

CRISPRseek user's guide

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June 2, 2015

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1 Introduction

CRISPR-Cas9 nucleases and their derivatives have rapidly become widely used tools for both genome modification and regulation of gene expression. They can create genetic changes with high efficiency in human stem cells, in model organisms such as mice and *Drosophila* and in a wide variety of other organisms. CRISPR-Cas9 nucleases create double strand DNA breaks that can facilitate a variety of genome modifications including short insertions and/or deletions (indels) or specific sequence changes introduced by homology directed repair with a DNA donor molecule. The high activity and relative ease of construction has made CRISPR-Cas9 nucleases a popular replacement for related technologies such as zinc finger nucleases and TALENs. Derivatives of CRISPR-Cas9 complexes include nickases, which only cleave one DNA strand, and gene expression regulators, which lack any DNA cleavage activity but can increase or decrease gene transcription. CRISPR-Cas9 nucleases are composed of an

RNA-protein complex that can target a variable sequence (guide RNA, abbreviated as "gRNA") that is directly adjacent to a constant motif (the "PAM" sequence). In the most widely use version from the species *Streptococcus pyogenes*, the gRNA is composed of a variable region of 20 bases and the preferred PAM sequence is an adjacent 3 base sequence of NGG (or NAG with lower activity). One potential limitation for CRISPR-Cas9 nucleases is that they can cleave at some sequences that do not precisely match the sequence targeted by the gRNA sequence. Thus, an important consideration for the design and application of CRISPR-Cas9 nucleases is the identification of gRNA regions with low rates of off-target cleavage.

Several computational analyses can assist with the construction and application of CRISPR-Cas9 nucleases with high on-target and low off-target cleavage. First, gRNA sequences can be evaluated for possible off-target sequences in the target genome. Second, sequences flanking possible off-target sequences can be reported to assist in the experimental analysis of off-target cleavage and to determine if these sequences are within critical regions for gene function such as exons. Third, specific arrangements of target sequences can be selected; one alternate approach to lower off-target rates is to introduce pairs of CRISPR-Cas9 nickases, which will only create double strand DNA breaks at genomic regions where the two sites have the proper spacing and orientation. Finally, in some applications, it is useful to use restriction enzyme sequences that overlap CRISPR-Cas9 target sites in order to monitor cleavage events. We developed [CRISPRseek](#) package that identifies candidate CRISPR-Cas9 target sequences within a given input sequence using a variety of experimentally useful constraints and also reports and ranks potential off-target sequences for each recovered target sequence. CRISPRseek will automatically find potential target sequences that are/are not present as pairs that can be used as double nickases or that have/don't have overlapping restriction enzyme cut site(s). It will then search genome-wide for off-targets with a user defined maximum number of mismatches, calculate the score of each off-target based on mismatches in the off-target and a penalty weight matrix, filter off-targets with user-defined criteria, and annotate off-targets with flanking sequences, and whether located in exon or not. Several reports are generated including a summary report with gRNAs ranked by total topN off-target score, restriction enzyme cut sites and possible paired gRNAs. Detailed paired gRNAs information, restriction enzyme cut sites, and off-target sequences and scores are stored in separate files in the output directory specified by the user. In total, four tab delimited files are generated in the output directory: OfftargetAnalysis.xls (off-target details), Summary.xls (gRNA summary), REcutDetails.xls (restriction enzyme cut sites of each gRNA), and pairedgRNAs.xls (potential paired gRNAs). These reports provide a comprehensive set of information to identify, select and utilize *Streptococcus pyogenes* CRISPR-Cas9 nucleases and their derivatives. The package can also be readily modified to accept different gRNA and PAM sequence requirements for CRISPR-Cas9 complexes from other bacterial species that can be used to target alternative genomic sequences. The package can also be modified to incorporate improved weight matrices for scoring off-target sequences as new experimental and computational results become available for CRISPR-Cas9 nucleases for *Streptococcus pyogenes* and other species.

2 Examples of using CRISPRseek

In this guide, we will illustrate five different gRNA search scenarios with a human sequence. First load [CRISPRseek](#), [BSgenome.Hsapiens.UCSC.hg19](#) and [TxDb.Hsapiens.UCSC.hg19.knownGene](#). Then

specify the sequence file path as `inputFilePath`, a fasta/fastq file containing a genomic sequence, restriction enzyme pattern file as `REpatternFile` and output directory as `outputDir`. Once the analysis is done, analysis results will be saved in the output directory.

