, the name of the output file to which stdout and stderr will go.
- 3. module load usc r uses Lmod to load the usc (required) and R modules.
- 4. Rscript ... a call to R to evaluate the expression paste(...). This will get the environment variable SLURMD_NODENAME (which sbatch creates) and print it on a message.

To do so, we will use **Secure copy protocol (scp)**, which allows us to copy data to and fro computers. In this case, we should do something like the following

```
scp 00-hello-world.slurm vegayon@hpc-transfer1.usc.edu:/home1/vegayon/
```

In words, "Using the username vegayon, connect to hpc-transfer1.usc.edu, take the file 00-hello-world.slurm and copy it to /home1/vegayon/. With the file now available in the cluster, we can submit this job using Slurm.

4.2 Step 2: Logging to HPC

- 1. Log-in using ssh. In the case of Windows users, download the **Putty** client. You have two options, the **discovery** or **endeavour** clusters.
- 2. To login, you will need to use your USC Net ID. If your USC email is flasname@usc.edu, your USC Net ID is flastname. Then:

```
ssh flastname@discovery.usc.edu
```

if you want to use the discovery cluster (available to all USC members), or

```
ssh flastname@endeavour.usc.edu
```

if you want to use the endeavour cluster (using private condos).

4.3 Step 3: Submitting the job

Overall, there are two ways to use the compute nodes: interactively (salloc) and in batch mode (sbatch). In this case, since we have a Slurm script, we will use the latter.

To submit the job, we can simple type the following:

```
sbatch 00-hello-world.slurm
```

And that's it!

In the case of interactive sessions, You can start one using the salloc command. For example, if you wanted to run R with 8 cores, using 16 Gigs of memory in total, you would need to do the following:

```
salloc -n1 --cpus-per-task=8 --mem-per-cpu=2G --time=01:00:00
```

Once your request is submitted, you will get access to a compute node. Within it, you can load the required modules and start R:

```
\begin{array}{ll} \text{module load usc r} \\ \textbf{R} \end{array}
```

Interactive sessions are not recommended for long jobs. Instead, use this resource if you need to inspect some large dataset, debug your code, etc.

5 Simulating pi (once more)

This is the same old example that lots of people (including me) have been using to ilustrate parallel computing with R. The example is very simple, we want to approximate pi by doing some Monte Carlo simulations.

We know that the area of a circle is $A = \pi r^2$, which is equivalent to say $\pi = A/r^2$, so, if we can approximate the Area of a circle, then we can approximate π . How do we do this?

Using montecarlo experiments, we have that the probability that a random point x falls within the unit circle can be approximated using the following formula

$$\hat{p} = \frac{1}{n} \sum_{i} \mathbf{1}(x \in \text{Circle})$$

This approximation, \hat{p} , multiplied by the area of the escribed square, which has an area equal to $(2 \times r)^2$, thus, we can finally write

$$\hat{\pi} = \hat{p} \times (2 \times r)^2 / r^2 = 4\hat{p}$$

5.1 Submitting jobs to Slurm

The main way that we will be working is submitting jobs using the sbatch function. This function takes as a main argument a bash file with the program to execute. In the case of R, a regular bash file looks something like this:

```
#!/bin/sh
#SBATCH --job-name=sapply
#SBATCH --time=00:10:00

module load usc r
Rscript --vanilla 01-sapply.R
```

This file has three components:

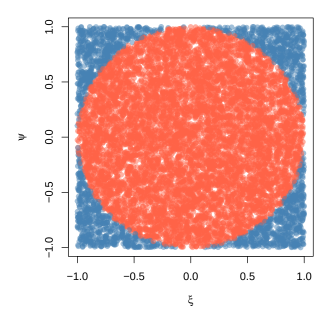


Figure 5.1: 10,000 random points drawn within the unit circle.

- The Slurm flags #SBATCH.
- Loading R module load usc and module load r.
- Executing the R script.

To submit a job the to queue, we need to enter the following:

```
sbatch 01-sapply.slurm
```

The following examples have two files, a bash script and a R script to be called by Slurm.

