Analysis of data from aCGH experiments using parallel computing and ff objects: long list of examples

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1 This vignette

We provide here example calls of all segmentation methods, with different options for methods, as well as different options for type of input object and clustering. This is provided here as both extended help and as a simple way of checking that all the functions can be run and yield the same results regardless of type of input and clustering.

2 Creating objects

We must ensure that we can run this vignette as stand alone. Thus, we load the package and create all necessary objects. This repeats work done in the main vignette.

We first try to move to the "/tmp" directory, if it exists. If it does not, the code will be executed in your current directory.

- > try(setwd("~/tmp"))
- > library(ADaCGH2)
- > ## loading in-RAM objects
- > data(inputEx)
- > summary(inputEx)

```
ΙD
                   chromosome
                                     position
                                                            L.1
Hs.101850:
                         :1.000
             1
                 Min.
                                  Min. : 1180411
                                                      Min.
                                                              :-1.07800
                                  1st Qu.: 36030889
Hs.1019 :
                 1st Qu.:1.000
                                                       1st Qu.:-0.22583
Hs.105460:
            1
                 Median :2.000
                                  Median : 70805790
                                                      Median :-0.01600
Hs.105656: 1
                 Mean
                        :2.284
                                  Mean
                                         : 92600349
                                                      Mean
                                                              :-0.03548
                 3rd Qu.:3.000
                                                       3rd Qu.: 0.16000
Hs.105941: 1
                                  3rd Qu.:149843856
Hs.106674: 1
                 Max.
                        :5.000
                                  Max.
                                         :243795357
                                                             : 0.88300
 (Other) :494
                                                      NA's
                                                              :5
     L.2
                                                                  L3
                           m4
                                              m5
Min.
        :-0.795000
                     Min.
                             :-0.1867
                                        Min.
                                               :-4.67275
                                                            Min.
                                                                   :-13.273
 1st Qu.:-0.139000
                     1st Qu.: 1.9790
                                        1st Qu.:-0.02025
                                                            1st Qu.:
                                                                      3.631
Median :-0.006000
                     Median : 2.2807
                                        Median : 0.43725
                                                            Median :
                                                                      3.925
Mean
        : 0.007684
                     Mean
                            : 3.4504
                                        Mean
                                               : 1.60159
                                                            Mean
                                                                      1.981
                     3rd Qu.: 5.8235
3rd Qu.: 0.134000
                                        3rd Qu.: 3.04475
                                                            3rd Qu.:
                                                                      4.110
        : 1.076000
                     Max.
                            : 6.6043
                                               : 9.60425
                                                                      6.374
Max.
                                        Max.
                                                            Max.
NA's
                                        NA's
                                               :41
                                                            NA's
        :15
                                                                   :9
      m6
Min.
        :-0.7655
 1st Qu.:-0.2260
Median :-0.0440
Mean
        :-0.0351
3rd Qu.: 0.1620
Max.
        : 0.7750
NA's
        :203
> head(inputEx)
                              ID chromosome position
                                                        L.1
                                                               L.2
                                                                        m4
                      Hs.212680
1*1180411*Hs.212680
                                          1
                                             1180411
                                                        NA 0.038 6.22625
1*1188041.5*Hs.129780 Hs.129780
                                             1188042
                                                        NA 0.028 6.17425
1*1194444*Hs.42806
                       Hs.42806
                                          1
                                             1194444
                                                        NA 0.042 6.17425
1*1332537*Hs.76239
                       Hs.76239
                                          1
                                             1332537
                                                        NA 0.285 5.62425
1*2362211*Hs.40500
                       Hs.40500
                                          1
                                             2362211
                                                        NA 0.058 5.85125
1*2372287*Hs.449936
                      Hs.449936
                                             2372287 0.294 -0.006 5.68525
                           m5
                                  L3 m6
1*1180411*Hs.212680
                      3.22625 6.038 NA
