VanillaICE: Hidden Markov Models for the Assessment of Chromosomal Alterations using High-throughput SNP Arrays

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Abstract

This package provides an implementation of a hidden Markov Model to identify copy number alterations from high throughput SNP arrays.

1 From processed SNP summaries

The starting point for this section of the vignette are B allele frequencies and log R ratios that are available from software such as GenomeStudio and the R package crlmm. In this section, we assume that the low-level summaries are available in a plain text file – one file per sample. For users of the crlmm package for preprocessing, please refer to the crlmmDownstream vignette.

To illustrate the workflow for copy number analysis, this package provides Illumina 610k array data for one Hapmap trio and one oral cleft trio preprocessed by GenomeStudio [1]. To keep the size of this package small, the GenomeStudio-processed files for these 6 samples have been parsed to include only columns relevant to copy number analyses and ≈ 11 k markers that are spanned by or border putative CNVs. The code chunks provided below would be the same if the full files were included.

```
> library(VanillaICE)
> library(foreach)
> registerDoSEQ()
> extdir <- system.file("extdata", package="VanillaICE", mustWork=TRUE)
> files <- list.files(extdir, pattern="FinalReport")</pre>
```

The first several lines of these files have the following information:

```
[Header]
BSGT Version,3.3.7
Processing Date,5/2/2009 12:08 AM
Content,,Human610-Quadv1_B.bpm
Num SNPs,620901
Total SNPs,620901
Num Samples,7599
Total Samples,7599
File ,1664 of 7599
[Data]
SNP Name,Allele1 - AB,Allele2 - AB,B Allele Freq,Log R Ratio
cnvi0005126,-,-,0.0085,-0.7214
cnvi0005127,-,-,0.0081,-0.8589
cnvi0005128,-,-,0.0063,-0.164
```

The physical annotation of the markers in the genome can be downloaded from on-line resources or from annotation packages available from Bioconductor. Here, we provide a plain-text file containing the genomic annotation for the markers based on build UCSC build hg18:

```
> list.files(extdir, pattern="SNP_info")
```

[1] "SNP_info.csv"

The annotation file contains the the name of the SNP or nonpolymorphic marker ('Name'), an indicator for whether the marker is a SNP ('Intensity Only'), the chromosome name ('Chr'), and the genomic physical position ('Position'). The columns for the SNP annotation can be in any order and have different variable labels than the file provided with this package. Below, we copy the header of the provided annotation file:

```
Chr,Position,Name,Intensity Only
11,45380778,cnvi0005126,1
11,45382079,cnvi0005127,1
11,46561032,cnvi0005128,1
```

1.1 Organizing marker annotation

We require that marker-level annotation is represented as a **GRanges**-derived class. To read the plain text annotation file, we use the **fread** provided in the *data.table* package. In addition, we define an indicator for whether the marker is polymophic using the 'Intensity Only' flag. The **SnpGRanges** class created from the **fgr** object in the following code-chunk ensures that this binary indicator is created and can be reliably accessed in downstream processing.

Ideally, one should also include the genome build and information on the chromosome lengths appropriate to the build. Here, we extract the metadata on the chromosomes using BSgenome Bioconductor package for the hg18 build. Finally, we sort the **fgr** object such that the chromosomes are ordered by their **seqlevels** and the markers are ordered by their genomic position along the chromosome.

```
> library(BSgenome.Hsapiens.UCSC.hg18)
> sl <- seqlevels(BSgenome.Hsapiens.UCSC.hg18)
> seqlevels(fgr) <- sl[sl %in% seqlevels(fgr)]
> seqinfo(fgr) <- seqinfo(BSgenome.Hsapiens.UCSC.hg18)[seqlevels(fgr),]
> fgr <- sort(fgr)</pre>
```

1.2 Organizing the marker-level summaries

The abbreviated plain text files included with this package and containing log R ratios and B allele frequencies from 2 trios are listed below.