5.1.1 Case 1: Single job, single core job

The most basic way is submitting a job using the sbatch command. Im this case you need to have 2 files: (1) An R script, and (2) a bash script. e.g.

The contents of the R script (01-sapply.R) are:

```
# Model parameters
nsims <- 1e3
n <- 1e4
# Function to simulate pi</pre>
```

```
simpi <- function(i) {

p <- matrix(runif(n*2, -1, 1), ncol = 2)
  mean(sqrt(rowSums(p^2)) <= 1) * 4

}

# Approximation
set.seed(12322)
ans <- sapply(1:nsims, simpi)

message("Pi: ", mean(ans))

saveRDS(ans, "01-sapply.rds")</pre>
```

The contents of the bashfile (01-sapply.slurm) are:

```
#!/bin/sh
#SBATCH --job-name=sapply
#SBATCH --time=00:10:00

module load usc r
Rscript --vanilla 01-sapply.R
```

5.1.2 Case 2: Single job, multicore job

Now, imagine that we would like to use more than one processor for this job, using something like the parallel::mclapply function from the parallel package. Then, besides of adapting the code, we need to tell Slurm that we are using more than one core per-task, as the following example:

R script (02-mclapply.R):

```
# Model parameters
nsims <- 1e3
n    <- 1e4
ncores <- 4L

# Function to simulate pi
simpi <- function(i) {</pre>
```

```
p \leftarrow matrix(runif(n*2, -1, 1), ncol = 2)
    mean(sqrt(rowSums(p^2)) \le 1) * 4
  }
  # Approximation
  set.seed(12322)
  ans <- parallel::mclapply(1:nsims, simpi, mc.cores = ncores)</pre>
  ans <- unlist(ans)</pre>
  message("Pi: ", mean(ans))
  saveRDS(ans, "02-mclpply.rds")
Bashfile (02-mclapply.slurm):
  #!/bin/sh
  #SBATCH --job-name=mclapply
  #SBATCH --time=00:10:00
  #SBATCH --cpus-per-task=4
  module load usc r
  Rscript --vanilla 02-mclapply.R
```

5.2 Jobs with the slurmR package

5.2.1 Case 3: Single job, multinode job

In this case, there is no simple way to submit a multinodal job to Slurm... unless you use the slurmR package (see installation instructions here)

Once you have the slurmR package in your system, you can proceed as follow

R script (03-parsapply-slurmr.R):

```
# Model parameters
nsims <- 1e3
n <- 1e4
ncores <- 4L</pre>
```

```
# Function to simulate pi
  simpi <- function(i) {</pre>
    p \leftarrow matrix(runif(n*2, -1, 1), ncol = 2)
    mean(sqrt(rowSums(p^2)) \le 1) * 4
  }
  # Setting up slurmR
  library(slurmR) # This also loads the parallel package
  # Making the cluster, and exporting the variables
  cl <- makeSlurmCluster(ncores)</pre>
  # Approximation
  clusterExport(cl, c("n", "simpi"))
  ans <- parSapply(cl, 1:nsims, simpi)</pre>
  # Closing connection
  stopCluster(cl)
  message("Pi: ", mean(ans))
  saveRDS(ans, "03-parsapply-slurmr.rds")
Bashfile (03-parsapply-slurmr.slurm):
  #!/bin/sh
  #SBATCH --job-name=parsapply
  #SBATCH --time=00:10:00
  module load usc r
  Rscript --vanilla 03-parsapply-slurmr.R
```

5.2.2 Case 4: Multi job, single/multi-core

Another way to submit jobs is using **job arrays**. A job array is essentially a job that is repreated njobs times with the same configuration. The main difference between replicates is what you do with the SLURM_ARRAY_TASK_ID environment variable. This variable is defined within each replicate and can be used to make the "subjob" depending on that.

