RIPSeeker: a statistical package for identifying protein-associated transcripts from RIP-seq experiments

Yue Li
yueli@cs.toronto.edu
May 3, 2016

1 Introduction

Ribonucleoprotein (RNP) immunoprecipitation (IP) followed by high-throughput sequencing (RIP-seq) has recently been developed to discover genome-wide RNA transcripts that interact with a protein or protein complex. RIP-seq is similar to both RNA-seq and ChIP-seq, but presents unique properties and challenges. Currently, no statistical tool is dedicated to RIP-seq analysis. We developed RIPSeeker, an HMM-based R package for de novo RIP peak predictions. RIPSeeker infers and discriminate RIP peaks from background or control (if available) using two-state HMM with negative binomial emission probability followed by statistical tests for model confidence. To evaluate the performance of RIPSeeker, we used the published RIP-seq data for PRC2 and also generated in-house data for CCNT1. Under an equal footing, RIPSeeker compares competitively with ChIP-seq and transcript-based methods in predicting canonical proteinassociated transcripts with high statistical confidence. Importantly, RIPSeeker has the ability to model reads on plus and minus strand separately, which allows identification of strand-specific noncoding RNA (e.g., antisense transcripts). Comparing to the published methods, RIPSeeker is adaptive to a larger dynamical range of peaks suitable for detecting the entire transcripts with various lengths rather than the punctuated binding sites

While *RIPSeeker* is specifically tailored for RIP-seq data analysis, it also provides a suite of bioinformatics tools integrated within this self-contained software package comprehensively addressing issues ranging from post-alignments processing to visualization and annotation. In addition, a rule-based approach is provided as an additional function named rulebaseRIPSeek for user to obtain RPKM/FPKM (and fold-change) for the gene/transcripts expressions in RIP (and control) based on automatically retrieved online Ensembl/UCSC annotation given single or paired-end alignments. This vignette provides a guide to the most common application of *RIPSeeker* package.

2 RIPSeeker overview

Figure 3 in the manuscript (in press) for peer-review presents the workflow of *RIPSeeker*. The input for RIPSeeker is a list of read alignments in BAM/BED/SAM format. Map-

ping reads to the reference genome can be performed by any RNA-seq aligner such as TopHat Trapnell et al. [2009]. After post-processing the alignment input by combineAlignGals, RIPSeeker first stratifies the genome into bins of automatically selected (selectBinSize) or a fixed user-defined size. Each bin may contain more than one aligned read. Multiple bins may together correspond to a single RNA transcript that binds to the protein of interest. Thus, these bins when treated as individual observations are not independent identically distributed (*i.i.d.*) and need to be treated as dependent events. Hidden Markov model (HMM) provides a sensible and efficient way to probabilistically model the dependence between sequential events through hidden variables [Rabiner, 1989, Bishop, 2006]. The adaptation of HMM is inspired by HPeak, which was specifically designed for ChIP-seq [Qin et al., 2010].

As an overview, RIPSeeker consists of two major steps: probabilistic inference of RIP regions (mainSeek) and significance test for the inferred RIP regions from HMM (seekRIP). In the first step, we apply a two-state HMM to model the background and RIP distributions (or emission probabilities) of RIP-seq read counts as negative binomial distributions, which has been shown by Anders and Huber [2010] to be a more realistic parametric model than Gaussian and Poisson models. The parameters of HMM are learned from the data by expectation-maximization (EM). The intermediate quantities required in the EM iterations are efficiently computed using forward-backward algorithm. After the optimization, Viterbi algorithm is applied to derive the most probable sequence of hidden states, which encodes whether each region is background (1) or RIP (2) across the genome. The consecutive RIP bins are merged into a single RIP region. In the second step, we compute the statistical significance of each RIP region with or without a control library based on the posterior probabilities derived directly from the HMM.