1*1188041.5*Hs.129780 3.17425 6.028 NA
1*1194444*Hs.42806
                      3.17425 6.042 NA
1*1332537*Hs.76239
                                  NA NA
                      2.62425
1*2362211*Hs.40500
                      2.85125
                                  NA NA
1*2372287*Hs.449936
                      2.68525
                                  NA NA
> cgh.dat <- inputEx[, -c(1, 2, 3)]</pre>
> chrom.dat <- as.integer(inputEx[, 2])</pre>
> pos.dat <- inputEx[, 3]</pre>
> ## choosing working dir for cluster
> originalDir <- getwd()</pre>
> if(!file.exists("ADaCGH2_vignette_tmp_dir"))
    dir.create("ADaCGH2_vignette_tmp_dir")
> setwd("ADaCGH2_vignette_tmp_dir")
> ## creating ff objects
> fnameRdata <- list.files(path = system.file("data", package = "ADaCGH2"),
                       full.names = TRUE, pattern = "inputEx.RData")
```

```
> inputToADaCGH(ff.or.RAM = "ff",
                        RDatafilename = fnameRdata)
   ... done reading; starting checks
      ... checking identical MidPos
      ... checking need to reorder inputData, data.frame version
   ... done with checks; starting writing
   ... done writing/saving probeNames
   ... done writing/saving chromData
   ... done writing/saving posData
   ... done writing/saving cghData
 Calling gc at end
          used (Mb) gc trigger (Mb) max used (Mb)
Ncells 1565102 83.6
                       2403845 128.4 1835812 98.1
Vcells 1544792 11.8
                       2481603 19.0 1966371 15.1
Files saved in current directory
/home/ramon/tmp/ADaCGH2_vignette_tmp_dir
with names :
chromData.RData, posData.RData, cghData.RData, probeNames.RData.
> ## setting random number generator for forking
> RNGkind("L'Ecuyer-CMRG")
> ## initializing cluster and setting up random number generator
> number.of.nodes <- detectCores()</pre>
> cl2 <- parallel::makeCluster(number.of.nodes, "PSOCK")</pre>
> parallel::clusterSetRNGStream(cl2)
> parallel::setDefaultCluster(cl2)
> parallel::clusterEvalQ(NULL, library("ADaCGH2"))
[[1]]
                             "bit"
 [1] "ADaCGH2"
                 "ff"
                                                                  "methods"
                                          "tools"
                                                      "parallel"
 [7] "stats"
                 "graphics" "grDevices" "utils"
                                                      "datasets"
                                                                  "base"
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                                                                  "methods"
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                 "graphics" "grDevices" "utils"
                                                                  "base"
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	"ADaCGH2" "stats"	"ff" "graphics"	"bit" "grDevices"	"tools" "utils"	"parallel" "datasets"	"methods" "base"
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                                                                   "base"
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                                                       "datasets"
                                                                   "base"
                 "graphics"
[[64]]
                                          "tools"
[1] "ADaCGH2"
                 "ff"
                              "bit"
                                                       "parallel"
                                                                   "methods"
[7] "stats"
                 "graphics" "grDevices" "utils"
                                                       "datasets"
                                                                   "base"
> ## verify we are using the right version of ADaCGH2
> parallel::clusterEvalQ(NULL,
                         library(help = ADaCGH2)$info[[1]][[2]])
\lceil \lceil 1 \rceil \rceil
[1] "Version:
                    2.3.10"
[[2]]
[1] "Version:
                    2.3.10"
[[3]]
[1] "Version:
                    2.3.10"
[[4]]
[1] "Version:
                    2.3.10"
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[[5]]

[1] "Version:	2.3.10
[[6]] [1] "Version:	2.3.10
[[7]] [1] "Version:	2.3.10
[[8]] [1] "Version:	2.3.10
[[9]] [1] "Version:	2.3.10
[[10]] [1] "Version:	2.3.10
[[11]] [1] "Version:	2.3.10
[[12]] [1] "Version:	2.3.10
[[13]] [1] "Version:	2.3.10
[[14]] [1] "Version:	2.3.10
[[15]] [1] "Version:	2.3.10
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[[17]] [1] "Version:	2.3.10
[[18]] [1] "Version:	2.3.10
[[19]] [1] "Version:	2.3.10
[[20]] [1] "Version:	2.3.10
[[21]] [1] "Version:	2.3.10

[[22]]

[1] "Version:

2.3.10"

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[1] "Version: 2.3.10"

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[1] "Version: 2.3.10"

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[[43]] [1] "Version:	2.3.10"
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[1] "Version: 2.3.10"
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[1] "Version:
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[1] "Version:
                    2.3.10"
[[63]]
[1] "Version:
                    2.3.10"
[[64]]
[1] "Version:
                    2.3.10"
> wdir <- getwd()</pre>
> parallel::clusterExport(NULL, "wdir")
> parallel::clusterEvalQ(NULL, setwd(wdir))
[[1]]
[1] "/home/ramon/tmp/ADaCGH2_vignette_tmp_dir"
[[2]]
[1] "/home/ramon/tmp/ADaCGH2_vignette_tmp_dir"
[[3]]
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[1] "/home/ramon/tmp/ADaCGH2_vignette_tmp_dir"
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[1] "/home/ramon/tmp/ADaCGH2_vignette_tmp_dir"

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[[27]]
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[1] "/home/ramon/tmp/ADaCGH2_vignette_tmp_dir"
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[1] "/home/ramon/tmp/ADaCGH2_vignette_tmp_dir"
[[35]]
[1] "/home/ramon/tmp/ADaCGH2_vignette_tmp_dir"
[[36]]
[1] "/home/ramon/tmp/ADaCGH2_vignette_tmp_dir"
[[37]]
[1] "/home/ramon/tmp/ADaCGH2_vignette_tmp_dir"
[1] "/home/ramon/tmp/ADaCGH2_vignette_tmp_dir"
[[39]]
[1] "/home/ramon/tmp/ADaCGH2_vignette_tmp_dir"
[[40]]
[1] "/home/ramon/tmp/ADaCGH2_vignette_tmp_dir"
[[41]]
[1] "/home/ramon/tmp/ADaCGH2_vignette_tmp_dir"
[[42]]
[1] "/home/ramon/tmp/ADaCGH2_vignette_tmp_dir"
[[43]]
```