Here is a quick example using R

```
ID <- Sys.getenv("SLURM_ARRAY_TASK_ID")
if (ID == 1) {
    ...[do this]...
} else if (ID == 2) {
    ...[do that]...
}</pre>
```

The slurm RR package makes submitting job arrays easy. Again, with the simulation of pi, we can do it in the following way:

R script (04-slurm_sapply.R):

```
# Model parameters
nsims <- 1e3
      <- 1e4
# ncores <- 4L
njobs <- 4L
# Function to simulate pi
simpi <- function(i, n.) {</pre>
  p \leftarrow matrix(runif(n.*2, -1, 1), ncol = 2)
  mean(sqrt(rowSums(p^2)) \le 1) * 4
}
# Setting up slurmR
library(slurmR) # This also loads the parallel package
# Approximation
ans <- Slurm_sapply(</pre>
  1:nsims, simpi,
  n.
           = n,
  njobs = njobs,
         = "collect",
  plan
  tmp_path = "/scratch/vegayon" # This is where all temp files will be exported
  )
message("Pi: ", mean(ans))
```

```
saveRDS(ans, "04-slurm_sapply.rds")
Bashfile (04-slurm_sapply.slurm):

#!/bin/sh
#SBATCH --job-name=slurm_sapply
#SBATCH --time=00:10:00

module load usc r
```

Rscript --vanilla 04-slurm sapply.R

One of the main benefits of using this approach instead of the the makeSlurmCluster function (and thus, working with a SOCK cluster) are:

- The number of jobs is not limited here (only by the admin, but not by R).
- If a job fails, then we can re-run it using sbatch once again (see example here).
- You can check the individual logs of each process using the function Slurm_lob().
- You can submit the job and quick the R session without waiting for it to finalize. You can always read back the job using the function read_slurm_job([path-to-the-temp])

5.2.3 Case 5: Skipping the .slurm file

The slurmR package has a function named sourceSlurm that can be used to avoid creating the .slurm file. The user can add the SBATCH options to the top of the R script (including the #!/bin/sh line) and submit the job from within R as follows:

R script (05-sapply.R):

```
mean(sqrt(rowSums(p^2)) <= 1) * 4
}

# Approximation
set.seed(12322)
ans <- sapply(1:nsims, simpi)

message("Pi: ", mean(ans))
saveRDS(ans, "05-sapply.rds")</pre>
```

From the R console (is OK if you are in the Head node)

```
slurmR::sourceSlurm("05-sapply.R")
```

And voilá! A temporary bash file will be generated and used submit the R script to the queue.

6 Rcpp

7 RcppArmadillo and OpenMP

- Friendlier than **RcppParallel**... at least for 'I-use-Rcpp-but-don't-actually-know-much-about-C++' users (like myself!).
- Must run only 'Thread-safe' calls, so calling R within parallel blocks can cause problems (almost all the time).
- Use arma objects, e.g. arma::mat, arma::vec, etc. Or, if you are used to them std::vector objects as these are thread safe.
- Pseudo Random Number Generation is not very straight forward... But C++11 has a nice set of functions that can be used together with OpenMP
- Need to think about how processors work, cache memory, etc. Otherwise you could get into trouble... if your code is slower when run in parallel, then you probably are facing false sharing
- If R crashes... try running R with a debugger (see Section 4.3 in Writing R extensions):

```
~$ R --debugger=valgrind
```

7.1 RcppArmadillo and OpenMP workflow

1. Tell Rcpp that you need to include that in the compiler:

```
#include <omp.h>
// [[Rcpp::plugins(openmp)]]
```

2. Within your function, set the number of cores, e.g.

```
// Setting the cores
omp_set_num_threads(cores);
```

7.2 RcppArmadillo and OpenMP workflow

3. Tell the compiler that you'll be running a block in parallel with OpenMP

```
#pragma omp [directives] [options]
{
   ...your neat parallel code...
}
```

You'll need to specify how OMP should handle the data:

- shared: Default, all threads access the same copy.
- private: Each thread has its own copy, uninitialized.
- firstprivate Each thread has its own copy, initialized.
- lastprivate Each thread has its own copy. The last value used is returned.

Setting default (none) is a good practice.