RIPSeeker is able to detect strand-specific RIP regions by running the same workflow on either plus and minus strand separately, making use of the strand-specific information retained in the original RIP-seq protocol [Cloonan et al., 2008, Zhao et al., 2010]. In addition, RIPSeeker takes advantage of modern computational architecture equipped with multiple processors by treating each chromosome as an independent thread and computing multiple threads in parallel using mclapply from multicore R package. Thus, the most time consuming step such as HMM inference operates on per-chromosome basis each running on a separate CPU core. The parallel computing is much more computationally and memory efficient than computing the entire genome all at once by treating it as a single concatenated sequence. RIPSeeker has numerous other features including disambiguating multihits (i.e., reads mapped to multiple loci with disambiguateMultihits), automatic annotation of RIP regions (annotateRIP), GO enrichment analysis (annotateRIP), and UCSC visualization (viewRIP).

Although the ultimate goal is to predict RIP regions, each step or component in the workflow has been implemented as a stand-alone function. Please referred to MATE-RIALS AND METHODS in the manuscript (may not be available during peer-review) for the underlying algorithms and the help manual for their usage:

```
> library(RIPSeeker)
```

> ?RIPSeeker

3 RIP-seq data

3.1 PRC2 in mouse embrynoic stem cell

Only a few RIP-seq datasets are available. In this tutorial, we will make use of part of the dataset from Zhao et al. [2010] (GEO accession: GSE17064). The data was generated to identify transcripts phycially associated with Ezh2 (a PRC2 unique subunit) in mouse embryonic stem cell (mESC). A negative control was also generated from mutant mESC depleted of Ezh2 (Ezh2 -/-) (MT). Here we will use the RIP data consisting of two techincal replicates (code ID: SRR039210-1, SRR039212) and the MT control (code ID: SRR039214). Notably, the library construction and strand-specific sequencing generated sequences from the opposite strand of the PRC2-bound RNA [Zhao et al., 2010], consequently, each read will be treated as if it were reverse complemented. After the quality control (QC) and alignments (Quality Control of Raw Read Library and Alignment of Filtered RIP-seq Read Library to Reference Genome in Supplementary Datafor the manuscript), the technical replicates were merged, resulting in 1,022,474 and 208,445 reads mapped to unique loci of the mouse reference genome (mm9 build) for RIP and control, respectively. To make the demonstration and the package size more managable, only the alignments to chromosome X (chrX) were extracted to generate the test data stored in the package. The full data (containing all of the processed alignments for the test data and another biological replicate SRR039213) are available as a Bioconductor data package RIPSeekerData. User can try the same command below on the full dataset.

```
> biocLite("RIPSeekerData")
> library(RIPSeekerData)
> extdata.dir <- system.file("extdata", package="RIPSeekerData")
> bamFiles <- list.files(extdata.dir, "\\.bam$",
+ recursive=TRUE, full.names=TRUE)
> bamFiles <- grep("PRC2/", bamFiles, value=TRUE)</pre>
```

3.2 CCNT1 in HEK293

The data for CCNT1 and GFP control were generated in-house in two experiments. The pilot experimental data contain 5,853 and 4,556 uniquely mapped read after the stringent QC for the CCNT1 and GFP control, respectively. Same as in the PRC2 data, the reads came from the second strand of the cDNA synthesis opposite to the original RNA strand. The non-strand-specific library from the second screen has deeper coverage with 26,859 and 45,024 uniquely aligned reads under QC for CCNT1 and GFP, respectively. CCNT1 is known to only associate with *RN7SK*, which can be used as an empirical measurement of senstivitiy and specificity of *RIPSeeker*. The data are in *RIPSeekerData* package.