[1] "/home/ramon/tmp/ADaCGH2_vignette_tmp_dir"

```
[[44]]
[1] "/home/ramon/tmp/ADaCGH2_vignette_tmp_dir"
[[45]]
[1] "/home/ramon/tmp/ADaCGH2_vignette_tmp_dir"
[1] "/home/ramon/tmp/ADaCGH2_vignette_tmp_dir"
[[47]]
[1] "/home/ramon/tmp/ADaCGH2_vignette_tmp_dir"
[[48]]
[1] "/home/ramon/tmp/ADaCGH2_vignette_tmp_dir"
[[49]]
[1] "/home/ramon/tmp/ADaCGH2_vignette_tmp_dir"
[[50]]
[1] "/home/ramon/tmp/ADaCGH2_vignette_tmp_dir"
[[51]]
[1] "/home/ramon/tmp/ADaCGH2_vignette_tmp_dir"
[[52]]
[1] "/home/ramon/tmp/ADaCGH2_vignette_tmp_dir"
[[53]]
[1] "/home/ramon/tmp/ADaCGH2_vignette_tmp_dir"
[[54]]
[1] "/home/ramon/tmp/ADaCGH2_vignette_tmp_dir"
[[55]]
[1] "/home/ramon/tmp/ADaCGH2_vignette_tmp_dir"
[[56]]
[1] "/home/ramon/tmp/ADaCGH2_vignette_tmp_dir"
[[57]]
[1] "/home/ramon/tmp/ADaCGH2_vignette_tmp_dir"
[[58]]
[1] "/home/ramon/tmp/ADaCGH2_vignette_tmp_dir"
[[59]]
[1] "/home/ramon/tmp/ADaCGH2_vignette_tmp_dir"
[[60]]
[1] "/home/ramon/tmp/ADaCGH2_vignette_tmp_dir"
[[61]]
```

```
[1] "/home/ramon/tmp/ADaCGH2_vignette_tmp_dir"
[[62]]
[1] "/home/ramon/tmp/ADaCGH2_vignette_tmp_dir"
[[63]]
[1] "/home/ramon/tmp/ADaCGH2_vignette_tmp_dir"
[[64]]
[1] "/home/ramon/tmp/ADaCGH2_vignette_tmp_dir"
>
```