> files <- list.files(extdir, full.names=TRUE, recursive=TRUE, pattern="FinalReport")</pre>

We will parse these files into a specific format such that all downstream steps for copy number estimation no longer depend on the specific format of the source files. At this point, we can encapsulate the the names of the source files ('sourcePaths'), the location of where we intend to store the parsed data ('parsedPath'), and the genomic marker annnotation created in the preceding section in a single object called an ArrayViews.

```
> ##
> ## A directory to keep parsed files
> ##
> parsedDir <- tempdir()
> views <- ArrayViews(rowRanges=fgr, sourcePaths=files, parsedPath=parsedDir)
> lrrFile(views) <- file.path(parsedDir, basename(fileName(views, "lrr")))
> views@bafFiles <- file.path(parsedDir, basename(fileName(views, "baf")))
> views@gtFiles <- file.path(parsedDir, basename(fileName(views, "gt")))
> colnames(views) <- gsub(".csv", "", colnames(views))
> show(views)
class 'ArrayViews'
No. files : 6
No. markers: 11780
```

Because the format of the source files depends on upstream software and version number within software, our approach to parsing these files is to read one file and to store the appropriate metadata from this file so that subsequent files can be parsed in a similar fashion. We use the **fread** to read in the first file.

```
> ## read the first file
> dat <- fread(files[1], skip="[Data]")</pre>
> head(dat,n=3)
      SNP Name Allele1 - AB Allele2 - AB B Allele Freq Log R Ratio
1: cnvi0005126
                                         _
                                                   0.0085
                                                               -0.7214
2: cnvi0005127
                                         _
                                                   0.0081
                                                               -0.8589
3: cnvi0005128
                                         _
                                                   0.0063
                                                               -0.1640
```

Next, we select which columns we plan to keep. Again, the required data for downstream processing is the name of the SNP identifier, the log R ratios, and B allele frequencies.

```
> ## information to store on the markers
> select_columns <- match(c("SNP Name", "Allele1 - AB", "Allele2 - AB",
+ "Log R Ratio", "B Allele Freq"), names(dat))</pre>
```

We also specify the order in which we will store the marker-level summaries by matching the rownames of the views object with the names of the markers in the source file:

> index_genome <- match(names(fgr), dat[["SNP Name"]])</pre>

Similar to the parameter classes defined in *Rsamtools*, we encapsulate the information for parsing the columns and rows of the source files in a class. In addition, we specify which variable names in the source file refers to log R ratios ('cnvar'), B allele frequences ('bafvar'), and genotypes ('gtvar').

```
> scan_params <- CopyNumScanParams(index_genome=index_genome, select=select_columns,
+ cnvar="Log R Ratio",
+ bafvar="B Allele Freq",
+ gtvar=c("Allele1 - AB", "Allele2 - AB"))
```

The parseSourceFile will parse a single file in the views object (by default, the first file) according to the parameters for reading the data in the scan_params object and write the output to the parsedPath directory. In particular, the parseSourceFile returns NULL.

```
> parseSourceFile(views, scan_params)
```

Apart from confirming their existance, the user should not have a need to directly access the parsed files. Utilities for querying these files are provided through the views object.

> head(list.files(parsedPath(views)), n=3)

```
[1] "FinalReport1664_baf.rds" "FinalReport1664_gt.rds"
```

[3] "FinalReport1664_lrr.rds"

1.3 Accessors for the parsed data

The preference for writing the parsed data to disk rather than keeping the data in RAM is simply that the latter does not scale to projects involving thousands of samples. For the former, slices of the parsed data easily be accessed from the parsedPath directory via methods defined for the ArrayViews class. For example, one can use accessors for the low-level summaries directly: 1rr, baf, and genotypes for log R ratios, B allele frequencies, and genotypes, respectively. The user has the option of either subsetting the views object or subsetting the matrix returned by the accessor to extract the appropriate data slice. In the following examples, we access data on the first 2 markers.

