ExomeDepth

Vincent Plagnol

December 11, 2012

Contents

1	What ExomeDepth does and tips for QC 1.1 What ExomeDepth does and does not do									
2	Create count data from BAM files 2.1 Count for the autosomes									
3	Load an example dataset									
4	Build the most appropriate reference set	3								
5	CNV calling									
6	Better annotation of CNV calls									
7	A visual example	6								
8	How to loop over the multiple samples	7								
9	Technical information about R session	8								

1 What ExomeDepth does and tips for QC

1.1 What ExomeDepth does and does not do

ExomeDepth uses read depth data to call CNVs from exome sequencing experiments. A key idea is that the test exome should be compared to a matched aggregate reference set. This aggregate reference set should combine exomes from the same batch and it should also be optimized for each exome. It will certainly differ from one exome to the next.

Importantly, ExomeDepth assumes that the CNV of interest is absent from the aggregate reference set. Hence related individuals should be excluded from the aggregate reference. It also means that ExomeDepth can miss common CNVs, if the call is also present in the aggregate reference. ExomeDepth is really suited to detect rare CNV calls (typically for rare Mendelian disorder analysis).

The ideas used in this package are of course not specific to exome sequencing and could be applied to other targeted sequencing datasets, as long as they contain a sufficiently large number of exons to estimate the parameters (at least 20 genes, say, but probably more would be useful). Also note that PCR based enrichment studies are often not well suited for this type of read depth analysis. The reason is that as the number of cycles is often set to a high number in order to equalize the representation of each amplicon, which can discard the CNV information.

1.2 Useful quality checks

Just to give some general expectations I usually obtain 150-280 CNV calls per exome sample (two third of them deletions). Any number clearly outside of this range is suspicious and suggests that either the model was inappropriate or that something went wrong while running the code. Less important and less precise, I also expect the

aggregate reference to contain 5-10 exome samples. While there is no set rule for this number, and the code may work very well with fewer exomes in the aggregate reference set, numbers outside of this range suggest potential technical artifacts.

2 Create count data from BAM files

2.1 Count for the autosomes

Firstly, to facilitate the generation of read count data, exon positions for the hg19 build of the human genome are available within ExomeDepth. This exons.hg19 data frame can be directly passed as an argument of getBAMCounts (see below).

```
> library(ExomeDepth)
> data(exons.hg19)
> print(head(exons.hg19))
  chromosome start
                     end
                                    name
           1 12011 12058 DDX11L10-201_1
1
2
           1 12180 12228 DDX11L10-201 2
3
           1 12614 12698 DDX11L10-201_3
4
           1 12976 13053 DDX11L10-201_4
5
           1 13222 13375 DDX11L10-201_5
6
           1 13454 13671 DDX11L10-201_6
```

To generate read count data, the function getBamCounts in ExomeDepth is set up to parse the BAM files. It generates an array of read count, stored in a GenomicRanges object. It is a wrapper around the function countBamInGRanges.exomeDepth which is derived from an equivalent function in the exomeCopy package. You can refer to the help page of getBAMCounts to obtain the full list of options. An example line of code (not evaluated here) would look like this:

my.bam is a set character vector of indexed BAM files. fasta is the reference genome in fasta format (only useful if one wants to obtain the GC content). exons.hg19 are the positions and names of the exons on the hg19 reference genome (as shown above). include.chr defaults to false: if the BAM file are aligned to a reference sequence with the convention chr1 for chromosomes instead of simply 1 (i.e. the UCSC convention vs. the Ensembl one) you need to set include.chr = TRUE, otherwise the counts will be equal to 0. Note that the data frame with exon locations provided with ExomeDepth uses the Ensembl location (i.e. no chr prefix before the chromosome names) but what matters to set this option is how the BAM files were aligned.