3 The examples

3.1 RAM objects and forking

```
> cbs.mergel.RAM.fork <- pSegmentDNAcopy(cgh.dat, chrom.dat,
                                              merging = "mergeLevels")
> cbs.mad.RAM.fork <- pSegmentDNAcopy(cgh.dat, chrom.dat,merging = "MAD")
> cbs.none.RAM.fork <- pSegmentDNAcopy(cgh.dat, chrom.dat, merging = "none")
> hmm.mergel.RAM.fork <- pSegmentHMM(cgh.dat, chrom.dat, merging = "mergeLevels")
> hmm.mad.RAM.fork <- pSegmentHMM(cgh.dat, chrom.dat, merging = "MAD")
> hs.mergel.RAM.fork <- pSegmentHaarSeg(cgh.dat, chrom.dat,
                               merging = "mergeLevels")
> hs.mad.RAM.fork <- pSegmentHaarSeg(cgh.dat, chrom.dat,
                               merging = "MAD")
> hs.none.RAM.fork <- pSegmentHaarSeg(cgh.dat, chrom.dat,
                               merging = "none")
> glad.RAM.fork <- pSegmentGLAD(cgh.dat, chrom.dat)
> biohmm.mergel.RAM.fork <- pSegmentBioHMM(cgh.dat,
                                  chrom.dat,
                                  pos.dat,
                                  merging = "mergeLevels")
> biohmm.mad.RAM.fork <- pSegmentBioHMM(cgh.dat,
                                           chrom.dat,
                                          pos.dat,
                                          merging = "MAD")
> biohmm.mad.bic.RAM.fork <- pSegmentBioHMM(cgh.dat,
                                               chrom.dat,
                                              pos.dat,
                                              merging = "MAD",
                                               aic.or.bic = "BIC")
> cghseg.mergel.RAM.fork <- pSegmentCGHseg(cgh.dat,
                                             chrom.dat,
                                             merging = "mergeLevels")
> cghseg.mad.RAM.fork <- pSegmentCGHseg(cgh.dat,
                                           chrom.dat,
                                           merging = "MAD")
> cghseg.none.RAM.fork <- pSegmentCGHseg(cgh.dat,
                                           chrom.dat,
                                           merging = "none")
```

3.2 ff objects and cluster

Compared to the section 3.1, the main differences are that we explicitly set the typeParall argument to "cluster" (the default is "fork") and the change in the names of the input data (which now refer to the names of the RData objects that contain the ff objects).

```
> cbs.mergel.ff.cluster <- pSegmentDNAcopy("cghData.RData", "chromData.RData",
                                    merging = "mergeLevels",
                                    typeParall = "cluster")
> cbs.mad.ff.cluster <- pSegmentDNAcopy("cghData.RData", "chromData.RData",
                                        merging = "MAD",
                                        typeParall = "cluster")
> cbs.none.ff.cluster <- pSegmentDNAcopy("cghData.RData", "chromData.RData",
                                         merging = "none",
                                          typeParall = "cluster")
> hmm.mergel.ff.cluster <- pSegmentHMM("cghData.RData", "chromData.RData",
                                merging = "mergeLevels",
                                typeParall = "cluster")
> hmm.mad.ff.cluster <- pSegmentHMM("cghData.RData", "chromData.RData",
                                    merging = "MAD",
                                    typeParall = "cluster")
> hs.mergel.ff.cluster <- pSegmentHaarSeg("cghData.RData", "chromData.RData",
                               merging = "mergeLevels",
                                       typeParall = "cluster")
> hs.mad.ff.cluster <- pSegmentHaarSeg("cghData.RData", "chromData.RData",
                               merging = "MAD", typeParall = "cluster")
> hs.none.ff.cluster <- pSegmentHaarSeg("cghData.RData", "chromData.RData",</pre>
                               merging = "none", typeParall = "cluster")
> glad.ff.cluster <- pSegmentGLAD("cghData.RData", "chromData.RData",
                                  typeParall = "cluster")
> biohmm.mergel.ff.cluster <- pSegmentBioHMM("cghData.RData",
                                   "chromData.RData",
                                   "posData.RData",
                                  merging = "mergeLevels",
                                       typeParall = "cluster")
> biohmm.mad.ff.cluster <- pSegmentBioHMM("cghData.RData",
                                           "chromData.RData",
                                           "posData.RData",
                                          merging = "MAD",
                                           typeParall = "cluster")
> biohmm.mad.bic.ff.cluster <- pSegmentBioHMM("cghData.RData",
                                               "chromData.RData",
                                               "posData.RData",
                                              merging = "MAD",
```

```
+
                                               aic.or.bic = "BIC",
                                               typeParall = "cluster")
> cghseg.mergel.ff.cluster <- pSegmentCGHseg("cghData.RData",
                                             "chromData.RData",
                                             merging = "mergeLevels",
                                             typeParall = "cluster")
> cghseg.mad.ff.cluster <- pSegmentCGHseg("cghData.RData",
                                           "chromData.RData",
                                           merging = "MAD",
                                           typeParall = "cluster")
> cghseg.none.ff.cluster <- pSegmentCGHseg("cghData.RData",
                                            "chromData.RData",
                                            merging = "none",
                                            typeParall = "cluster")
> waves.mergel.ff.cluster <- pSegmentWavelets("cghData.RData",
                                              "chromData.RData",
                                              merging = "mergeLevels",
                                              typeParall = "cluster")
> waves.mad.ff.cluster <- pSegmentWavelets("cghData.RData",
                                            "chromData.RData",
                                            merging = "MAD",
                                            typeParall = "cluster")
> waves.none.ff.cluster <- pSegmentWavelets("cghData.RData",
                                             "chromData.RData",
                                             merging = "none",
                                             typeParall = "cluster")
```