3.3 User-supplied data

For the following example, user may replace the extdata.dir with the directory containing their own alignment data and change the cNAME to point to the specific control (if applicable).

```
> # Retrieve system files
> # OR change it to the extdata.dir from the code chunk above
```

> # to get RIP predictions on the full alignment data

311119111110111		WICH SIOSI		CIICD	arra 1	mecadae		o ± anni •		
		seqnames	strand		cigar	qwi	dth	start	end	
		<rle></rle>	<rle></rle>	<cha< td=""><td>racter></td><td><integ< td=""><td>er></td><td><integer></integer></td><td><integer></integer></td><td></td></integ<></td></cha<>	racter>	<integ< td=""><td>er></td><td><integer></integer></td><td><integer></integer></td><td></td></integ<>	er>	<integer></integer>	<integer></integer>	
SRR0392	10.4322179	chrX	+		361	1	36	3000326	3000361	
SRR0392	10.5524106	chrX	_		361	1	36	3001293	3001328	
SRR0392	10.5069294	chrX	_		361	1	36	3002790	3002825	
SRR0392	10.1476279	chrX	_		36N	1	36	3016328	3016363	
SRR0392	210.711491	chrX	+		361	l	36	3021161	3021196	
SRR0392	11.1427139	chrX	_		361	l	36	166443604	166443639	
SRR0392	211.447965	chrX	_		361	l	36	166445241	166445276	
SRR0392	11.1352670	chrX	_		361	l	36	166446255	166446290	
SRR0392	211.918096	chrX	_		361	l	36	166446315	166446350	
SRR03	9211.67451	chrX	_		361	l	36	166446621	166446656	
		width	n n	junc	uniqu	ıeHit				
		<integer></integer>	<integ< td=""><td>ger></td><td> <logi< td=""><td>cal></td><td></td><td></td><td></td><td></td></logi<></td></integ<>	ger>	<logi< td=""><td>cal></td><td></td><td></td><td></td><td></td></logi<>	cal>				
SRR0392	10.4322179	36)	0	E	ALSE				
SRR0392	10.5524106	36)	0	E	ALSE				
SRR0392	10.5069294	36)	0	E	ALSE				
SRR0392	10.1476279	36)	0		TRUE				
SRR0392	210.711491	. 36)	0	E	ALSE				
SRR0392	11.1427139	36		0	E	ALSE				

SRR039211.1352670 36 0 | SRR039211.918096 36 0 | SRR039211.67451 36 0 |

36

seqinfo: 22 sequences from mm9 genome

> ctlGal

SRR039211.447965

GAlignments object with 8276 alignments and 1 metadata column:

	seqnames	strand	cigar	qwidth	start	end
	<rle></rle>	<rle></rle>	<character></character>	<integer></integer>	<integer></integer>	<integer></integer>
SRR039214.4949641	chrX	+	36M	36	3028539	3028574
SRR039214.5310910	chrX	_	35M	35	3039791	3039825
SRR039214.4625677	chrX	_	36M	36	3084567	3084602

0 |

FALSE

FALSE

FALSE

TRUE

SRR039214.1854227	chrX	+	36M	36	3139109	3139144
SRR039214.5753635	chrX	+	36M	36	3139131	3139166
• • •						• • •
SRR039214.904524	chrX	_	32M	32	166445788	166445819
SRR039214.2473565	chrX	_	21M	21	166445960	166445980
SRR039214.576581	chrX	_	36M	36	166446074	166446109
SRR039214.3756853	chrX	_	36M	36	166446781	166446816
SRR039214.2347055	chrX	+	36M	36	166650104	166650139
	width	njunc	uniqueHit			
	<integer></integer>	<integer></integer>	<logical></logical>			
SRR039214.4949641	36	0	FALSE			
SRR039214.5310910	35	0	FALSE			
SRR039214.4625677	36	0	FALSE			
SRR039214.1854227	36	0	FALSE			
SRR039214.5753635	36	0	FALSE			
SRR039214.904524	32	0	FALSE			
SRR039214.2473565	21	0	FALSE			
SRR039214.576581	36	0	FALSE			
SRR039214.3756853	36	0	FALSE			
SRR039214.2347055	36	0	TRUE			