getBAMCounts creates an object of the GRanges class which can easily be converted into a matrix or a data frame (which is the input format for ExomeDepth). An example of GenomicRanges output generated by getBAMCounts is provided in this package (chromosome 1 only to keep the size manageable). Here is how this object could for example be used to obtain a more generic data frame:

```
> library(ExomeDepth)
> data(ExomeCount)
> ExomeCount.dafr <- as(ExomeCount[, colnames(ExomeCount)], 'data.frame')
> ExomeCount.dafr$chromosome <- gsub(as.character(ExomeCount.dafr$space),
                                           pattern = 'chr',
                                           replacement = '')
                                                              ##remove the annoying chr letters
> print(head(ExomeCount.dafr))
                end width
                                                 GC Exome1 Exome2 Exome3 Exome4
  space start
                                   names
                                                         0
      1 12012 12058
                       47 DDX11L10-201_1 0.6170213
                                                                0
                                                                        0
                                                                               0
```

```
2
      1 12181 12228
                         48 DDX11L10-201_2 0.5000000
                                                              0
                                                                      0
                                                                             0
                                                                                     0
3
                         84 DDX11L10-201_3 0.5952381
                                                                    242
      1 12615 12698
                                                            118
                                                                           116
                                                                                   170
4
      1 12977 13053
                         77 DDX11L10-201_4 0.6103896
                                                            198
                                                                    48
                                                                           104
                                                                                   118
5
                        153 DDX11L10-201_5 0.5882353
                                                                           530
      1 13223 13375
                                                            516
                                                                   1112
                                                                                   682
6
      1 13455 13671
                        217 DDX11L10-201_6 0.5898618
                                                            272
                                                                    762
                                                                           336
                                                                                   372
  chromosome
1
            1
2
            1
3
            1
4
            1
5
            1
6
            1
```

2.2 Counts for chromosome X

Calling CNVs on the X chromosome can create issues if the exome sample of interest and the reference exome samples it is being compared to (what I call the aggregate reference) are not gender matched. For this reason the chromosome X exonic regions are not included by default in the data frame exons.hg19, mostly to avoid users getting low quality CNV calls because of this issue. However, loading the same dataset in R also brings another object called exons.hg19.X that containts the chromosome X exons.

```
> data(exons.hg19.X)
> head(exons.hg19.X)
       chromosome start
                             end
                                             name
238055
                X 170411 170514 NCRNA00108-001_1
238056
                X 171605 171759 NCRNA00108-001_2
238057
                X 172683 172713 NCRNA00108-001_3
238058
                X 192992 193062
                                     PLCXD1-201_1
238059
                X 198150 198352
                                     PLCXD1-201_2
238060
                X 200835 200982
                                     PLCXD1-201_3
```

This object can be used to generate CNV counts and further down the line CNV calls, in the same way as exons.hg19. While this is not really necessary, I would recommend calling CNV on the X separately from the autosomes. Also make sure that the genders are matched properly (i.e. do not use male as a reference for female samples and vice versa).

3 Load an example dataset

We have already loaded a dataset of chromosome 1 data for four exome samples. We run a first test to make sure that the model can be fitted properly. Note the use of the subset.for.speed option that subsets some rows purely to speed up this computation.

4 Build the most appropriate reference set

A key idea behing ExomeDepth is that each exome should not be compared to all other exomes but rather to an optimized set of exomes that are well correlated with that exome. This is what I call the optimized aggregate reference set, which is optimized for each exome sample. So the first step is to select the most appropriate reference sample. This step is demonstrated below.

Note that the drop = FALSE option is just used in case the reference set contains a single sample. If this is the case, it makes sure that the subsetted object is a data frame, not a numeric vector.

5 CNV calling

4

575

576

deletion

> all.exons <- new('ExomeDepth',

test = my.test,

Now the following step is the longest one as the beta-binomial model is applied to the full set of exons:

```
reference = my.reference.selected,
                    formula = 'cbind(test, reference) ~ 1')
   We can now call the CNV by running the underlying hidden Markov model:
> all.exons <- CallCNVs(x = all.exons,</pre>
                         transition.probability = 10^-4,
                         chromosome = ExomeCount.dafr$space,
                         start = ExomeCount.dafr$start,
                         end = ExomeCount.dafr$end,
                         name = ExomeCount.dafr$names)
Number of hidden states: 3
Number of data points: 26547
Initializing the HMM
Done with the first step of the HMM, now running the trace back
Total number of calls: 23
> print(head(all.exons@CNV.calls))
                                                    end chromosome
  start.p end.p
                        type nexons
                                        start
       25
                                        89553
1
             27
                    deletion
                                                 91106
2
       52
             66
                    deletion
                                  15
                                       324290
                                                523834
                                                                 1
3
      100
            103 duplication
                                   4
                                       743956
                                                745551
                                                                 1
```