3.3 ff objects and forking

The main difference with section 3.2 is the argument typeParall; we did not need to pass it explicitly (since the default is fork), but we will do for clarity.

```
> cbs.mergel.ff.fork <- pSegmentDNAcopy("cghData.RData", "chromData.RData",
                                        merging = "mergeLevels",
                                    typeParall = "fork")
> cbs.mad.ff.fork <- pSegmentDNAcopy("cghData.RData", "chromData.RData",
                                     merging = "MAD",
                                        typeParall = "fork")
> cbs.none.ff.fork <- pSegmentDNAcopy("cghData.RData", "chromData.RData",
                                      merging = "none", typeParall = "fork")
> hmm.mergel.ff.fork <- pSegmentHMM("cghData.RData", "chromData.RData",
                             merging = "mergeLevels", typeParall = "fork")
> hmm.mad.ff.fork <- pSegmentHMM("cghData.RData", "chromData.RData",
                                 merging = "MAD", typeParall = "fork")
> hs.mergel.ff.fork <- pSegmentHaarSeg("cghData.RData", "chromData.RData",
                               merging = "mergeLevels", typeParall = "fork")
> hs.mad.ff.fork <- pSegmentHaarSeg("cghData.RData", "chromData.RData",
                               merging = "MAD", typeParall = "fork")
> hs.none.ff.fork <- pSegmentHaarSeg("cghData.RData", "chromData.RData",
                               merging = "none", typeParall = "fork")
> glad.ff.fork <- pSegmentGLAD("cghData.RData", "chromData.RData",</pre>
```

```
typeParall = "fork")
> biohmm.mergel.ff.fork <- pSegmentBioHMM("cghData.RData",
                                   "chromData.RData",
                                   "posData.RData",
                                   merging = "mergeLevels",
                                    typeParall = "fork")
> biohmm.mad.ff.fork <- pSegmentBioHMM("cghData.RData",
                                           "chromData.RData",
                                           "posData.RData",
                                           merging = "MAD",
                                        typeParall = "fork")
> biohmm.mad.bic.ff.fork <- pSegmentBioHMM("cghData.RData",
                                                "chromData.RData",
                                                "posData.RData",
                                               merging = "MAD",
                                               aic.or.bic = "BIC",
                                            typeParall = "fork")
> cghseg.mergel.ff.fork <- pSegmentCGHseg("cghData.RData",
                                              "chromData.RData",
                                             merging = "mergeLevels",
                                          typeParall = "fork")
> cghseg.mad.ff.fork <- pSegmentCGHseg("cghData.RData",</pre>
                                            "chromData.RData",
                                           merging = "MAD", typeParall = "fork")
 cghseg.none.ff.fork <- pSegmentCGHseg("cghData.RData",</pre>
                                            "chromData.RData",
                                            merging = "none", typeParall = "fork")
> waves.merge.ff.fork <- pSegmentWavelets("cghData.RData",
                                            "chromData.RData",
                                           merging = "mergeLevels",
                                           typeParall = "fork")
> waves.mad.ff.fork <- pSegmentWavelets("cghData.RData",
                                            "chromData.RData",
                                         merging = "MAD",
                                         typeParall = "fork")
> waves.none.ff.fork <- pSegmentWavelets("cghData.RData",
                                           "chromData.RData",
                                          merging = "none",
                                          typeParall = "fork")
```

3.4 Comparing output

Here we verify that using different input and clustering methods does not change the results. Before carrying out the comparisons, however, we open the ff objects gently.