6

9 0 9

seqinfo: 22 sequences from mm9 genome

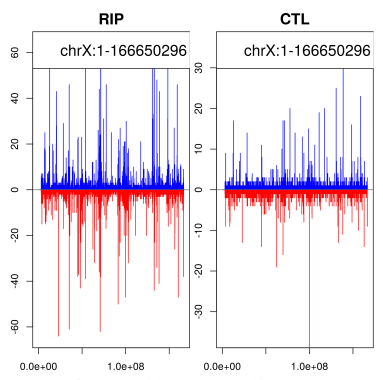
Note from the commands above that the RIP alignment files corresponding to the technical replicates are combined into a single *GAlignments* object[Aboyoun et al.]. The uniqueHit column from the output above indicate whether the read mapped to a single locus or multiple loci. The latter is commonly referred as the "multihits". The ambiguous alignments arise from the repetitive elements or gene duplication events in the mammalian genomes. By default, the unique and multihits are flagged with binary TRUE and FALSE value in column uniqueHit, respectively. In the later step, each multihit will be assigned by the function disambiguateMultihits to a unique locus based on the posterior probabilities of the loci of being at the background or read enriched states of the two-state HMM (Disambiguate multihits using posterior decoding from HMM in the manuscript).

4 RIP-seq analysis

4.1 High-level visuzliation of alignments at chromosomal level

After processing the alignment files, we could first visualize the alignments by calling plotStrandedCoverage to get a rough idea about the alignment quality and where the reads aggregate within each chromosome. plotStrandedCoverage first bins each chromosome by nonoverlapping windows of fixed size (1 kb in the example below) and plots the number of alignments fall within these bins across the whole chromosome. The function is implemented based on the vignette *GenomicRanges Use Cases* from the *GenomicRanges* package [Aboyoun et al.].

```
> ripGR <- as(ripGal, "GRanges")
> ripGRList <- split(ripGR, seqnames(ripGR))</pre>
```



Here we first converted the *GAlignments* object ripGal and ctlGal to *GenomicRangesList* to have each list item represented by a chromosome. We then lapply the function to all nonempty chromosomes, which in our case is *chrX* for both RIP (left) and control (right). RIP represents the pooled alignments of the two technical replicates. Read count on + and - strand are displayed as blue and red bars on the positive and negative y-axis, respectively. Read counts are drawn to scale within a chromosome. Unlike ChIP-seq on double-stranded DNA, no symmetry is observed for the peak for the strand-specific sequencing data. However, considerable noise is observed within the Ezh2 -/- mutant library (right panel), which ideally should not have any aligned reads. This may also imply considerable noise within the wild type library and thus a high false discovery rate if the RIP regions were simply determined based on read counts. Please refer to Figure S3 and S4 in Supplementary Datafor similar RIP-seq

results on all chromosomes, which motivated the development of RIPSeeker.

4.2 ripSeek: the all-in-one function

The front-end main function ripSeek is in many cases the only function that users need to run to get the RIP predictions and most of the relevant information out of the alignment data. It requires a few important parameter settings:

```
> # specify control name
> cNAME <- "SRR039214"
> # output file directory
> outDir <- file.path(getwd(), "RIPSeeker_vigenette_example_PRC2")</pre>
> # Parameters setting
> binSize <- NULL
                       # set to NULL to automatically determine bin size
> minBinSize <- 10000 # min bin size in automatic bin size selection
> maxBinSize <- 10100</pre>
                               # max bin size in automatic bin size selection
> multicore <- TRUE
                                     # use multicore
> strandType <- "-"
                                     # set strand type to minus strand
> biomart <- "ENSEMBL_MART_ENSEMBL"</pre>
                                               # use archive to get ensembl 65
> biomaRt_dataset <- "mmusculus_gene_ensembl" # mouse dataset id name
                                               # use ensembl 65 for annotation 1
> host <- "dec2011.archive.ensembl.org"</pre>
> goAnno <- "org.Mm.eg.db"</pre>
                                                # GO annotation database
> ############## run main function ripSeek to predict RIP ###############
> seekOut.PRC2 <- ripSeek(bamPath = bamFiles, cNAME = cNAME,
                     reverseComplement = TRUE, genomeBuild = "mm9",
                     strandType = strandType,
                     uniqueHit = TRUE, assignMultihits = TRUE,
                     rerunWithDisambiguatedMultihits = TRUE,
                     binSize=binSize, minBinSize = minBinSize,
                     maxBinSize = maxBinSize,
                     biomart=biomart, host=host,
                     biomaRt_dataset = biomaRt_dataset, goAnno = goAnno,
                     multicore=multicore, outDir=outDir)
```