2 1569583

1570002

5	587	591	deleti	ion	5	1592941	1	603069		1	
6	2324	2327	deleti	ion	4	12976452	12	980570		1	
			id	BF	read	ds.expecte	ed	reads.	observed	reads.rat	io
1	ch	r1:895	53-91106	12.40		22	24		68	0.3	04
2	chr1	:324290	0-523834	13.40		38	30		190	0.5	00
3	chr1	:743956	3-745551	7.67		20	01		336	1.6	70
4	chr1:1	569583-	-1570002	5.53		6	86		24	0.3	53
5	chr1:1	592941-	-1603069	13.90		113	36		434	0.3	82
6	chr1:129	76452-1	12980570	12.10		78	30		342	0.4	38

Now the last thing to do is to save it in an easily readable format (csv in this example, which can be opened in excel if needed):

Note that it is probably best to annotate the calls before creating that csv file (see below for annotation tools).

6 Better annotation of CNV calls

Much can be done to annotate CNV calls and this is an open problem. While this is a work in progress, I have started adding basic options. Importantly, the key function uses the more recent syntax from the package GenomicRanges. Hence the function will only return a warning and not add the annotations if you use a version of GenomicRanges prior to 1.8.10. The best way to upgrade is probably to use R 2.15.0 or later and let Bioconductor scripts do the install for you. If you use R 2.14 or earlier the annotation steps described below will probably only return a warning and not annotate the calls.

Perhaps the most useful step is to identify the overlap with a set of common CNVs identified in Conrad et al, Nature 2010. If one is looking for rare CNVs, these should probably be filtered out. The first step is to load these reference data from Conrad et al. To make things as practical as possible, these data are now available as part of ExomeDepth.

```
> data(Conrad.hg19)
> head(Conrad.hg19.common.CNVs)
```

GRanges with 6 ranges and 1 elementMetadata col:

```
seqnames
                       ranges strand |
                                           names
       <Rle>
                               <Rle> | <factor>
                    <IRanges>
[1]
           1 [10499, 91591]
                                         CNVR1.1
[2]
           1 [10499, 177368]
                                         CNVR1.2
[3]
           1 [82705, 92162]
                                         CNVR1.5
[4]
           1 [85841,
                     91967]
                                         CNVR1.4
[5]
           1 [87433, 89163]
                                         CNVR1.6
[6]
           1 [87446, 109121]
                                         CNVR1.7
```

seqlengths:

Then one can use this information to annotate our CNV calls with the function AnnotateExtra.

The min.overlap argument set to 0.5 requires that the Conrad reference call overlaps at least 50% of our CNV calls to declare an overlap. The column.name argument simply defines the name of the column that will store the overlap information. The outcome of this procedure can be checked with:

```
> print(head(all.exons@CNV.calls))
  start.p end.p
                                                    end chromosome
                        type nexons
                                        start
       25
1
             27
                    deletion
                                   3
                                        89553
                                                  91106
2
       52
                                  15
                                       324290
                                                 523834
                                                                  1
             66
                    deletion
3
      100
            103 duplication
                                  4
                                       743956
                                                 745551
                                                                  1
4
      575
            576
                    deletion
                                   2 1569583
                                               1570002
                                                                  1
5
      587
            591
                    deletion
                                   5 1592941
                                               1603069
6
     2324
                                   4 12976452 12980570
           2327
                    deletion
                                                                  1
                       id
                             BF reads.expected reads.observed reads.ratio
        chr1:89553-91106 12.40
                                             224
                                                                       0.304
1
                                                              68
2
      chr1:324290-523834 13.40
                                             380
                                                             190
                                                                       0.500
3
      chr1:743956-745551 7.67
                                             201
                                                             336
                                                                       1.670
    chr1:1569583-1570002 5.53
4
                                              68
                                                             24
                                                                       0.353
5
    chr1:1592941-1603069 13.90
                                           1136
                                                             434
                                                                       0.382
6 chr1:12976452-12980570 12.10
                                             780
                                                             342
                                                                       0.438
                                Conrad.hg19
1 CNVR1.1, CNVR1.2, CNVR1.5, CNVR1.4, CNVR1.7
2
                                    CNVR2.4
3
                                       <NA>
4
                                   CNVR17.1
5
                                   CNVR17.1
6
                CNVR72.3, CNVR72.4, CNVR72.2
```