4. Compile!

7.3 Ex 5: RcppArmadillo + OpenMP

Our own version of the dist function... but in parallel!

```
#include <omp.h>
#include <RcppArmadillo.h>
// [[Rcpp::depends(RcppArmadillo)]]
// [[Rcpp::plugins(openmp)]]
using namespace Rcpp;
// [[Rcpp::export]]
arma::mat dist_par(const arma::mat & X, int cores = 1) {

    // Some constants
    int N = (int) X.n_rows;
    int K = (int) X.n_cols;

    // Output
    arma::mat D(N,N);
    D.zeros(); // Filling with zeros

    // Setting the cores
```

```
omp_set_num_threads(cores);

#pragma omp parallel for shared(D, N, K, X) default(none)
  for (int i=0; i<N; ++i)
    for (int j=0; j<i; ++j) {
      for (int k=0; k<K; k++)
            D.at(i,j) += pow(X.at(i,k) - X.at(j,k), 2.0);

      // Computing square root
      D.at(i,j) = sqrt(D.at(i,j));
      D.at(j,i) = D.at(i,j);
    }

    // My nice distance matrix
    return D;
}</pre>
```

```
# Simulating data
set.seed(1231)
K <- 5000
n <- 500
x <- matrix(rnorm(n*K), ncol=K)
# Are we getting the same?
table(as.matrix(dist(x)) - dist_par(x, 4)) # Only zeros</pre>
```

250000

```
# Benchmarking!
microbenchmark::microbenchmark(
  dist(x),  # stats::dist
  dist_par(x, cores = 1), # 1 core
```

```
dist_par(x, cores = 2), # 2 cores
    dist_par(x, cores = 4), # 4 cores
    times = 1,
    unit = "ms"
Unit: milliseconds
                                                  median
                  expr
                           min
                                     lq
                                            mean
                                                               uq
                                                                      max
               dist(x) 2223.023 2223.023 2223.023 2223.023 2223.023
 dist_par(x, cores = 1) 2414.402 2414.402 2414.402 2414.402 2414.402 2414.402
 dist_par(x, cores = 2) 1865.621 1865.621 1865.621 1865.621 1865.621
 dist_par(x, cores = 4) 1223.261 1223.261 1223.261 1223.261 1223.261
```

1

neval

1

1

1

7.4 Ex 6: The future

- **future** is an R package that was designed "to provide a very simple and uniform way of evaluating R expressions asynchronously using various resources available to the user."
- future class objects are either resolved or unresolved.
- If queried, **Resolved** values are return immediately, and **Unresolved** values will block the process (i.e. wait) until it is resolved.
- Futures can be parallel/serial, in a single (local or remote) computer, or a cluster of them.

Let's see a brief example

7.5 Ex 6: The future (cont'd)

```
library(future)
plan(multicore)
# We are creating a global variable
a <- 2</pre>
```

```
# Creating the futures has only the overhead (setup) time
system.time({
 x1 \%<-\% {Sys.sleep(3);a^2}
 x2 \% -\% {Sys.sleep(3);a^3}
})
##
      user system elapsed
     0.023
           0.008
                     0.030
# Let's just wait 5 seconds to make sure all the cores have returned
Sys.sleep(3)
system.time({
 print(x1)
 print(x2)
})
## [1] 4
## [1] 8
##
     user system elapsed
     0.003 0.000 0.003
```

7.6 See also

- Package parallel
- Using the iterators package
- Using the foreach package
- 32 OpenMP traps for C++ developers
- The OpenMP API specification for parallel programming
- 'openmp' tag in Rcpp gallery
- OpenMP tutorials and articles

For more, checkout the CRAN Task View on HPC

7.7 Session info

```
R version 4.2.3 (2023-03-15)

Platform: x86_64-pc-linux-gnu (64-bit)

Running under: Ubuntu 22.04.2 LTS

Matrix products: default

BLAS: /usr/lib/x86_64-linux-gnu/blas/libblas.so.3.10.0

LAPACK: /usr/lib/x86_64-linux-gnu/lapack/liblapack.so.3.10.0
```

locale:

```
[1] LC_CTYPE=en_US.UTF-8 LC_NUMERIC=C
[3] LC_TIME=en_US.UTF-8 LC_COLLATE=en_US.UTF-8
[5] LC_MONETARY=en_US.UTF-8 LC_MESSAGES=en_US.UTF-8
[7] LC_PAPER=en_US.UTF-8 LC_NAME=C
```