- bamPath saves the paths to the bam files;
- cNAME is a character string that distinguishes the control from the RIP among the bamFiles;
- reverseComplement=TRUE indicates that the reads were sequenced from the opposite strand of the original RNA molecule such that the original strand signs of the alignments are switched (i.e. + to -, - to +);
- genomeBuild="mm9" specifies the fact that the previous mouse genome (mm9/NCBIM37) build was used as reference for the alignments;
- strandType is set to "-" to search for transcripts only on the minus strand of chromosomes such as Xist, which is known to physically interact with PRC2 Zhao et al. [2010].
- uniqueHit = TRUE requires training HMM with only the unique hits.

- assignMultihits = TRUE enables disambiguateMultihits to assign each multihit to a unique locus based on the posterior probabilities derived from HMM.
- rerunWithDisambiguatedMultihits = TRUE tells RIPSeeker to retrain the HMM using the dataset with disambiguated multihits.
- binSize = NULL enables automatic bin size selection by the routine selectBinSize.
- minBinSize=10000, maxBinSize=12000 For demonstration puprose, we set the lower and upper bounds of the search space for the optimal bin size to 10 kb and 12 kb, respectively. Such choice may suffice if one just wants to quickly look for some known long noncoding RNA (lncRNA) such as Xist (22,843 nt) or Tsix (53,441 nt).
- biomart, host, biomart_dataset, goAnno are set to enable automatic online annotation of RIP predictions via annotateRIP, which makes use of the functions from *ChIPpeakAnno* [Zhu et al., 2010] and *biomaRt* [Durinck et al., 2005, 2009]. Since the genome we used for the alignments is mm9 not the most recent build mm10, we need to retreive the archive annotation from Ensembl by setting the host and biomart. If the user uses the most recent assembly as reference genome for the alignments, then biomart should be set to "ensembl" and host may be omitted.
- multicore = TRUE enables parallel computing on each chromosome separately. It will greatly improve the computation time when used on a cluster.
- outDir the output directory to save the results (See Value section in ?ripSeek)

4.3 ripSeek outputs

The main function ripSeek returns a *list* which comprises four items outlined below. The user may find annotatedRIPGR the most useful. In most cases, the other three items can be considered as intermediate results.

```
> names(seekOut.PRC2)
> df <- elementMetadata(seekOut.PRC2$annotatedRIPGR$sigGRangesAnnotated)
> # order by eFDR
> df <- df[order(df$eFDR), ]
> # get top 100 predictions
> df.top100 <- head(df, 100)
> head(df.top100)
> # examine known PRC2-associated lncRNA transcripts
> df.top100[grep("Xist", df$external_gene_id), ]
```

- mainSeekOutputRIPA (inner) list containing three items:
 - nbhGRList A GRangesList object of the HMM trained parameters for each chromosome on RIP;
 - alignGal, alignGalFiltered GAlignments objects of the RIP alignment outputs from combineAlignGals and disambiguateMultihits, respectively. The former may contain multiple alignments due the same reads whereas the latter contains a one-to-one mapping from read to alignment after disambiguating the multihits;

- mainSeekOutputCTL Same as mainSeekOutputRIP but for the control library;
- RIPGRList The peaks in *GRangesList*. Each list item represents the RIP peaks on a chromosome accompanied with statistical scores including (read) count, logOddScore, pval, pvalAdj, eFDR for the RIP and control (if available).
- annotatedRIPGR A *list* containing two items:
 - sigGRangesAnnotatedGRanges containing peaks and their scores (same as in RIPGRList) accompanied with genomic information retrieved from Ensembl database;
 - enrichedGOA *data.frame* containing enriched GO terms associated with the RIP peaks.