To find `BSgenome` of other species, please refer to available.genomes in the `BSgenome` package. For example, `BSgenome.Hsapiens.UCSC.hg19` for hg19, `BSgenome.Mmusculus.UCSC.mm10` for mm10, `BSgenome.Celegans.UCSC.ce6` for ce6, `BSgenome.Rnorvegicus.UCSC.rn5` for rn5, `BSgenome.Drerio.UCSC.danRer7` for Zv9, and `BSgenome.Dmelanogaster.UCSC.dm3` for dm3

To create and use `TxDb` objects, please refer to the `GenomicFeatures` package. For a list of existing `TxDb` objects, please search for annotation package starting with `Txdb` at <http://www.bioconductor.org/packages/release/BiocViews.html>, such as `Txdb.Rnorvegicus.UCSC.rn5.refGene` for rat, `Txdb.Mmusculus.UCSC.mm10.knownGene` for mouse, `Txdb.Hsapiens.UCSC.hg19.knownGene` for human, `Txdb.Dmelanogaster.UCSC.dm3.ensGene` for Drosophila and `Txdb.Celegans.UCSC.ce6.ensGene` for C.elegans

`org.Hs.eg.db` is the gene ID mapping package for human. For a list of existing `OrgDb` packages, please search for `OrgDb` at <http://www.bioconductor.org/packages/release/BiocViews.html>, such as `org.Rn.eg.db` for rat `org.Mm.eg.db` for mouse `org.Dm.eg.db` for Drosophila `org.Ce.eg.db` for C.elegans

```
> library(CRISPRseek)
> library(BSgenome.Hsapiens.UCSC.hg19)
> library(Txdb.Hsapiens.UCSC.hg19.knownGene)
> library(org.Hs.eg.db)
> outputDir <- getwd()
> inputFilePath <- system.file('extdata', 'inputseq.fa', package = 'CRISPRseek')
> REpatternFile <- system.file('extdata', 'NEBenzymes.fa', package = 'CRISPRseek')
```

2.1 Scenario 1: Finding paired gRNAs with restriction enzyme cut site(s)

Paired gRNAs in proper spacing and orientation give more specificity and gRNAs overlap with restriction enzyme cut sites facilitates cleavage monitoring. Calling the function `offTargetAnalysis` with `findPairedgRNAOnly=TRUE` and `findgRNAsWithREcutOnly=TRUE` results in searching, scoring and annotating gRNAs that are in paired configuration and at least one of the pairs overlap a restriction enzyme cut site. To be considered as a pair, gap between forward gRNA and the corresponding reverse gRNA needs to be (`min.gap`, `max.gap`) inclusive and the reverse gRNA must sit before the forward gRNA. The default (`min.gap`, `max.gap`) is (0,20). Please note that `chromToSearch` is set to `chrX` here for speed purpose, usually you would set it to `all`, which is the default. In order for a gRNA to be considered overlap with restriction enzyme cut site, the enzyme cut pattern must overlap with one of the gRNA positions specified in `overlap.gRNA.positions`, default position 17 and 18. Please note that `max.mismatch` allowed for off-target finding is set to 4 by default, set it to a larger number will significantly slow down the search. `org.Hs.egSYMBOL` is `entrezID` to gene symbol mapping in `org.Hs.eg.db` package for human. For detailed parameter settings using function `offTargetAnalysis`, please type `help(offTargetAnalysis)`

```
> results <- offTargetAnalysis(inputFilePath, findgRNAsWithREcutOnly = TRUE,
+ REpatternFile = REpatternFile, findPairedgRNAOnly = TRUE,
+ BSgenomeName = Hsapiens, chromToSearch = "chrX", min.gap = 0, max.gap = 20,
+ txdb = Txdb.Hsapiens.UCSC.hg19.knownGene, orgAnn = org.Hs.egSYMBOL,
```

```
+ max.mismatch = 0,overlap.gRNA.positions = c(17, 18),
+ outputDir = outputDir,overwrite = TRUE)

Validating input ...
Searching for gRNAs ...
>>> Finding all hits in sequence chrX ...
>>> DONE searching
Building feature vectors for scoring ...
Calculating scores ...
Annotating, filtering and generating reports ...
Done annotating
Add paired information...
Add RE information...
write gRNAs to bed file...
Scan for REsites in flanking region...
[1] "MY_ERROR: Error in library(GeneRfold): there is no package called 'GeneRfold'\n"
Done. Please check output files in directory /tmp/RtmpRuu5ZV/Rbuild49a8704896c4/CRISPRseek/vignettes/
```