> lrr(views)[1:2,]

```
FinalReport1664.txt FinalReport1675.txt
rs12789205
                          0.271
                                              -0.169
rs2114088
                          0.114
                                              -0.182
           FinalReport1686.txt FinalReport6841.txt
rs12789205
                          0.191
                                              -0.084
                         -0.114
rs2114088
                                               0.253
           FinalReport6872.txt FinalReport6903.txt
rs12789205
                          0.067
                                              -0.101
rs2114088
                         -0.076
                                              -0.043
> ## or
> lrr(views[1:2, ])
           FinalReport1664.txt FinalReport1675.txt
rs12789205
                          0.271
                                              -0.169
rs2114088
                          0.114
                                              -0.182
           FinalReport1686.txt FinalReport6841.txt
rs12789205
                          0.191
                                              -0.084
rs2114088
                         -0.114
                                               0.253
           FinalReport6872.txt FinalReport6903.txt
rs12789205
                          0.067
                                              -0.101
rs2114088
                         -0.076
                                              -0.043
```

```
> ## B allele frequencies
```

```
> baf(views[1:2, ])
```

	FinalReport1664.txt	FinalReport1675.txt
rs12789205	0.999	0.529
rs2114088	0.000	0.000
	FinalReport1686.txt	FinalReport6841.txt
rs12789205	0.549	0.481
rs2114088	0.000	0.000
	FinalReport6872.txt	FinalReport6903.txt
rs12789205	0.482	0.488
rs2114088	0.000	0.493
	CE::genotypes(views)	by function of the same name in crlmm [1:2,]
	CE::genotypes(views)	-
	CE::genotypes(views)	[1:2,]
> VanillaI	CE::genotypes(views) FinalReport1664.txt	[1:2,] FinalReport1675.txt
> VanillaI rs12789205	CE::genotypes(views) FinalReport1664.txt 3 1	[1:2,] FinalReport1675.txt 2
> VanillaI rs12789205	CE::genotypes(views) FinalReport1664.txt 3 1	[1:2,] FinalReport1675.txt 2 1
> VanillaI rs12789205 rs2114088	CE::genotypes(views) FinalReport1664.txt 3 FinalReport1686.txt 2 1	[1:2,] FinalReport1675.txt 2 1 FinalReport6841.txt 2 1
<pre>> VanillaI rs12789205 rs2114088 rs12789205 rs2114088</pre>	CE::genotypes(views) FinalReport1664.txt 3 FinalReport1686.txt 2 1 FinalReport6872.txt	[1:2,] FinalReport1675.txt 2 1 FinalReport6841.txt 2 1 FinalReport6903.txt
<pre>> VanillaI rs12789205 rs2114088 rs12789205</pre>	CE::genotypes(views) FinalReport1664.txt 3 FinalReport1686.txt 2 1	[1:2,] FinalReport1675.txt 2 1 FinalReport6841.txt 2 1

More often, it is useful to extract the low-level data in a RangedSummarizedExperiment-derived object such that meta-data on the samples remains bound to the columns of the assay data (log R ratios / B allele frequencies) and meta-data on the rows remains bound to the rows of the assay data. This is accomplished by applying the SnpExperiment function to a views object. In the following example, we create a RangedSummarizedExperiment-derived object for the first three samples in the views object.

```
> snp_exp <- SnpExperiment(views[, 4:5])
> show(snp_exp)

class: SnpArrayExperiment
dim: 11780 2
metadata(0):
assays(2): cn baf
rownames(11780): rs12789205 rs2114088 ... rs5994329 rs1055232
rowData names(1): isSnp
colnames(2): FinalReport6841.txt FinalReport6872.txt
colData names(0):
```

2 Hidden Markov model

2.1 HMM parameters

The hidden Markov model for inference of germline copy number from SNP array data was originally described in Scharpf *et al.* [3] but has evolved to focus more on B allele frequencies than genotype call probabilities [the latter requires a dependency on the *crlmm* package]. Many of the the updates for computing emission probabilities have been described more recently [2].