In addition, the (annotated) peaks and enriched GO annotations are saved in outDir as text files in tab-delimted (viewable with Excel) or GFF3 (General Feature Format 3) formats (?rtracklayer:io for details for the formats.).

```
> list.files(outDir)
```

File (1) and (4) can be imported to a dedicated genome browser such as Integrative Genomic Viewer (IGV) Robinson et al. [2011] to visualize and interact with the putative RIP regions accompanied with all of the scores. Files (2) and (3) provide the most detailed information directly viewable in Excel. File (5) stores all of the intermediate results as RData for future reference or retrieval (with load command).

4.4 Visualization with UCSC browser

In addition, we could visualize the results using <code>viewRIP</code>, which launches online UCSC genome browser with uploaded custom track corresponding to the loci of RIP regions and scores (RIPScore, -log10(p-value), -log10(q-value), or -log10(eFDR)) generated from <code>ripSeek</code>. This task is accomplished seamlessly within the R console by making use of the available functions from <code>rtracklayer</code> Lawrence et al.

```
> viewRIP(seekOut.PRC2$RIPGRList$chrX,
+ seekOut.PRC2$mainSeekOutputRIP$alignGalFiltered,
+ seekOut.PRC2$mainSeekOutputCTL$alignGalFiltered, scoreType="eFDR")
```

4.5 Compute RPKM and fold-change of RIP over control for known genes or transcripts

User may want to know the abundance of all of the known coding or noncoding genes or transcripts in their RIP libraries and how their expressions compare with the control. Given a list of single-end or paired-end read alignment files in BAM/SAM/BED format, the function computeRPKM computes the read counts and normalized read counts as expression of annotated transcripts in the unit of "reads per kilobase of exon per million mapped reads" (RPKM) or "fragments per kilobase of exon per million mapped reads" (FPKM), respectively. Furthermore, rulebaseRIPSeek computes

¹The "fragments" refers to the long fragment sequneced on both ends by the short read pairs.

the fold-change difference of gene expression in terms of RPKM/FPKM between the RIP and control libraries. rulebaseRIPSeek reports putative RIP genes/transcripts if their RPKM expression and the ratio of RPKM[RIP]/RPKM[control] (on either + or - strand) are above t_1 and t_2 . By default, $t_1=0.4$ and $t_2=3.0$, consistent with the thresholds applied in original method Zhao et al. [2010].

```
> # Retrieve system files
> extdata.dir <- system.file("extdata", package="RIPSeeker")
> bamFiles <- list.files(extdata.dir, ".bam$", recursive=TRUE, full.names=TRUE)
> bamFiles <- grep("PRC2", bamFiles, value=TRUE)</pre>
> # use biomart
> txDbName <- "biomart"
> biomart <- "ENSEMBL_MART_ENSEMBL"</pre>
                                              # use archive to get ensembl 65
> dataset <- "mmusculus_gene_ensembl"</pre>
> host <- "dec2011.archive.ensembl.org"</pre>
                                                  # use ensembl 65 for annotation
> # compute transcript abundance in RIP
> ripRPKM <- computeRPKM(bamFiles=grep(pattern="SRR039214", bamFiles, value=TRU
                          dataset=dataset, moreGeneInfo=TRUE, justRPKM=FALSE,
+
                          idType="ensembl_transcript_id", txDbName=txDbName,
                          biomart=biomart, host=host, by="tx")
> # compute transcript abundance in RIP and control as well as
> # foldchnage in RIP over control
> rulebase.results <- rulebaseRIPSeek(bamFiles=bamFiles, cNAME=cNAME, myMin=1,
                                        featureGRanges=ripRPKM$featureGRanges,
                                       biomart=biomart, host=host, dataset=datas
> head(ripRPKM$rpkmDF)
> df <- rulebase.results$rpkmDF</pre>
> df <- df[order(df$foldchange, decreasing=TRUE), ]</pre>
> # top 10 transcripts
> head(df, 10)
```