2.2 Scenario 2: Finding paired gRNAs with/without restriction enzyme cut site(s)

Calling the function `offTargetAnalysis` with `findPairedgRNAOnly = TRUE` and `findgRNAsWithREcutOnly = FALSE` results in searching, scoring and annotating gRNAs that are in paired configuration without requiring overlap any restriction enzyme cut site. The gRNAs will be annotated with restriction enzyme cut sites for users to review later.

```
> results <- offTargetAnalysis(inputFilePath, findgRNAsWithREcutOnly = FALSE,
+ REpatternFile = REpatternFile,findPairedgRNAOnly = TRUE,
+ BSgenomeName = Hsapiens, chromToSearch = "chrX",
+ txdb = TxDb.Hsapiens.UCSC.hg19.knownGene,
+ orgAnn = org.Hs.egSYMBOL,
+ max.mismatch = 1, outputDir = outputDir, overwrite = TRUE)

Validating input ...
Searching for gRNAs ...
>>> Finding all hits in sequence chrX ...
>>> DONE searching
Building feature vectors for scoring ...
Calculating scores ...
Annotating, filtering and generating reports ...
Done annotating
Add paired information...
Add RE information...
write gRNAs to bed file...
Scan for REsites in flanking region...
[1] "MY_ERROR: Error in library(GeneRfold): there is no package called 'GeneRfold'\n"
Done. Please check output files in directory /tmp/RtmpRuu5ZV/Rbuild49a8704896c4/CRISPRseek/vignettes/
```

2.3 Scenario 3: Finding all gRNAs with restriction enzyme cut site(s)

Calling the function `offTargetAnalysis` with `findPairedgRNAOnly=FALSE` and `findgRNAsWithREcutOnly = TRUE` results in searching, scoring and annotating all gRNAs (paired and not paired) overlap restriction enzyme cut site(s) and off-targets. The gRNAs will be annotated with paired information for users to review later.

```
> results <- offTargetAnalysis(inputFilePath, findgRNAsWithREcutOnly = TRUE,
+ REpatternFile = REpatternFile, findPairedgRNAOnly = FALSE,
+ BSgenomeName = Hsapiens, chromToSearch = "chrX",
```

```
+ txdb = TxDb.Hsapiens.UCSC.hg19.knownGene,
+ orgAnn = org.Hs.egSYMBOL,
+ max.mismatch = 1, outputDir = outputDir, overwrite = TRUE)

Validating input ...
Searching for gRNAs ...
>>> Finding all hits in sequence chrX ...
>>> DONE searching
Building feature vectors for scoring ...
Calculating scores ...
Annotating, filtering and generating reports ...
Done annotating
Add paired information...
Add RE information...
write gRNAs to bed file...
Scan for REsites in flanking region...
[1] "MY_ERROR: Error in library(GeneRfold): there is no package called 'GeneRfold'\n"
Done. Please check output files in directory /tmp/RtmpRuu5ZV/Rbuild49a8704896c4/CRISPRseek/vignettes/
```

2.4 Scenario 4: Finding all gRNAs

Calling the function `offTargetAnalysis` with `findPairedgRNAOnly = FALSE` and `findgRNAsWithREcutOnly = FALSE` results in searching, scoring and annotating all gRNAs and off-targets. The gRNAs will be annotated with paired information and restriction enzyme cut sites for users to review later. Please note that this search will be the slowest among all type of searches aforementioned.

```
> results <- offTargetAnalysis(inputFilePath, findgRNAsWithREcutOnly = FALSE,
+ REpatternFile = REpatternFile, findPairedgRNAOnly = FALSE,
+ BSgenomeName = Hsapiens, chromToSearch = "chrX",
+ txdb = TxDb.Hsapiens.UCSC.hg19.knownGene,
+ orgAnn = org.Hs.egSYMBOL,
+ max.mismatch = 1, outputDir = outputDir, overwrite = TRUE)

Validating input ...
Searching for gRNAs ...
>>> Finding all hits in sequence chrX ...
>>> DONE searching
Building feature vectors for scoring ...
Calculating scores ...
Annotating, filtering and generating reports ...
Done annotating
Add paired information...
Add RE information...
write gRNAs to bed file...
Scan for REsites in flanking region...
[1] "MY_ERROR: Error in library(GeneRfold): there is no package called 'GeneRfold'\n"
Done. Please check output files in directory /tmp/RtmpRuu5ZV/Rbuild49a8704896c4/CRISPRseek/vignettes/
```