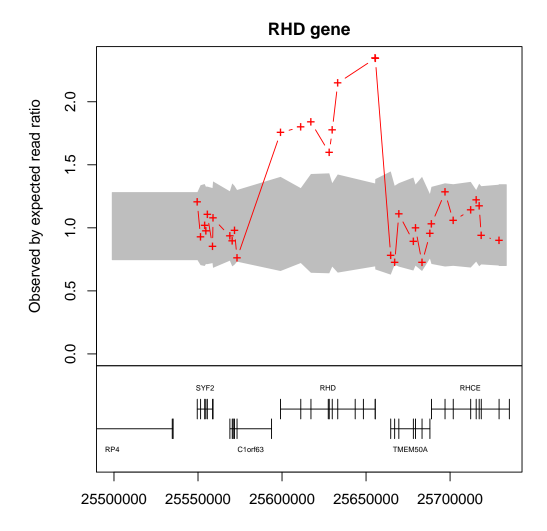
I have processed the Conrad et al data in the GRanges format. Potentially any other reference dataset could be converted as well. See for example the exon information:

```
exons.hg19.GRanges <- GRanges(seqnames = exons.hg19$chromosome,
+
                                   IRanges(start=exons.hg19$start,end=exons.hg19$end),
                                   names = exons.hg19$name)
 all.exons <- AnnotateExtra(x = all.exons,
                              reference.annotation = exons.hg19.GRanges,
                              min.overlap = 0.0001,
+
                              column.name = 'exons.hg19')
> all.exons@CNV.calls[3:6,]
  start.p end.p
                       type nexons
                                       start
                                                   end chromosome
3
      100
            103 duplication
                                  4
                                      743956
                                               745551
                                                                1
4
      575
            576
                   deletion
                                  2 1569583 1570002
                                                                1
5
      587
            591
                   deletion
                                  5 1592941 1603069
                                                                1
6
     2324
                                  4 12976452 12980570
           2327
                   deletion
                                                                1
                            BF reads.expected reads.observed reads.ratio
                      id
3
      chr1:743956-745551 7.67
                                           201
                                                           336
                                                                     1.670
4
    chr1:1569583-1570002 5.53
                                            68
                                                            24
                                                                     0.353
5
    chr1:1592941-1603069 13.90
                                          1136
                                                           434
                                                                     0.382
6 chr1:12976452-12980570 12.10
                                           780
                                                           342
                                                                     0.438
                 Conrad.hg19
3
                         <NA>
4
                    CNVR17.1
5
                    CNVR17.1
6 CNVR72.3, CNVR72.4, CNVR72.2
                                                                                exons.hg19
3
                ENST00000447500_1, ENST00000447500_2, ENST00000412115_3, ENST00000435300_2
                                                                MMP23B-202_8,MMP23B-202_9
5 RP11-345P4-003_6,RP11-345P4-003_7,RP11-345P4-003_8,RP11-345P4-003_9,RP11-345P4-003_10
                                 PRAMEF7-201_1, PRAMEF7-201_2, PRAMEF7-201_3, PRAMEF7-201_4
6
```

This time I report any overlap with an exon by specifying a value close to 0 in the min.overlap argument. Note the metadata column names which MUST be specified for the annotation to work properly.