Fitting the hidden Markov model to the marker-level summaries requires extracting the relevant data from a views object using the SnpExperiment function illustrated in the previous section. All

user-level parameters relevant for fitting the HMM are specified in a parameter object for the emission probabilities and a parameter object for the transition probabilities. In the following code-chunk, we create a parameter object for the emission probabilities.

```
> param <- EmissionParam()</pre>
```

The easiest way to specify or change parameters for computing emission probabilities is through the EmissionParam constructor. For example, here we create a parameter object with the temper parameter set to 1/2:

```
> param <- EmissionParam(temper=0.5)
> show(param)
EmissionParam :
CN mean: -2, -0.4, 0, 0, 0.4, 1
CN sd: 0.6, 0.3, 0.3, 0.3, 0.3, 0.3
BAF mean: 0, 0.25, 0.33, 0.5, 0.67, 0.75, 1
BAF sd: 0.1, 0.1, 0.1, 0.1, 0.1, 0.1
max # of EM updates: 5
initial state probabilities: 0.17, 0.17, 0.17, 0.17, 0.17
tempering scalar: 0.5
model homozygous regions: FALSE
See temper(), cn_means(), cn_sds(),...
```

Similarly, the constructor for the TransitionParam class can be used to specify alternatives to the defaults. By not specifying an TransitionParam object, default values created by the constructor are used by the HMM in the following example.

2.2 Fitting the HMM

The function hmm2 fits the 6-state HMM to the snp_exp object, returning a list-derived class (a HMMList) of the same length as the number of samples.

```
> fit <- hmm2(snp_exp, param)</pre>
> show(fit)
An object of class 'HMMList'
 Length: 2
 Use [[j]] to retrieve results for the jth sample.
 Use unlist to coerce to a 'GRanges' object.
> length(fit)
[1] 2
> ## HMM object for the first sample
> fit[[1]]
Object of class 'HMM'
  granges (no. segments) : 7
 no. duplications
                          : 2
 no. hemizygous deletions: 0
 no. homozygous deletions: 0
See posterior(), segs(), cnvSegs(), getHmmParams(),
```

3 Inspecting, Filtering, and plotting HMM results

Several methods are defined for the HMMList class to facilitate the selection of candidate CNVs, the filtering of technical artifacts, and to enable visualization of the HMM calls in the context of the marker-level summaries.

The fit object is an instance of the HMMList object of length three, each element corresponding to a sample:

```
> names(fit)
```

[1] "FinalReport6841.txt" "FinalReport6872.txt"

The R method $\$ can be used to access the HMM summary for the results for the jth sample. For example, here we extract the HMM summary for the 2nd sample:

```
> show(fit[[2]])
```

```
Object of class 'HMM'
granges (no. segments) : 13
no. duplications : 1
no. hemizygous deletions: 4
no. homozygous deletions: 0
See posterior(), segs(), cnvSegs(), getHmmParams(),
```

Alternatively, we can use unlist to create a single GRanges-derived class containing the segmentation of all three samples

```
> head(unlist(fit), n=3)
```

HmmGRanges object with 3 ranges and 4 metadata columns: seqnames ranges strand | numberFeatures state <Rle> <IRanges> <Rle> | <integer> <integer> [1] chr11 43132366-55116789 1445 3 * | 5 [2] chr11 55124465-55204003 * | 29 [3] chr11 55209499-67164817 * | 1963 3 prCall id <numeric> <character> [1] 1 FinalReport6841.txt [2] 1 FinalReport6841.txt [3] 1 FinalReport6841.txt

```
seqinfo: 3 sequences from hg18 genome
```

or a GRangesList by splitting on the sample id:

```
> grl <- split(unlist(fit), unlist(fit)$id)</pre>
```