2.5 Scenario 5: Target and off-target analysis for user specified gRNAs

Calling the function `offTargetAnalysis` with `findgRNAs = FALSE` results in target and off-target searching, scoring and annotating for the input gRNAs. The gRNAs will be annotated with restriction enzyme cut sites for users to review later. However, paired information will not be available.

```
> gRNAFilePath <- system.file('extdata', 'testHsap_GATA1_ex2_gRNA1.fa',
+ package = 'CRISPRseek')
> results <- offTargetAnalysis(inputFilePath = gRNAFilePath,
+ findgRNAsWithREcutOnly = FALSE, REpatternFile = REpatternFile,
+ findPairedgRNAOnly = FALSE, findgRNAs = FALSE,
```

```
+ BSgenomeName = Hsapiens, chromToSearch = 'chrX',
+ txdb = TxDb.Hsapiens.UCSC.hg19.knownGene,
+ orgAnn = org.Hs.egSYMBOL,
+ max.mismatch = 1, outputDir = outputDir, overwrite = TRUE)

Validating input ...
>>> Finding all hits in sequence chrX ...
>>> DONE searching
Building feature vectors for scoring ...
Calculating scores ...
Annotating, filtering and generating reports ...
Done annotating
Add paired information...
Add RE information...
write gRNAs to bed file...
Scan for REsites in flanking region...
[1] "MY_ERROR: Error in library(GeneRfold): there is no package called 'GeneRfold'\n"
Done. Please check output files in directory /tmp/RtmpRuu5ZV/Rbuild49a8704896c4/CRISPRseek/vignettes/
```

2.6 Scenario 6. Quick gRNA finding without target or off-target analysis

Calling the function `offTargetAnalysis` with `chromToSearch = ""` results in quick gRNA search without performing on-target and off-target analysis. Parameters `findgRNAsWithREcutOnly` and `findPairedgRNAOnly` can be tuned to indicate whether searching for gRNAs overlap restriction enzyme cut sites or not, and whether searching for gRNAs in paired configuration or not.

```
> results <- offTargetAnalysis(inputFilePath, findgRNAsWithREcutOnly = TRUE,
+ REpatternFile = REpatternFile, findPairedgRNAOnly = TRUE,
+ chromToSearch = "", outputDir = outputDir, overwrite = TRUE)

Validating input ...
Searching for gRNAs ...
Done. Please check output files in directory /tmp/RtmpRuu5ZV/Rbuild49a8704896c4/CRISPRseek/vignettes/
```

2.7 Scenario 7. Quick gRNA finding with gRNA efficiency prediction

Calling the function `offTargetAnalysis` with `max.mismatch = 0` results in quick gRNA search with gRNA efficiency prediction without off-target analysis. Parameters `findgRNAsWithREcutOnly` and `findPairedgRNAOnly` can be tuned to indicate whether searching for gRNAs overlap restriction enzyme cut sites or not, and whether searching for gRNAs in paired configuration or not.

```
> results <- offTargetAnalysis(inputFilePath, findgRNAsWithREcutOnly = TRUE,
+ annotateExon = FALSE, findPairedgRNAOnly = TRUE, chromToSearch = "chrX",
+ max.mismatch = 0, BSgenomeName = Hsapiens, outputDir = outputDir, overwrite = TRUE)

Validating input ...
Searching for gRNAs ...
>>> Finding all hits in sequence chrX ...
>>> DONE searching
Building feature vectors for scoring ...
Calculating scores ...
Annotating, filtering and generating reports ...
Done annotating
Add paired information...
Add RE information...
write gRNAs to bed file...
Scan for REsites in flanking region...
[1] "MY_ERROR: Error in library(GeneRfold): there is no package called 'GeneRfold'\n"
Done. Please check output files in directory /tmp/RtmpRuu5ZV/Rbuild49a8704896c4/CRISPRseek/vignettes/
```

2.8 Scenario 8. Find potential gRNAs preferentially targeting one of two alleles without running time-consuming off-target analysis on all possible gRNAs.

