# Introduction to GeneBreak

Evert van den Broek & Stef van Lieshout

July 13, 2015

# 1 Running GeneBreak

This is a short tutorial on how to use the GeneBreak package. It covers an example run using included copy number data of one chromosome from 200 samples. The samples are part of the CAIRO study described in REF. Let's start by loading the package.

> library(GeneBreak)

### 1.1 Loading copy number data

Copy number data can be loaded in two ways. Either from a cghCall/QDNAseq object (ouput of bioconductor packages CGHcall or QDNAseq) or by providing a data.frame with at least 5 columns: Chromosome, Start, End and FeatureName (usually probe identifier). Note: when using the data.frame input the column names must be exactly as described here and in the same order! In this tutorial we will use a built-in dataset of chromosome 20:

```
> data( "copynumber.data.chr20" )
```

By inspecting the dataset, we see that we are dealing with an R object of class "cghCall" with 3653 features (aCGH probes in this case) and 200 samples.

```
> copynumber.data.chr20
cghCall (storageMode: lockedEnvironment)
assayData: 3653 features, 200 samples
  element names: calls, copynumber, probamp, probgain, probloss, probnorm, segmented
protocolData: none
phenoData
  sampleNames: sample_1 sample_2 ... sample_200 (200 total)
  varLabels: Cellularity
  varMetadata: labelDescription
featureData
  featureNames: A_16_P03469195 A_14_P136138 ... A_18_P13856091 (3653 total)
  fvarLabels: Chromosome Start End
```

```
fvarMetadata: labelDescription
experimentData: use 'experimentData(object)'
Annotation:
```

# 1.2 Obtaining data from a cghCall object (ouput from packages CGHcall or QDNAseq)

Then we need to obtain copy number data.

```
> breakpoints <- getBreakpoints( data = copynumber.data.chr20 )
Breakpoint detection started...</pre>
```

#### 1.3 Loading gene annotation data

Then we need to obtain gene annotations. For hg18 (and hg19, hg38) reference sequence these are included and can be loaded:

```
> data( "ens.gene.ann.hg18" )
```

Inspect the annotation.

#### > head( ens.gene.ann.hg18 )

Gene	EnsID	${\it Chromosome}$	Start	End	band	strand
MIRN1302-2	ENSG00000221311	1	20229	20366	p36.33	1
FAM138E	ENSG00000222027	1	24417	25944	p36.33	-1
FAM138E	ENSG00000222003	1	24417	25944	p36.33	-1
FAM138A	ENSG00000222003	1	24417	25944	p36.33	-1
OR4F5	ENSG00000177693	1	58954	59871	p36.33	1
0R4F29	ENSG00000177799	1	357522	358460	p36.33	1

#### 1.4 Filtering

Next we filter breakpoints. Different filters can be set with different threshold. Default here is "deltaSeg" filter with a threshold of 0.2. This means that only breakpoints which...

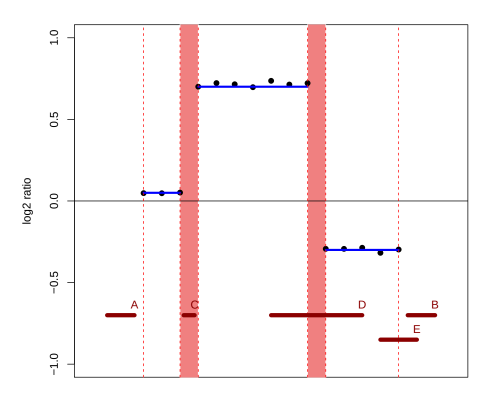
```
> breakpointsFiltered <- bpFilter( breakpoints )</pre>
```

```
Applying BP selection...
```

Next we will add the gene annotation information to the GeneBreak object. No analysis is done here yet.

```
> breakpointsAnnotated <- addGeneAnnotation( breakpointsFiltered, ens.gene.ann.hg18 )
```

```
Adding of gene annotation started on 659 genes by 200 samples 0% \dots 25% \dots 50% \dots 75% \dots Adding gene annotation DONE
```



Chromosome

Figure 1: GeneBreak geneAnnotation

Next we perform the gene analysis. This overlaps the genomic locations of the genes with the copy number data to find breakpoints within genes.

#### > breakpointGenes <- bpGenes( breakpointsAnnotated )</pre>

```
Running bpGenes: 659 genes and 200 samples 0% ... 25% ... 50% ... 75% ... bpGenes DONE A total of 1029 gene breaks in 241 genes detected
```

Next we determine the significantly recurring breakpoints. This be done at the "gene" or "feature" level and using one of two different methods ("Benjamini Hochberg" or "Gilbert"). The advantage of using...