3.1 Filters

There are several meta-data columns stored in the **GRanges**-derived summary useful for filtering the set of genomic intervals. All the available parameters for filtering the **fit** object are stored in the parameter class **FilterParam** of which an instance can be created by its constructor of the same name without any arguments.

```
> filter_param <- FilterParam()
> show(filter_param)
An object of class 'FilterParam'
   min. posterior probability of CNV call: 0.99
   min. no. of markers spanned by segment: 10
   min. width of segment : 1
   selected HMM states : 1, 2, 3, 4, 5, 6
   selected seqnames : chr1, chr2, chr3, chr4, chr5, chr6 ...
```

To apply the default filter parameters to the fit object, we use the cnvFilter function. The cnvFilter function returns only the set of genomic ranges satisfying the filters. For example,

```
> cnvFilter(fit, filter_param)
```

HmmGRa	nges obje	ct with	18	ranges	and 4 m	net	adata d	columns:	
	seqnames			ranges	s strand	1	number	Features	state
	<rle></rle>		<]	[Ranges>	<rle></rle>	>	<	<pre>integer></pre>	<integer></integer>
[1]	chr11	4313236	6-5	55116789) *	<		1445	3
[2]	chr11	5512446	65-5	5204003	} *	<		29	5
[3]	chr11	5520949	9-6	67164817	′ *	<		1963	3
[4]	chr12	2105	54-1	17482570) *	<		4591	3
[5]	chr22	1445641	2-2	23994408	} *	<		2244	3
						•			
[14]	chr22	1871089	95-1	19048116	; *	<		28	3
[15]	chr22	1906631	.5-1	19819918	} *	<		211	2
[16]	chr22	1984233	33-2	23983992	2 *	<		1099	3
[17]	chr22	2399172	25-2	24236803	} *	<		63	5
[18]	chr22	2423790)5-2	29381906	; *	<		1441	3
	prCall	1			id				
	<numeric2< td=""><td>></td><td><</td><td><charact< td=""><td>er></td><td></td><td></td><td></td><td></td></charact<></td></numeric2<>	>	<	<charact< td=""><td>er></td><td></td><td></td><td></td><td></td></charact<>	er>				
[1]		1 FinalF	lepo	ort6841.	txt				
[2]		1 FinalF	lepo	ort6841.	txt				
[3]		1 FinalF	lepo	ort6841.	txt				
[4]	:	1 FinalF	lepo	ort6841.	txt				
[5]		1 FinalF	lepo	ort6841.	txt				
[14]	:	1 FinalF	lepo	ort6872.	txt				
[15]	:	1 FinalF	lepo	ort6872.	txt				
[16]	:	1 FinalF	lepo	ort6872.	txt				
[17]	:	1 FinalF	lepo	ort6872.	txt				
[18]	:	1 FinalF	lepo	ort6872.	txt				

seqinfo: 3 sequences from hg18 genome

To select only the segments with altered copy number states (states 1, 2, 5, and 6) on chromosome 22, we can define the filter parameters as follows:

```
> select_cnv <- FilterParam(state=c("1", "2", "5", "6"), seqnames="chr22")
> cnvs <- cnvFilter(fit, select_cnv)
> cnvs
```

HmmGRa	nges obje	ct with 6	ranges	and 4 me	etadata	columns:	
	seqnames		ranges	strand	numbe	erFeatures	state
	<rle></rle>	<1	Ranges>	<rle></rle>	1	<integer></integer>	<integer></integer>
[1]	chr22	23999142-2	24240667	*	1	65	5
[2]	chr22	17202486-1	7388108	*		44	2
[3]	chr22	17405381-1	7633332	*		48	2
[4]	chr22	17690812-1	.8693299	*	1	281	2
[5]	chr22	19066315-1	9819918	*	1	211	2
[6]	chr22	23991725-2	24236803	*	1	63	5
	prCall			id			
	<numeric></numeric>	<	charact	er>			
[1]	1	FinalRepo	rt6841.	txt			
[2]	1	FinalRepo	rt6872.	txt			
[3]	1	FinalRepo	rt6872.	txt			
[4]	1	FinalRepo	rt6872.	txt			
[5]	1	FinalRepo	rt6872.	txt			
[6]	1	FinalRepo	rt6872.	txt			
	mfat 2 da	averea fr	am haio	~~~~~			