Below is an example to search for all gRNAs that target at least one of the alleles. Two files are provided containing sequences that differ by a single nucleotide polymorphism (SNP). The results are saved in file scoresFor2InputSequences.xls in outputDir directory.

```
> inputFile1Path <- system.file("extdata", "rs362331C.fa", package = "CRISPRseek")
> inputFile2Path <- system.file("extdata", "rs362331T.fa", package = "CRISPRseek")
> REpatternFile <- system.file("extdata", "NEBenzymes.fa", package = "CRISPRseek")
> seqs <- compare2Sequences(inputFile1Path, inputFile2Path,
+   outputDir = outputDir , REpatternFile = REpatternFile,
+   overwrite = TRUE)

search for gRNAs for input file1...
Validating input ...
Searching for gRNAs ...
Done. Please check output files in directory /tmp/RtmpRuu5ZV/Rbuild49a8704896c4/CRISPRseek/vignettes/rs362331C.fa/
search for gRNAs for input file2...
Validating input ...
Searching for gRNAs ...
Done. Please check output files in directory /tmp/RtmpRuu5ZV/Rbuild49a8704896c4/CRISPRseek/vignettes/rs362331T.fa/
[1] "Scoring ..."
finish off-target search in sequence 2
finish off-target search in sequence 1
finish feature vector building
finish score calculation
[1] "MY_ERROR: Error in library(GeneRfold): there is no package called 'GeneRfold'"
[1] "Done!"

> seqs

      name          gRNAPlusPAM          targetInSeq1
1  rs362331T_gR9r  GTAGATGAGGGAGCAGGCGTGGG  GTGGATGAGGGAGCAGGCGTGGG
2  rs362331T_gR38f  CTA CTGTGTGCACTTCATCCTGG  CCACTGTGTGCACTTCATCCTGG
3  rs362331T_gR22r  TGAAGTGCACACAGTAGATGAGG  TGAAGTGCACACAGTGGATGAGG
4  rs362331T_gR21r  GAAAGTGCACACAGTAGATGAGG  GAAAGTGCACACAGTGGATGAGG
5  rs362331T_gR15r  CACACAGTAGATGAGGGAGCAGG  CACACAGTGGATGAGGGAGCAGG
6  rs362331T_gR10r  AGTAGATGAGGGAGCAGGCGTGG  AGTGGATGAGGGAGCAGGCGTGG
7   rs362331C_gR9r  GTGGATGAGGGAGCAGGCGTGGG  GTGGATGAGGGAGCAGGCGTGGG
8  rs362331C_gR38f  CCACTGTGTGCACTTCATCCTGG  CCACTGTGTGCACTTCATCCTGG
9  rs362331C_gR28r  CCAGGATGAAGTGCACACAGTGG  CCAGGATGAAGTGCACACAGTGG
10 rs362331C_gR22r  TGAAGTGCACACAGTGGATGAGG  TGAAGTGCACACAGTGGATGAGG
11 rs362331C_gR21r  GAAAGTGCACACAGTGGATGAGG  GAAAGTGCACACAGTGGATGAGG
12 rs362331C_gR15r  CACACAGTGGATGAGGGAGCAGG  CACACAGTGGATGAGGGAGCAGG
13 rs362331C_gR10r  AGTGGATGAGGGAGCAGGCGTGG  AGTGGATGAGGGAGCAGGCGTGG

      targetInSeq2  guideAlignment20offTarget  offTargetStrand  scoreForSeq1
1  GTAGATGAGGGAGCAGGCGTGGG  ..G.....  -  98.6
2  CTA CTGTGTGCACTTCATCCTGG  .C.....  +  100
3  TGAAGTGCACACAGTAGATGAGG  .....G....  -  17.2
4  GAAAGTGCACACAGTAGATGAGG  .....G....  -  26.8
5  CACACAGTAGATGAGGGAGCAGG  .....G.....  -  61.1
6  AGTAGATGAGGGAGCAGGCGTGG  ..G.....  -  100
7  GTAGATGAGGGAGCAGGCGTGGG  ..A.....  -  100
8  CTA CTGTGTGCACTTCATCCTGG  .T.....  +  100
9  CCAGGATGAAGTGCACACAGTAG  .....G....  -  100
10 TGAAGTGCACACAGTAGATGAGG  .....A....  -  100
11 GAAAGTGCACACAGTAGATGAGG  .....A....  -  100
12 CACACAGTAGATGAGGGAGCAGG  .....A.....  -  100
13 AGTAGATGAGGGAGCAGGCGTGG  ..A.....  -  100
      scoreForSeq2  n.mismatch  offTarget  targetSeqName  gRNAefficacy
1  100  1  1  rs362331C:3-25  rs362331T extended sequence too short
2  100  1  1  rs362331C:22-44  rs362331T extended sequence too short
3  100  1  1  rs362331C:16-38  rs362331T 0.118513953473509
4  100  1  1  rs362331C:15-37  rs362331T 0.229116108934864
```