First, we will open the objects created above (same objects as were also created in the main vignette, in section "Carrying out segmentation and calling"). Instead of inserting many calls to each individual object, we open all available objects that match ff.cluster. To do that quickly we store the names of the objects

```
> ff.cluster.obj <- ls(pattern = "*.ff.cluster")</pre>
```

pages with the string "TRUE")

```
> tmpout <-
+ capture.output(
+ lapply(ff.cluster.obj, function(x) lapply(get(x), open))
+ )

We repeat that operation with the output from section 3.3:

> ff.fork.obj <- ls(pattern = "*.ff.fork")
> tmpout <-
+ capture.output(
+ lapply(ff.fork.obj, function(x) lapply(get(x), open))
+ )
>
```

And we create the list of results from the RAM and forking runs (no need for special opening here, since these are not ff objects)

```
> RAM.fork.obj <- ls(pattern = "*.RAM.fork")
```

We can now compare the output. We want to compare the output from three different methods, so we need to run three comparisons (this is what we did explicitly in the help for pSegment). Since this is a very repetitive operation, we define a small utility function that will return TRUE if both components (outSmoothed and outState) of all three objects are identical. (Since the function will take as input not an actual object, but a name, we use get inside the function.)

We use all.equal to compare the output from the smoothing, to allow for possible numerical fuzz (that could result from differences in storage). When comparing the assigned state, however, we check for exact identity.

```
> identical3 <- function(x, y, z) {</pre>
    comp1 <- all.equal(get(x)$outSmoothed[ , ], get(y)$outSmoothed[ , ])</pre>
    comp2 <- all.equal(get(y)$outSmoothed[ , ], get(z)$outSmoothed[ , ])</pre>
    comp3 <- identical(get(x)$outState[ , ], get(y)$outState[ , ])</pre>
    comp4 <- identical(get(y)$outState[ , ], get(z)$outState[ , ])</pre>
    if (!all(isTRUE(comp1), isTRUE(comp2), comp3, comp4)) {
      cat(paste("Comparing ", x, y, z, "\n",
             "not equal: some info from comparisons.\n",
                 "\n comp1 = ", paste(comp1, sep = " ", collapse = "\n
                                                                               "),
                 "\n comp2 = ", paste(comp2, sep = " ", collapse = "\n
                                                                               "),
                 "\n comp3 = ", paste(comp3, sep = " ", collapse = "\n
                                                                               "),
                 "\n comp4 = ", paste(comp4, sep = " ", collapse = "\n
                                                                               "),
                 "\langle n \rangle ")
      return (FALSE)
+
    } else {
      TRUE
    }
+ }
```

You should expect most (though not necessarily all) the comparisons to yield a TRUE. In some cases, however, different runs of the same method might not yield the same results (e.g., CBS, HMM, etc). If you get non-identical results, you can try running those methods a few

times, to check for differences. You can also disable load balancing, and try using reproducible streams for the random number generators (see the vignette of package **parallel**).

Let's check those results then:

>

```
> mapply(identical3, RAM.fork.obj,
             ff.fork.obj, ff.cluster.obj)
Comparing cbs.none.RAM.fork cbs.none.ff.fork cbs.none.ff.cluster
not equal: some info from comparisons.
 comp1 =
         TRUE
          Component 4: Mean relative difference: 0.03339901
 comp3 =
         TRUE
 comp4 = FALSE
Comparing glad.RAM.fork glad.ff.fork glad.ff.cluster
not equal: some info from comparisons.
comp1 =
         TRUE
 comp2 =
          Component 4: Mean relative difference: 0.5858093
 comp3 =
         TRUE
 comp4 = FALSE
biohmm.mad.bic.RAM.fork
                            biohmm.mad.RAM.fork
                                                  biohmm.mergel.RAM.fork
                   TRUE
                                            TRUE
                                                                     TRUE
       cbs.mad.RAM.fork
                            cbs.mergel.RAM.fork
                                                       cbs.none.RAM.fork
                   TRUE
                                            TRUE
                                                                   FALSE
    cghseg.mad.RAM.fork
                         cghseg.mergel.RAM.fork
                                                    cghseg.none.RAM.fork
                   TRUE
                               hmm.mad.RAM.fork
          glad.RAM.fork
                                                     hmm.mergel.RAM.fork
                  FALSE
                                            TRUE
                                                                     TRUE
        hs.mad.RAM.fork
                                                        hs.none.RAM.fork
                             hs.mergel.RAM.fork
                   TRUE
                                            TRUE
                                                                     TRUE
     waves.mad.RAM.fork
                          waves.mergel.RAM.fork
                                                     waves.none.RAM.fork
                   TRUE
                                            TRUE
                                                                     TRUE
```

(Of course, we depend on the lists of names of objects having the output from the same method and option in the same position, which is the case in these examples).