[9] LC_ADDRESS=C LC_TELEPHONE=C

[11] LC_MEASUREMENT=en_US.UTF-8 LC_IDENTIFICATION=C

attached base packages:

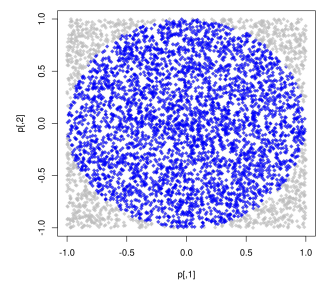
```
[1] stats graphics grDevices utils datasets methods base
```

loaded via a namespace (and not attached):

```
[1] compiler_4.2.3 magrittr_2.0.2 fastmap_1.1.0 cli_3.6.1 [5] tools_4.2.3 htmltools_0.5.4 yaml_2.3.5 Rcpp_1.0.10 [9] stringi_1.7.6 rmarkdown_2.20 knitr_1.37 stringr_1.4.0 [13] jsonlite_1.7.3 xfun_0.37 digest_0.6.29 rlang_1.1.0 [17] evaluate_0.15
```

7.8 Bonus track 1: Simulating π

- We know that $\pi = \frac{A}{r^2}$. We approximate it by randomly adding points x to a square of size 2 centered at the origin.
- So, we approximate π as $\Pr{\|x\| \le 1\} \times 2^2}$



The R code to do this

```
pisim <- function(i, nsim) { # Notice we don't use the -i-
    # Random points
    ans <- matrix(runif(nsim*2), ncol=2)

# Distance to the origin
    ans <- sqrt(rowSums(ans^2))

# Estimated pi
    (sum(ans <= 1)*4)/nsim
}</pre>
```

7.9 Bonus track 1: Simulating π (cont'd)

```
library(parallel)
  # Setup
  cl <- makePSOCKcluster(4L)</pre>
  clusterSetRNGStream(cl, 123)
  # Number of simulations we want each time to run
  nsim <- 1e5
  # We need to make -nsim- and -pisim- available to the
  # cluster
  clusterExport(cl, c("nsim", "pisim"))
  # Benchmarking: parSapply and sapply will run this simulation
  # a hundred times each, so at the end we have 1e5*100 points
  # to approximate pi
  microbenchmark::microbenchmark(
    parallel = parSapply(cl, 1:100, pisim, nsim=nsim),
    serial = sapply(1:100, pisim, nsim=nsim),
    times
          = 1
  )
Unit: milliseconds
              min
                        lq
                               mean
                                      median
parallel 268.7455 268.7455 268.7455 268.7455 268.7455
  serial 329.4686 329.4686 329.4686 329.4686 329.4686
```

```
ans_par <- parSapply(cl, 1:100, pisim, nsim=nsim)
ans_ser <- sapply(1:100, pisim, nsim=nsim)
stopCluster(cl)

par ser R</pre>
```

3.141762 3.141266 3.141593

8 Misc

8.1 General resources

The Center for Advanced Research Computing (formerly HPCC) has tons of resources online. Here are a couple of useful links:

- Center for Advanced Research Computing Website https://carc.usc.edu
- User forum (very useful!) https://hpc-discourse.usc.edu/categories
- Monitor your account https://hpcaccount.usc.edu/
- Slurm Jobs Templates https://carc.usc.edu/user-information/user-guides/high-performance-computing/slurm-templates
- Using R https://carc.usc.edu/user-information/user-guides/software-and-programming/r

8.2 Data Pointers

IMHO, these are the most important things to know about data management at USC's HPC:

- 1. Do your data transfer using the transfer nodes (it is faster).
- 2. Never use your home directory as a storage space (use your project's allotted space instead).
- 3. Use the scratch filesystem for temp data only, i.e., never save important files in scratch.
- 4. Finally, besides of **Secure copy protocol (scp)**, if you are like me, try setting up a GUI client for moving your data (see this).