5	100	1	rs362331C:9-31	rs362331T	0.0546177253478725
6	100	1	rs362331C:4-26	rs362331T	0.161377523263354
7	98.6	1	rs362331T:3-25	rs362331C	extended sequence too short
8	100	1	rs362331T:22-44	rs362331C	extended sequence too short
9	100	0	rs362331T:22-44	rs362331C	extended sequence too short
10	17.2	1	rs362331T:16-38	rs362331C	0.0978516718170484
11	26.8	1	rs362331T:15-37	rs362331C	0.129816665166513
12	61.1	1	rs362331T:9-31	rs362331C	0.0169291602963948
13	100	1	rs362331T:4-26	rs362331C	0.140082863263818

scoreDiff	
1	-1.4
2	0
3	-82.8
4	-73.2
5	-38.9
6	0
7	1.4
8	0
9	0
10	82.8
11	73.2
12	38.9
13	0

rs362331C.fa and rs362331T.fa are the names of the two input files. The output file will list all of the possible gRNA sequences for each of the two input sequences and provide a cleavage score for each of the two input sequences. To preferentially target one allele, select gRNA sequences that have the lowest score for the other allele. Selected gRNAs can then be examined for off-target sequences as described in Step 6.

3 References

References

- [1] Mali P. et al., CAS9 transcriptional activators for target specificity screening and paired nickases for cooperative genome engineering. Nat Biotechnol. 2013. 31(9):833-8
- [2] Hsu, P.D. et al., DNA targeting specificity of rNA-guided Cas9 nucleases. Nat Biotechnol. 2013. 31:827-834.
- [3] . Rational design of highly active sgRNAs for CRISPR-Cas9-mediated gene inactivation. Nat Biotechnol. 2014 Sep 3. doi: 10.1038 nbt.3026
- [4] Lihua Julie Zhu, Benjamin R. Holmes, Neil Aronin and Michael Brodsky. CRISPRseek: a Bioconductor package to identify target-specific guide RNAs for CRISPR-Cas9 genome-editing systems. Plos One Sept 23rd 2014

4 Session Info

```
> sessionInfo()
```

```
R version 3.2.0 (2015-04-16)
Platform: x86_64-unknown-linux-gnu (64-bit)
Running under: Ubuntu 14.04.2 LTS
```

```
locale:
```

```
[1] LC_CTYPE=en_US.UTF-8      LC_NUMERIC=C              LC_TIME=en_US.UTF-8
[4] LC_COLLATE=C              LC_MONETARY=en_US.UTF-8  LC_MESSAGES=en_US.UTF-8
[7] LC_PAPER=en_US.UTF-8     LC_NAME=C                 LC_ADDRESS=C
[10] LC_TELEPHONE=C           LC_MEASUREMENT=en_US.UTF-8 LC_IDENTIFICATION=C
```

```
attached base packages:
```

```
[1] stats4      parallel    stats      graphics  grDevices  utils      datasets  methods
[9] base
```

```
other attached packages:
```

```
[1] org.Hs.eg.db_3.1.2          RSQLite_1.0.0
[3] DBI_0.3.1                  TxDb.Hsapiens.UCSC.hg19.knownGene_3.1.2
[5] GenomicFeatures_1.20.1     AnnotationDbi_1.30.1
[7] Biobase_2.28.0             BSgenome.Hsapiens.UCSC.hg19_1.4.0
[9] CRISPRseek_1.8.1          seqinr_3.1-3
[11] ade4_1.7-2                 BSgenome_1.36.0
[13] rtracklayer_1.28.4         GenomicRanges_1.20.4
[15] GenomeInfoDb_1.4.0        Biostrings_2.36.1
[17] XVector_0.8.0              IRanges_2.2.3
[19] S4Vectors_0.6.0           BiocGenerics_0.14.0
```

```
loaded via a namespace (and not attached):
```

```
[1] zlibbioc_1.14.0           GenomicAlignments_1.4.1 BiocParallel_1.2.2
[4] tools_3.2.0               lambda.r_1.1.7          futile.logger_1.4.1
[7] futile.options_1.0.0     bitops_1.0-6           biomaRt_2.24.0
[10] RCurl_1.95-4.6           Rsamtools_1.20.4       XML_3.98-1.2
[13] BiocStyle_1.6.0
```