seqinfo: 3 sequences from hg18 genome

3.2 Visualization

3.2.1 Trellis graphics for low-level summaries

Visualization of the CNVs in the context of the lower-level summaries is acheived through a combination of grid and the grid-derived graphics provided by *lattice*. The xyplotList and constructor HmmTrel-lisParam should be sufficient for producing a decent graphic using the default settings. In the following code chunk, we create a trellis object for each CNV segment.

```
> trellis_param <- HmmTrellisParam()
> cnvList <- split(cnvs, cnvs$id)
> figList <- xyplotList(cnvList, snp_exp, trellis_param)
> names(figList)
```

[1] "FinalReport6841.txt" "FinalReport6872.txt"

Each element in figList is a trellis object and can be displayed using print.

```
> class(figList[["FinalReport6841.csv"]][[1]])
```

```
[1] "NULL"
```

3.2.2 Layout using grid

```
> fig1 <- figList[["FinalReport6841.csv"]][[1]]</pre>
```

```
> vps <- viewports()
> xygrid(fig1, vps, cnvList[[1]][1])
```

NULL

A hemizygous deletion on chromosome 22 is split into too many regions.

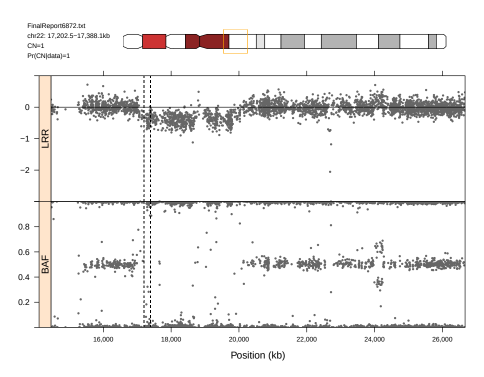
```
> cnvs_sample2 <- cnvList[[2]]
> cnvs_sample2
```

FinalReport6841.txt				
chr22: 23,999.1-24,240.7kb				٦
CN=3				J
Pr(CN data)=1	L_			

Figure 1: A single-copy duplication (state 5).

HmmGRar	nges object w	ith 5 ranges a	and 4 me	etadata	columns:			
2	seqnames	ranges	strand	numb	erFeatures	state		
	<rle></rle>	<iranges></iranges>	<rle></rle>	I	<integer></integer>	<integer></integer>		
[1]	chr22 1720	2486-17388108	*		44	2		
[2]	chr22 1740	5381-17633332	*		48	2		
[3]	chr22 1769	0812-18693299	*	1	281	2		
[4]	chr22 1906	6315-19819918	*	1	211	2		
[5]	chr22 2399	1725-24236803	*	1	63	5		
	prCall		id					
<	<numeric></numeric>	<characte< td=""><td>er></td><td></td><td></td><td></td></characte<>	er>					
[1]	1 Fina	alReport6872.	txt					
[2]	1 Fina	alReport6872.	txt					
[3]	1 Fina	alReport6872.	txt					
[4]	1 Fina	alReport6872.	txt					
[5]	1 Fina	alReport6872.	txt					
seqir	seqinfo: 3 sequences from hg18 genome							

> xygrid(figList[[2]][[1]], vps, cnvs_sample2[1])



Combining the adjacent hemizygous deletions through **reduce** provides a more satisfying segmentation of this data.

```
> cnvs_sample2r <- reduce(cnvs_sample2, min.gapwidth=500e3)
> fig2 <- xyplotList(cnvs_sample2r, snp_exp)</pre>
```

Parallelization

As the HMM is fit independently to each sample, parallelization is straightforward. In the following unevaluated code chunk, we set up a parallel environment using the R packages *snow* and *foreach*.

```
> library(foreach)
> library(snow)
> library(doSNOW)
> cl <- makeCluster(2, type="SOCK")
> registerDoSNOW(cl)
> results <- hmm2(snp_exp)
> stopCluster(cl)
```

Citing this software

Robert B Scharpf, Giovanni Parmigiani, Jonathan Pevsner, and Ingo Ruczinski. Hidden Markov models for the assessment of chromosomal alterations using high-throughput SNP arrays. *Annals of Applied Statistics*, 2(2):687–713, 2008.

