# Parallel Computation

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```
## -- Attaching packages ------ tidyverse 1.3.0 --
## v ggplot2 3.2.1 v purrr 0.3.3
## v tibble 2.1.3 v dplyr 0.8.3
## v tidyr 1.0.0 v stringr 1.4.0
## v readr 1.3.1 v forcats 0.4.0
## -- Conflicts ------- tidyverse_conflicts() --
```

# **Parallel Computation**

## Implict Parallel

## x dplyr::lag()

• BLAS (Basic Linear Algebra Subroutines)

## x dplyr::filter() masks stats::filter()

- CRAN R shippings with a version of single threaded BLAS library.
- Microsoft R Open ships with Intel MKL (Win/Linux) / Accelerate ML (macOS) BLAS libraries.
- on macOS, R could be configured to use the optimized BLAS from Apple's Accelerate framework
- We could only install R with different BLAS libraries such as openblas or ATLAS

#### **Embarrassingly Parallel**

Also called perfectly parallel, delightfully parallel or pleasingly parallel.

masks stats::lag()

An embarrassingly parallel task can be considered a trivial case - little or no manipulation is needed to separate the problem into a number of parallel tasks.

A bit deroute first - revisit some of our old friends map and map\_\* in purrr.

```
1:4 %>% map(function(x) x^2)

## [[1]]
## [1] 1
## [2]]
## [1] 4
```

```
##
## [[3]]
## [1] 9
##
## [[4]]
## [1] 16
1:4 %>% map_dbl(function(x) x^2)
## [1] 1 4 9 16
1:4 %>% map_dbl(~ .^2)
## [1] 1 4 9 16
These are the base R equivalence.
1:4 %>% lapply(function(x) x^2)
## [[1]]
## [1] 1
##
## [[2]]
## [1] 4
##
## [[3]]
## [1] 9
## [[4]]
## [1] 16
1:4 %>% sapply(function(x) x^2)
## [1] 1 4 9 16
Suppose we have a list of vectors and we want to operation some operation on each vector.
# it is a slow operation, imagine that in real applications, it could take a few minutes
slow_task <- function(x) {</pre>
  sum(x %o% x)
}
list_of_vectors <- replicate(10, list(rnorm(5000)))</pre>
list_of_vectors %>% glimpse()
## List of 10
## $ : num [1:5000] 0.775 0.639 0.107 0.533 -0.15 ...
## $ : num [1:5000] 0.313 0.879 -0.194 -2.059 1.447 ...
## $ : num [1:5000] 0.402 0.3066 0.5407 -0.5085 -0.0382 ...
## $ : num [1:5000] 0.802 -0.218 1.087 -0.222 0.774 ...
```

```
## $ : num [1:5000] -0.9784 1.1896 0.0203 0.3901 0.6679 ...
## $ : num [1:5000] 0.862 1.317 0.539 0.695 -0.761 ...
## $ : num [1:5000] -2.271 -0.991 0.563 1.537 0.515 ...
## $ : num [1:5000] -1.472 0.469 -1.147 1.536 -0.544 ...
## $ : num [1:5000] 0.662 0.318 0.734 -0.716 -1.115 ...
## $ : num [1:5000] -0.364 -0.215 0.259 0.668 -0.102 ...
list_of_vectors %>% map_dbl(slow_task)
```

However, these commands only run in a single process, it means, if the list is doubled, the time is also at least doubled.

```
system.time({
  list_of_vectors %>% map_dbl(slow_task)
})
##
      user system elapsed
##
     2.468
            1.141
                     3.683
# double the list
longer_list_of_vectors <- c(list_of_vectors, list_of_vectors)</pre>
system.time({
  longer_list_of_vectors %>% map_dbl(slow_task)
})
##
      user system elapsed
##
     4.241
             2.146
                     6.631
```

We are hoping to use multiple processes to speed up the job. The traditional way is to use the parallel package.

### The package parallel

## [1] 6466.426

```
library(parallel)

Consider again the above list_vector example,

# the number of cores we have
detectCores()

## [1] 8

# it will create a socket cluster on my own computer
cl <- makeCluster(4)
parLapply(cl, list_of_vectors, slow_task)

## [[1]]</pre>
```

```
##
## [[2]]
## [1] 27662.34
##
## [[3]]
## [1] 2389.771
##
## [[4]]
## [1] 39.54569
##
## [[5]]
## [1] 817.6046
##
## [[6]]
## [1] 403.4123
##
## [[7]]
## [1] 10655.5
##
## [[8]]
## [1] 1159.628
##
## [[9]]
## [1] 3966.624
##
## [[10]]
## [1] 434.5246
# or if you want simplified result
parSapply(cl, list_of_vectors, slow_task)
    [1] 6466.42626 27662.34303
                                                39.54569
                                                            817.60457
                                                                         403.41228
                                  2389.77135
    [7] 10655.50161 1159.62790
                                  3966.62361
                                                434.52465
# stop the cluster after use
stopCluster(cl)
```

Remark: you don't have to make and stop clusters for every operation, you could make a cluster in the very beginning of your script and close it at the very end.

Let's test the speed improvement