4 Exercising the code for the load balancing options

This section simply exercises the load balancing options. We use Haar as it is the fastest method, and one unlikely to be affected by the order in which different columns are run (in contrast to, say, HMM), so we need not worry about random numbers here.

```
[1] TRUE
> hs.none.ff.cluster <- pSegmentHaarSeg("cghData.RData", "chromData.RData",</pre>
                                        merging = "none", typeParall = "cluster")
> hs.none.ff.cluster.lb <- pSegmentHaarSeg("cghData.RData", "chromData.RData",
                                        merging = "none", typeParall = "cluster",
                                        loadBalance = TRUE)
> hs.none.ff.cluster.nlb <- pSegmentHaarSeg("cghData.RData", "chromData.RData",</pre>
                                        merging = "none", typeParall = "cluster",
                                        loadBalance = FALSE)
> ## do not show all the opening ... messages
> tmpout <-
   capture.output(
      lapply("hs.none.ff.cluster", function(x) lapply(get(x), open))
> tmpout <-
   capture.output(
      lapply("hs.none.ff.cluster.lb", function(x) lapply(get(x), open))
> tmpout <-
   capture.output(
      lapply("hs.none.ff.cluster.nlb", function(x) lapply(get(x), open))
> identical3("hs.none.ff.cluster", "hs.none.ff.cluster.lb",
             "hs.none.ff.cluster.nlb")
[1] TRUE
> hs.none.ff.fork <- pSegmentHaarSeg("cghData.RData", "chromData.RData",
                               merging = "none", typeParall = "fork")
> hs.none.ff.fork.lb <- pSegmentHaarSeg("cghData.RData", "chromData.RData",
                                     merging = "none", typeParall = "fork",
                                      loadBalance = TRUE)
> hs.none.ff.fork.nlb <- pSegmentHaarSeg("cghData.RData", "chromData.RData",
                                     merging = "none", typeParall = "fork",
                                     loadBalance = FALSE)
> tmpout <-
   capture.output(
      lapply("hs.none.ff.fork", function(x) lapply(get(x), open))
> tmpout <-
   capture.output(
      lapply("hs.none.ff.fork.lb", function(x) lapply(get(x), open))
      )
> tmpout <-
```

+ lapply("hs.none.ff.fork.nlb", function(x) lapply(get(x), open))
+)
> identical3("hs.none.ff.fork", "hs.none.ff.fork.lb", "hs.none.ff.fork.nlb")

[1] TRUE

capture.output(

>

(There is no need to compare between ff.fork, ff.cluster, RAM.fork, as those were already shown to be identical.)

5 Clean up actions

These are not strictly necessary, but we will explicitly stop the cluster. In this vignette, we will not execute the code below to remove the directory we created or the objects, in case you want to check them out or play around with them, but the code is below.

To make sure there are no file permission problems, we add code below to explicitly delete some of the "ff" files and objects (and we wait a few seconds to allow pending I/O operations to happen before we delete the directory).

```
> parallel::stopCluster(cl2)
> ## This is the code to remove all the files we created
> ## and the temporary directory.
> ## We are not executing it!
> load("chromData.RData")
> load("posData.RData")
> load("cghData.RData")
> delete(cghData); rm(cghData)
> delete(posData); rm(posData)
> delete(chromData); rm(chromData)
> tmpout <-
    capture.output(
      lapply(ff.fork.obj, function(x) {
        lapply(get(x), delete)}))
> rm(list = ff.fork.obj)
> tmpout <-
    capture.output(
      lapply(ff.cluster.obj, function(x) {
        lapply(get(x), delete)}))
> rm(list = ff.cluster.obj)
> setwd(originalDir)
> print(getwd())
> Sys.sleep(3)
> unlink("ADaCGH2_vignette_tmp_dir", recursive = TRUE)
> Sys.sleep(3)
```