5 RIP-seq analysis on CCNT1 data

As promised earlier, we will apply ripSeek to the CCNT1 data and annotate the *de novo* putative RIP regions with the latest online Ensembl annotation based on human genome reference NCBI37/hg19.

```
> maxBinSize <- NULL
                              # max bin size in automatic bin size selection
> multicore <- TRUE
                                     # use multicore
> strandType <- "+"
                                     # set strand type to minus strand
> biomart <- "ensembl"</pre>
                                  # use archive to get ensembl 65
> biomaRt_dataset <- "hsapiens_gene_ensembl" # human dataset id name
> goAnno <- "org.Hs.eg.db"</pre>
                                                # GO annotation database
> ############### run main function ripSeek to predict RIP #####################
> seekOut.CCNT1 <- ripSeek(bamPath = bamFiles, cNAME = cNAME,
                      reverseComplement = TRUE, genomeBuild = "hg19",
                      strandType = strandType,
                      uniqueHit = TRUE, assignMultihits = TRUE,
                      rerunWithDisambiguatedMultihits = TRUE,
                     binSize=binSize, minBinSize = minBinSize,
                     maxBinSize = maxBinSize,
                     biomart=biomart, goAnno = goAnno,
                     biomaRt_dataset = biomaRt_dataset,
                     multicore=multicore, outDir=outDir)
> df <- elementMetadata(seekOut.CCNT1$annotatedRIPGR$sigGRangesAnnotated)
> # order by eFDR
> df <- df[order(df$eFDR), ]</pre>
> # get top 100 predictions
> df.top20 <- head(df, 20)
> # examine known PRC2-associated lncRNA transcripts
> df.top20[grep("RN7SK", df$external_gene_id)[1], ]
> list.files(outDir)
> viewRIP(seekOut.CCNT1$RIPGRList$chr6, seekOut.CCNT1$mainSeekOutputRIP$alignGa
> # use biomart
> txDbName <- "biomart"
> biomart <- "ensembl"</pre>
> dataset <- "hsapiens_gene_ensembl"</pre>
> # compute transcript abundance in RIP
> ripRPKM <- computeRPKM(bamFiles=bamFiles[1],</pre>
                          dataset=dataset, moreGeneInfo=TRUE, justRPKM=FALSE,
                          idType="ensembl_transcript_id", txDbName=txDbName,
                         biomart=biomart, by="tx")
> # compute transcript abundance in RIP and control as well as
> # foldchnage in RIP over control
> rulebase.results <- rulebaseRIPSeek(bamFiles=bamFiles, cNAME=cNAME, myMin=1,
                                       featureGRanges=ripRPKM$featureGRanges,
                                       biomart=biomart, dataset=dataset)
> head(ripRPKM$rpkmDF)
> df <- rulebase.results$rpkmDF</pre>
> df <- df[order(df$foldchange, decreasing=TRUE), ]</pre>
> # top 10 transcripts
> head(df, 10)
```

6 Session Info

```
> sessionInfo()
R version 3.3.0 (2016-05-03)
Platform: x86_64-pc-linux-gnu (64-bit)
Running under: Ubuntu 14.04.4 LTS
locale:
 [1] LC_CTYPE=en_US.UTF-8
                               LC NUMERIC=C
 [3] LC_TIME=en_US.UTF-8
                               LC_COLLATE=C
 [5] LC_MONETARY=en_US.UTF-8
                               LC_MESSAGES=en_US.UTF-8
 [7] LC_PAPER=en_US.UTF-8
                               LC NAME=C
 [9] LC_ADDRESS=C
                               LC_TELEPHONE=C
[11] LC_MEASUREMENT=en_US.UTF-8 LC_IDENTIFICATION=C
attached base packages:
[1] parallel stats4 stats
                                 graphics grDevices utils
                                                               datasets
[8] methods
             base
other attached packages:
 [1] RIPSeeker_1.12.0
                               rtracklayer_1.32.0
 [3] GenomicAlignments_1.8.0
                               Rsamtools_1.24.0
 [5] Biostrings 2.40.0
                               XVector_0.12.0
 [7] SummarizedExperiment_1.2.0 Biobase_2.32.0
 [9] GenomicRanges_1.24.0 GenomeInfoDb_1.8.0
[11] IRanges_2.6.0
                               S4Vectors_0.10.0
[13] BiocGenerics_0.18.0
loaded via a namespace (and not attached):
                                                            BiocParallel_1.6.0
[1] XML_3.98-1.4 bitops_1.0-6
                                        zlibbioc_1.18.0
[5] tools_3.3.0
                      RCurl 1.95-4.8
```

