## Parallel package

This document gives a brief introduction to one of R's most useful standard packages – the parallel package.

\*\*Motivating example

Lets start with a motivating example. Suppose we want to find the number of primes between **2** and **5,000,000** (inclusive). A sensible approach would consists of the following steps:

- 1. Write a function to detect if a given integer is a prime.
- 2. Apply the function in #1 on all integer between 2 and 5,000,000.
- 3. The total number of primes between 2 and 5,000,000 is the number of primes detected in step #2.

Here is a function that can be used to check if a given integer is a prime:

```
> is.prime <- function(n) ifelse(n == 2 || sum(!(n %% 1:ceiling(sqrt(n)))) == 1, TRUE, FALSE)
> is.prime(2) ## 2 is a prime
[1] TRUE
> is.prime(197) ## 197 is a prime
[1] TRUE
> is.prime(19791) ## 19791 is not a prime
[1] FALSE
```

The following code gives the number of prime numbers between 2 and 5,000,000 while avoiding the use of for loops.

```
> system.time(print(sum(sapply(2:5e6, is.prime))))
[1] 348513
  user system elapsed
45.732  0.168  45.904
```

By default, R handles loops sequentially. What it means is, R will compute the first task at hand and moves on to the next when the first is complete.

$$\boxed{\operatorname{Task} 1} \to \boxed{\operatorname{Task} 2} \to \boxed{\operatorname{Task} 3} \to \cdots \to \boxed{\operatorname{Task} 10000}$$

There are R packages (e.g., snow, multicore, parallel, Segue, etc.) allow R to take advantage of multiple processes to compute tasks in a parallel fashion

Of these packages in parallel computing, only the parallel package is a standard package, meaning you can load the package without having to install it.

```
> library(parallel)
```

The first step is to know how many CPU cores (clusters or workers) are available. This gives us an idea on how to divide the jobs

```
> detectCores()
```

[1] 8

This means I can assign jobs to up to 8 clusters. To assign jobs, the following steps are needed: 1. Define clusters 2. Setup clusters 3. Export functions/libraries to each cluster 4. Assign jobs to each cluster 5. Shut down clusters The following codes give an example for the above prime example:

<sup>\*\*</sup>Parallel computing

```
> cl <- makePSOCKcluster(detectCores())
> setDefaultCluster(cl)
> clusterExport(NULL, "is.prime")
> system.time(print(sum(parSapply(NULL, 2:5e6, FUN = is.prime))))

[1] 348513
    user system elapsed
    2.812    0.224    15.966
> stopCluster(cl)
```

Within the parallel environment, sapply is replaced with parSapply. The parallel also offers functions like parLapply and parApply to replace with lapply and apply.

Note that the computing time with parSapply in this example is not proportional to the computing time without. The lost in computing time is due to the communication between processes. A more efficient way for our example is to submit a batch of jobs to each cluster. For example:

```
c1 <- makePSOCKcluster(detectCores())
> setDefaultCluster(c1)
> clusterExport(NULL, "is.prime")
> system.time(print(sum(parApply(NULL, matrix(1:5e6, 5e4), 1, FUN = function(x) sapply(x, is.prime))) - 1))
[1] 348513
    user system elapsed
    0.192    0.012    11.851
> stopCluster(c1)
```

The first example assigns 5,000,000 jobs to 8 clusters while the second example assigns 50,000 jobs (each job has 100 numbers to compute) to 8 clusters.