```
run_each <- function(x, fum, n_cores) {
  cl <- makeCluster(n_cores)
  result <- parLapply(cl, x, fun)
  stopCluster(cl)
  result
}
system.time(run_each(longer_list_of_vectors, slow_task, 2))</pre>
```

```
## user system elapsed
## 0.009 0.007 3.158
```

```
system.time(run_each(longer_list_of_vectors, slow_task, 8))
##
     user system elapsed
##
    0.033
           0.041 6.759
Processing Chunk
```

The iteratable is divided into chunks before sending the chunks to the workers. Sys.getpid() tells us the process id of a worker.

```
cl <- makeCluster(4)</pre>
parSapply(cl, 1:10, function(x) {
  Sys.getpid()
})
## [1] 8600 8600 8600 8617 8617 8629 8629 8641 8641 8641
parSapply(cl, 1:10, function(x) {
  Sys.getpid()
},
chunk.size = 2
)
   [1] 8600 8600 8617 8617 8629 8629 8641 8641 8600 8600
parSapply(cl, 1:10, function(x) {
  Sys.getpid()
},
chunk.size = 1
    [1] 8600 8617 8629 8641 8600 8617 8629 8641 8600 8617
stopCluster(cl)
```

## Load balancing

parLapply pre-schedules the tasks to each work. It could be suboptimal when different tasks require different amount of time to complete.

```
cl <- makeCluster(4)</pre>
x \leftarrow c(3, 3, 1, 1, 1, 1, 1, 1)
pause <- function(x) {</pre>
  Sys.sleep(x)
system.time({
  map(x, pause)
})
```

```
##
      user system elapsed
            0.000 12.016
##
     0.005
system.time({
  parLapply(cl, x, pause)
})
##
           system elapsed
      user
            0.000
                     6.010
##
     0.003
system.time({
  parLapply(cl, x, pause, chunk.size = 2)
})
##
      user system elapsed
                     6.006
     0.002
            0.001
##
system.time({
  parLapply(cl, x, pause, chunk.size = 1)
})
##
      user system elapsed
##
     0.003
            0.000
                     4.009
```

Instead of presheeduling the tasks, a task could also be assigned to a free worker dynamically using parLapplyLB.

```
system.time({
   parLapplyLB(cl, x, pause)
})

## user system elapsed
## 0.006 0.001 3.007
Note that it only takes 3 seconds now.
```

```
stopCluster(cl)
```

### Interact directly with the workers

We just saw an quick example on using parLapply. Let's try a few more things.

```
cl <- makeCluster(4)</pre>
```

We could run some arbitrary commands on each of the workers

```
clusterEvalQ(cl, {
    x <- rnorm(100)
    mean(x)
})</pre>
```

```
## [[1]]
## [1] 0.02233895
##
## [[2]]
## [1] -0.1053939
##
## [[3]]
## [1] -0.03107646
##
## [[4]]
## [1] 0.003768593
clusterEvalQ(cl, {
  Sys.getpid()
})
## [[1]]
## [1] 8719
##
## [[2]]
## [1] 8732
##
## [[3]]
## [1] 8744
##
## [[4]]
## [1] 8756
Global variables in master are not exported to the worker automatically (the same is true for parLapply)
y <- 3
clusterEvalQ(cl, {
})
## Error in checkForRemoteErrors(lapply(cl, recvResult)): 4 nodes produced errors; first error: object
clusterExport exports the global variables to each worker.
clusterExport(cl, "y")
clusterEvalQ(cl, {
  y + 1
})
## [[1]]
## [1] 4
##
## [[2]]
## [1] 4
##
## [[3]]
## [1] 4
```

```
##
## [[4]]
## [1] 4
```

## [[4]] ## [1] 6

Note: if you use clusterExport inside a function, you may want to specify the envir parameter.