7 A visual example

The ExomeDepth object includes a plot function. This function shows the ratio between observed and expected read depth. The 95% confidence interval is marked by a grey shaded area. Here we use a common CNV located in the *RHD* gene as an example. We can see that the individual in question has more copies than the average (in fact two functional copies of *RHD*, which corresponds to rhesus positive).



8 How to loop over the multiple samples

A FAQ is a way to deal with a set of a significant number of exomes, i.e. how to loop over all of them using ExomeDepth. This can be done with a loop. I show below an example of how I would code things. The code is not executed in the vignette to save time when building the package, but it can give some hints to users who do not have extensive experience with R.

```
> #### get the annotation datasets to be used later
> data(Conrad.hg19)
```

```
> exons.hg19.GRanges <- GRanges(seqnames = exons.hg19$chromosome,
                                IRanges(start=exons.hg19$start,end=exons.hg19$end),
                                names = exons.hg19$name)
> ### prepare the main matrix of read count data
> ExomeCount.mat <- as.matrix(ExomeCount.dafr[, grep(names(ExomeCount.dafr), pattern = 'Exome.*')])
> nsamples <- ncol(ExomeCount.mat)</pre>
> ### start looping over each sample
> for (i in 1:nsamples) {
+ #### Create the aggregate reference set for this sample
    my.choice <- select.reference.set (test.counts = ExomeCount.mat[,i],
                                        reference.counts = ExomeCount.mat[,-i],
                                        bin.length = (ExomeCount.dafr$end - ExomeCount.dafr$start)/1000,
+
                                        n.bins.reduced = 10000)
   my.reference.selected <- apply(X = ExomeCount.mat[, my.choice$reference.choice, drop = FALSE],</pre>
                                   MAR = 1,
                                    FUN = sum)
    message('Now creating the ExomeDepth object')
    all.exons <- new('ExomeDepth',</pre>
                     test = ExomeCount.mat[,i],
                     reference = my.reference.selected,
                     formula = 'cbind(test, reference) ~ 1')
+ ############ Now call the CNVs
    all.exons <- CallCNVs(x = all.exons,
                          transition.probability = 10^-4,
                          chromosome = ExomeCount.dafr$space,
                          start = ExomeCount.dafr$start,
+
                          end = ExomeCount.dafr$end,
                          name = ExomeCount.dafr$names)
+ ###################### Now annotate the ExomeDepth object
    all.exons <- AnnotateExtra(x = all.exons,
                               reference.annotation = Conrad.hg19.common.CNVs,
                               min.overlap = 0.5,
                                column.name = 'Conrad.hg19')
   all.exons <- AnnotateExtra(x = all.exons,
                               reference.annotation = exons.hg19.GRanges,
                                min.overlap = 0.0001,
+
                               column.name = 'exons.hg19')
+
   output.file <- paste('Exome_', i, 'csv', sep = '')</pre>
    write.csv(file = output.file, x = all.exons@CNV.calls, row.names = FALSE)
+
+ }
>
```

9 Technical information about R session

```
> sessionInfo()
R version 2.15.1 (2012-06-22)
Platform: x86_64-unknown-linux-gnu (64-bit)
```

locale:

[5] LC_MONETARY=en_US.iso885915 LC_MESSAGES=en_US.iso885915

[7] LC_PAPER=C LC_NAME=C
[9] LC_ADDRESS=C LC_TELEPHONE=C

[11] LC_MEASUREMENT=en_US.iso885915 LC_IDENTIFICATION=C

attached base packages:

[1] stats4 splines stats graphics grDevices utils datasets

[8] methods base

other attached packages:

[1] ExomeDepth_0.9.7 Rsamtools_1.8.5 Biostrings_2.24.1 [4] GenomicRanges_1.8.11 IRanges_1.14.4 BiocGenerics_0.2.0

[7] VGAM_0.8-7 aod_1.3

loaded via a namespace (and not attached):

[1] bitops_1.0-4.1 tools_2.15.1 zlibbioc_1.2.0