References

P. Aboyoun, H. Pages, and M. Lawrence. *GenomicRanges: Representation and manipulation of genomic intervals*. R package version 1.8.9.

Simon Anders and Wolfgang Huber. Differential expression analysis for sequence count data. *Genome Biology*, 11(10):R106, October 2010.

Christopher Bishop. *Pattern recognition and machine learning*. Number 605–631 in Information Science and Statistics. Springer Science, 2006.

Nicole Cloonan, Alistair R R Forrest, Gabriel Kolle, Brooke B A Gardiner, Geoffrey J Faulkner, Mellissa K Brown, Darrin F Taylor, Anita L Steptoe, Shivangi Wani, Graeme Bethel, Alan J Robertson, Andrew C Perkins, Stephen J Bruce, Clarence C Lee, Swati S Ranade, Heather E Peckham, Jonathan M Manning, Kevin J Mckernan, and Sean M Grimmond. Stem cell transcriptome profiling via massive-scale mRNA sequencing. Nature Methods, 5(7):613–619, July 2008. Strand specific protocol.

- S Durinck, PT Spellman, and E Birney. Mapping identifiers for the integration of genomic datasets with the R/Bioconductor package biomaRt. *Nature protocols*, 4 (8):1184–1191, 2009.
- Steffen Durinck, Yves Moreau, Arek Kasprzyk, Sean Davis, Bart De Moor, Alvis Brazma, and Wolfgang Huber. BioMart and Bioconductor: a powerful link between biological databases and microarray data analysis. *Bioinformatics (Oxford, England)*, 21(16):3439–3440, August 2005.
- Michael Lawrence, Vince Carey, and Robert Gentleman. *rtracklayer: R interface to genome browsers and their annotation tracks.* R package version 1.16.1.
- Zhaohui S Qin, Jianjun Yu, Jincheng Shen, Christopher A Maher, Ming Hu, Shanker Kalyana-Sundaram, Jindan Yu, and Arul M Chinnaiyan. HPeak: an HMM-based algorithm for defining read-enriched regions in ChIP-Seq data. *BMC Bioinformatics*, 11:369, 2010.
- L.R Rabiner. A tutorial on hidden Markov models and selected applications in speech recognition. In *Proceedings of the IEEE*, pages 257–286, 1989.
- James T Robinson, Helga Thorvaldsdóttir, Wendy Winckler, Mitchell Guttman, Eric S Lander, Gad Getz, and Jill P Mesirov. Integrative genomics viewer. *Nature Publishing Group*, 29(1):24–26, 2011.
- C Trapnell, L Pachter, and S L Salzberg. TopHat: discovering splice junctions with RNA-Seq. *Bioinformatics (Oxford, England)*, 25(9):1105–1111, May 2009.
- Jing Zhao, Toshiro K Ohsumi, Johnny T Kung, Yuya Ogawa, Daniel J Grau, Kavitha Sarma, Ji Joon Song, Robert E Kingston, Mark Borowsky, and Jeannie T Lee. Genome-wide Identification of Polycomb-Associated RNAs by RIP-seq. *Molecular Cell*, 40(6):939–953, December 2010.
- Lihua J Zhu, Claude Gazin, Nathan D Lawson, Hervé Pagès, Simon M Lin, David S Lapointe, and Michael R Green. ChIPpeakAnno: a Bioconductor package to annotate ChIP-seq and ChIP-chip data. *BMC Bioinformatics*, 11:–, 2010.