```
doCalc <- function() {</pre>
  z \leftarrow 5
  # export z from the current scope
  clusterExport(cl, "z", envir = environment())
  clusterEvalQ(cl, {
    z + 1
  })
}
doCalc()
## [[1]]
## [1] 6
##
## [[2]]
## [1] 6
## [[3]]
## [1] 6
```

If you want to set a random seed, the following doesn't work because each work returns the same result.

```
# wrong
set.seed(123)
clusterEvalQ(cl, {
  rnorm(5)
})
## [1] -0.88326009 -2.16480736 1.77660023 -0.13725486 0.06906422
##
## [[2]]
## [1] -0.8128060   0.4333262 -0.2760510 -1.1120264 -1.0612642
## [[3]]
## [1] -0.03124709 2.15854565 -2.47045119 -0.01678581 -0.41465096
## [[4]]
## [1] 0.4309572 -0.2502550 1.6120490 0.8735261 -1.6402650
# wrong again
clusterEvalQ(cl, {
  set.seed(123)
  rnorm(5)
})
```

```
## [[1]]
## [1] -0.56047565 -0.23017749 1.55870831 0.07050839 0.12928774
##
## [[2]]
## [1] -0.56047565 -0.23017749 1.55870831 0.07050839 0.12928774
##
## [[3]]
## [1] -0.56047565 -0.23017749 1.55870831 0.07050839 0.12928774
##
## [[4]]
## [1] -0.56047565 -0.23017749 1.55870831 0.07050839 0.12928774
clusterSetRNGStream(cl, 123)
clusterEvalQ(cl, {
 rnorm(5)
})
## [[1]]
## [1] -0.9685927  0.7061091  1.4890213 -1.8150926  0.3304096
##
## [[2]]
## [1] -0.4094454   0.8909694 -0.8653704   1.4642711   1.2674845
## [[3]]
##
## [[4]]
## [1] -1.0388664 1.5745125 0.7470820 0.6718720 0.2691436
# do not forget to close the cluster
stopCluster(cl)
```

#### Another pouplar package: foreach

As we have seen a bit earlier, we might need to use clusterExport to export certain gloabl variables. It would be a bit cumbersome. To reduce the extra steps, we could consider foreach and doParallel. They will send all the globals to the workers before running the tasks. (However, I don't personally recommand foreach though it is (was!?) popular, I perfer furrer in the below)

```
library(foreach)

##

## Attaching package: 'foreach'

## The following objects are masked from 'package:purrr':

##

## accumulate, when

library(doParallel)

## Loading required package: iterators
```

```
cl <- makeCluster(4)
registerDoParallel(cl)
m <- matrix(rnorm(9), 3, 3)
# matrix m is sent to the workers implictly
# in fact, all globals are sent to the workers in default
foreach(i = seq_len(nrow(m)), .combine = c) %dopar% {
    sum(m[i, ])
}</pre>
```

```
## [1] -0.02905105 -1.36595099 2.58692045
```

```
stopCluster(cl)
```

However, it is not quite "functional" as map, lapply and parLapply.

## map or lapply like syntax

```
mclapply from parallel (unix / macOS only)
```

Remark: mclapply relies on forking, it means that it doesn't work on Windows. We will discuss a cross platform approach.)

```
# in default, `mclapply` uses 2 cores
system.time({
    mclapply(
        c(2, 2, 2, 2),
        function(x) Sys.sleep(x)
    )
})
```

```
## user system elapsed
## 0.005 0.012 4.015
```

```
system.time({
  mclapply(
    c(3, 3, 1, 1, 1, 1, 1),
    function(x) Sys.sleep(x),
  mc.preschedule = FALSE, # set FALSE to enable load balancing
  mc.cores = 4
)
})
```

```
## user system elapsed
## 0.020 0.034 3.028
```

#### package furrr

furr provides functions which are very similar to those in purr. One nice thing about furr is that it doesn't send absolutely every global variables to the workers as foreach and mclapply. It does lexical analysis to deduce what objects are needed and only transfer them to the workers. It also loads libraries automatically in the workers.

```
library(furrr)
## Loading required package: future
# to use 4 workers, `plan(multiprocess)` will use all the avaliable workers
plan(multiprocess, workers = 4)
system.time({
  future_map(
    c(2, 2, 2, 2),
    ~ Sys.sleep(.)
})
##
      user system elapsed
##
     0.089
            0.003
                      2.388
future_map has a family of type speific functions. For example,
future_map_dbl(list(1:10, 11:20, 21:30, 31:41), ~ sum(.))
## [1] 55 155 255 396
load balanacing in future_map
# without load balanacing
system.time({
  future_map(
    c(3, 3, 1, 1, 1, 1, 1, 1),
    ~ Sys.sleep(.)
})
##
      user system elapsed
##
     0.111
            0.005
                     6.046
# with load balanacing
system.time({
  future_map(
    c(3, 3, 1, 1, 1, 1, 1, 1),
    ~ Sys.sleep(.),
    .options = future_options(scheduling = FALSE)
  )
})
##
      user system elapsed
##
     0.171
             0.009
                      4.008
```

### The honorable mention multidplyr

The package multidplyr is tidyverse solution to multiprocess data management. However, it is not released to CRAN yet, it means we need to install it from its github repo by using the package remotes.