#### > breakpointStatistics <- bpStats( breakpointGenes )</pre>

Applying statistical test over 200 samples for: gene breakpoints: BH test...

#### > breakpointStatistics

This will return an object of class CopyNumberBreakPointGenes. By using recurrentGenes() we can observe the recurrent affected genes.

#### > head( recurrentGenes( breakpointStatistics ) )

A total of 14 recurrent breakpoint genes (at FDR < 0.1)

	Gene s	samp]	LeCount	feature	eTotal
13886	PCMTD2		64		4
13898	C20orf69		33		3
4268	BFSP1		8		5
5473	ABHD12		10		9
4780	C20orf26		7		18
3493	HAO1		5		5
	pva	alue		FDR	
13886	1.350385e	-103	8.89903	35e-101	
13898	5.522293	e-44	1.819	595e-41	
4268	3.941447	e-07	8.6580	045e-05	
5473	5.756361	e-05	9.4836	605e-03	
4780	2.748743	e-04	3.6228	343e-02	
3493	6.528175	e-04	3.9617	727e-02	

## 2 Gene Annotation creation

The steps below are a description of how the gene annotations in this package were created.

```
> # gene annotations obtained via Biomart.
> # HUGO gene names (HGNC symbol), Ensembl_ID and chromosomal location
> # Used (and most) recent releases:
> # HG18: release54
> # HG19: release75
> # HG38: release80 (date: 150629)
> library(biomaRt)
> ensembl54 = useMart( host='may2009.archive.ensembl.org', biomart='ENSEMBL_MART_ENSEMBL', o
> ensembl75 = useMart( host='feb2014.archive.ensembl.org', biomart='ENSEMBL_MART_ENSEMBL', o
> ensembl80 = useMart( "ensembl", dataset="hsapiens_gene_ensembl" )
> createAnnotationFile <- function( biomartVersion ) {</pre>
    biomart_result <- getBM(attributes = c("hgnc_symbol", "ensembl_gene_id", "chromosome_na
    biomart_result[ ,3] <- as.vector( biomart_result[ ,3] )</pre>
    biomart_result$chromosome_name[ biomart_result$chromosome_name=="X" ] <- "23"
    biomart_result$chromosome_name[ biomart_result$chromosome_name=="Y" ] <- "24"
    biomart_genes <-biomart_result[ which(biomart_result[ ,1]!="" & biomart_result[ ,3] %in
    colnames(biomart_genes)[1:5]<-c("Gene", "EnsID", "Chromosome", "Start", "End")</pre>
    cat( c("Biomart version:", biomartVersion@host, "including:", dim(biomart_genes)[1], "ge
    return( biomart_genes )
+ }
> ens.gene.ann.hg18 <- createAnnotationFile( ensembl54 )</pre>
> ens.gene.ann.hg19 <- createAnnotationFile( ensemb175 )</pre>
> ens.gene.ann.hg38 <- createAnnotationFile( ensembl80 )</pre>
```

## 3 Session Information

R version 3.1.1 (2014-07-10)

The version number of R and packages loaded for generating the vignette were:

```
Platform: x86_64-pc-linux-gnu (64-bit)
locale:
 [1] LC_CTYPE=en_US.UTF-8
 [2] LC_NUMERIC=C
 [3] LC_TIME=en_US.UTF-8
 [4] LC_COLLATE=en_US.UTF-8
 [5] LC_MONETARY=en_US.UTF-8
 [6] LC_MESSAGES=en_US.UTF-8
 [7] LC_PAPER=en_US.UTF-8
 [8] LC_NAME=C
 [9] LC_ADDRESS=C
[10] LC_TELEPHONE=C
[11] LC_MEASUREMENT=en_US.UTF-8
[12] LC_IDENTIFICATION=C
attached base packages:
[1] parallel stats
                        graphics
[4] grDevices utils
                        datasets
[7] methods base
other attached packages:
[1] GeneBreak_0.99.0
[2] CGHbase_1.26.0
[3] marray_1.44.0
[4] limma_3.22.7
[5] Biobase_2.26.0
[6] BiocGenerics_0.12.1
loaded via a namespace (and not attached):
[1] devtools_1.7.0 digest_0.6.8
[3] magrittr_1.5 Rcpp_0.11.6
[5] roxygen2_4.1.1 stringi_0.4-1
[7] stringr_1.0.0 tools_3.1.1
```