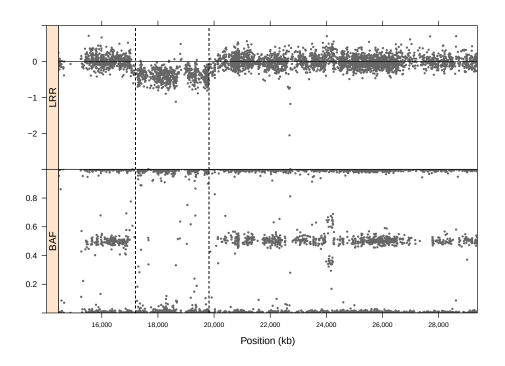


Figure 2: A hemizygous deletion (state 2) on chromosome 22 after merging adjacent hemizygous CNV calls.

Session Information

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

- R version 3.5.0 (2018-04-23), x86_64-pc-linux-gnu
- Locale: LC_CTYPE=en_US.UTF-8, LC_NUMERIC=C, LC_TIME=en_US.UTF-8, LC_COLLATE=C, LC_MONETARY=en_US.UTF-8, LC_MESSAGES=en_US.UTF-8, LC_PAPER=en_US.UTF-8, LC_NAME=C, LC_ADDRESS=C, LC_TELEPHONE=C, LC_MEASUREMENT=en_US.UTF-8, LC_IDENTIFICATION=C
- Running under: Ubuntu 16.04.4 LTS
- Matrix products: default
- BLAS: /home/biocbuild/bbs-3.7-bioc/R/lib/libRblas.so
- LAPACK: /home/biocbuild/bbs-3.7-bioc/R/lib/libRlapack.so
- Base packages: base, datasets, grDevices, graphics, methods, parallel, stats, stats4, utils
- Other packages: BSgenome 1.48.0, BSgenome.Hsapiens.UCSC.hg18 1.3.1000, Biobase 2.40.0, BiocGenerics 0.26.0, BiocParallel 1.14.1, Biostrings 2.48.0, DelayedArray 0.6.0, GenomeInfoDb 1.16.0, GenomicRanges 1.32.2, IRanges 2.14.8, S4Vectors 0.18.1, SummarizedExperiment 1.10.1, VanillaICE 1.42.4, XVector 0.20.0, data.table 1.11.2, foreach 1.4.4, matrixStats 0.53.1, rtracklayer 1.40.2

Loaded via a namespace (and not attached): BiocInstaller 1.30.0, DBI 1.0.0, GenomeInfoDbData 1.1.0, GenomicAlignments 1.16.0, Matrix 1.2-14, RCurl 1.95-4.10, Rcpp 0.12.16, RcppEigen 0.3.3.4.0, Rsamtools 1.32.0, VGAM 1.0-5, XML 3.98-1.11, affyio 1.50.0, base64 2.0, beanplot 1.2, bit 1.1-12, bitops 1.0-6, codetools 0.2-15, compiler 3.5.0, crlmm 1.38.0, ellipse 0.4.1, ff 2.2-13, grid 3.5.0, illuminaio 0.22.0, iterators 1.0.9, lattice 0.20-35, limma 3.36.1, mvtnorm 1.0-7, oligoClasses 1.42.0, openssl 1.0.1, preprocessCore 1.42.0, splines 3.5.0, tools 3.5.0, zlibbioc 1.26.0

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