```
remotes::install_github("tidyverse/multidplyr")
```

Since it is still under developement, use it with caution!

```
library(multidplyr)
```

There are two ways to get data to the workers in cluster:

- partition() a data frame that already loaded in the interactive process.
- Load a different subset of the data in each worker.

Use partition() to send data to workers

```
library(nycflights13)
cluster <- new_cluster(4)</pre>
flights %>%
  group_by(dest, origin) %>%
  partition(cluster) %>%
  summarize(air_time = mean(air_time, na.rm = TRUE))
## Source: party_df [224 x 3]
## Groups: dest
## Shards: 4 [43--67 rows]
##
##
     dest origin air_time
##
                     <dbl>
     <chr> <chr>
## 1 ABQ
           JFK
                     249.
## 2 ATL
           JFK
                     112.
## 3 AVL EWR
                      89.8
## 4 AVL
                      92.2
           LGA
## 5 BDL
           EWR
                      25.5
## 6 BHM
           JFK
                     117
## # ... with 218 more rows
```

Load data in each worker

```
cluster <- new_cluster(4)

cluster %>% cluster_library(c("tidyverse", "nycflights13"))

cluster %>%
    cluster_assign_partition(destination = unique(flights$dest))

# let's check the `destimation` variable in the workers

cluster %>%
    cluster_call(destination)
```

```
## [[1]]
## [1] "IAH" "MIA" "BQN" "ATL" "ORD" "FLL" "IAD" "MCO" "PBI" "TPA" "LAX" "SFO"
```

```
## [13] "DFW" "BOS" "LAS" "MSP" "DTW" "RSW" "SJU" "PHX" "BWI" "CLT" "BUF" "DEN"
  [25] "SNA" "MSY" "SLC"
##
## [[2]]
  [1] "XNA" "MKE" "SEA" "ROC" "SYR" "SRQ" "RDU" "CMH" "JAX" "CHS" "MEM" "PIT"
## [13] "SAN" "DCA" "CLE" "STL" "MYR" "JAC" "MDW" "HNL" "BNA" "AUS" "BTV" "PHL"
## [25] "STT" "EGE"
##
## [[3]]
   [1] "AVL" "PWM" "IND" "SAV" "CAK" "HOU" "LGB" "DAY" "ALB" "BDL" "MHT" "MSN"
## [13] "GSO" "CVG" "BUR" "RIC" "GSP" "GRR" "MCI" "ORF" "SAT" "SDF" "PDX" "SJC"
## [25] "OMA" "CRW"
## [[4]]
## [1] "OAK" "SMF" "TUL" "TYS" "OKC" "PVD" "DSM" "PSE" "BHM" "CAE" "HDN" "BZN"
## [13] "MTJ" "EYW" "PSP" "ACK" "BGR" "ABQ" "ILM" "MVY" "SBN" "LEX" "CHO" "TVC"
## [25] "ANC" "LGA"
cluster %>% cluster_send({
 df <- flights %>% filter(dest %in% destination)
})
cluster %>%
 party_df("df") %>%
  group_by(dest, origin) %>%
 summarize(air_time = mean(air_time, na.rm = TRUE)) %>%
collect()
## # A tibble: 224 x 3
## # Groups: dest [105]
##
     dest origin air_time
##
      <chr> <chr>
                     <dbl>
   1 ATL
##
           EWR
                     112.
   2 ATL
            JFK
                     112.
## 3 ATL
           LGA
                     114.
## 4 BOS
           EWR
                      40.3
## 5 BOS
           JFK
                      38.5
## 6 BOS
           LGA
                      37.9
## 7 BQN
           EWR
                     196.
## 8 BQN
           JFK
                     194.
## 9 BUF
           EWR
                      51.2
## 10 BUF
           JFK
                      57.1
## # ... with 214 more rows
```

#### Reference:

 $\bullet \ \ R \ Programming \ for \ Data \ Science \ https://bookdown.org/rdpeng/rprogdatascience/parallel-computation. \ html$