8.3 The Slurm options they forgot to tell you about...

First of all, you have to be aware that the only thing Slurm does is allocate resources. If your application uses parallel computing or not, that's another story.

Here some options that you need to be aware of:

- ntasks (default 1) This tells Slurm how many processes you will have running. Notice that processes need not to be in the same node (so Slurm may reserve space in multiple nodes)
- cpus-per-task (defatult 1) This is how many CPUs each task will be using. This is what you need to use if you are using OpenMP (or a package that uses that), or anything you need to keep within the same node.
- nodes the number of nodes you want to use in your job. This is useful mostly if you care about the maximum (I would say) number of nodes you want your job to work. So, for example, if you want to use 8 cores for a single task and force it to be in the same node, you would add the option --nodes=1/1.
- mem-per-cpu (default 1GB) This is the MINIMUM amount of memory you want Slurm to allocate for the task. Not a hard barrier, so your process can go above that.
- time (default 30min) This is a hard limit as well, so if you job takes more than the specified time, Slurm will kill it.
- partition (default "") and account (default"") these two options go along together, this tells Slurm what resources to use. Besides of the private resources we have the following:
 - quick partition: Any job that is small enough (in terms of time and memory) will go this way. This is usually the default if you don't specify any memory or time options.
 - main partition: Jobs that require more resources will go in this line.
 - scavenge partition: If you need a massive number resources, and have a job that shouldn't, in principle, take too long to finalize (less than a couple of hours), and you are OK with someone killing it, then this queue is for you. The Scavenge partition uses all the idle resources of the private partitions, so if any of the owners requests the resources, Slurm will cancel your job, i.e. you have no priority (see more).
 - largemem partition: If you need lots of memory, we have 4 1TB nodes for that.

More information about the partitions here

8.4 Good practices (recomendations)

This is what you should use as a minimum:

```
#SBATCH --output=simulation.out

#SBATCH --job-name=simulation

#SBATCH --time=04:00:00

#SBATCH --mail-user=[you]@usc.edu

#SBATCH --mail-type=END,FAIL
```

- output is the name of the logfile to which Slurm will write.
- job-name is that, the name of the job. You can use this to either kill or at least be able to identify what is what you are running when you use myqueue
- time Try always to set a time estimate (plus a little more) for your job.
- mail-user, mail-type so Slurm notifies you when things happen

Also, in your R code

• Any I/O should be done to either Scratch (/scratch/[your usc net id]) or Tmp Sys.getenv("TMPDIR").

8.5 Running R interactively

- 1. The HPC has several pre-installed pieces of software. R is one of those.
- 2. To access the pre-installed software, we use the **Lmod module system** (more information **here**)
- 3. It has multiple versions of R installed. Use your favorite one by running

```
module load usc r/[version number]
```

Where [version number] can be 3.5.6 and up to 4.0.3 (the latest update). The usc module automatically loads gcc/8.3.0, openblas/0.3.8, openmpi/4.0.2, and pmix/3.1.3.

4. It is never a good idea to use your home directory to install R packages, that's why you should try using a **symbolic link instead**, like this

```
cd ~
mkdir -p /path/to/a/project/with/lots/of/space/R
ln -s /path/to/a/project/with/lots/of/space/R R
```

This way, whenever you install your R packages, R will default to that location

5. You can run interactive sessions on HPC, but this recommended to be done using the salloc function in Slurm, in other words, NEVER EVER USE R (OR ANY SOFT-WARE) TO DO DATA ANALYSIS IN THE HEAD NODES! The options passed to salloc are the same options that can be passed to sbatch (see the next section.) For example, if need to do some analyses in the thomas partition (which is private and I have access to), I would type something like

```
salloc --account=lc_pdt --partition=thomas --time=02:00:00 --mem-per-cpu=2G
```

This would put me in a single node allocating 2 gigs of memory for a maximum of 2 hours.

8.6 NoNos when using R

- Do computation on the head node (compile stuff is OK)
- Request a number of nodes (unless you know what you are doing)
- Use your home directory for I/O
- Save important information in Staging/Scratch

9 Summary

In summary, this book has no content whatsoever.

1 + 1

[1] 2

References