# Introduction to crisprDesign

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2022-08-30

## 1 Introduction

crisprDesign is the core package of the crisprVerse, and plays the role of a one-stop shop for designing and annotating CRISPR guide RNA (gRNA) sequences. This includes the characterization of on-targets and off-targets using different aligners, on- and off-target scoring, gene context annotation, SNP annotation, sequence feature characterization, repeat annotation, and many more.

The software was developed to be as applicable and generalizable as possible.

It currently support five types of CRISPR modalities (modes of perturbations): CRISPR knockout (CRISPRko), CRISPR activation (CRISPRa), CRISPR interference (CRISPRi), CRISPR base editing (CRISPRbe), and CRISPR knockdown (CRISPRkd) (see Kampmann (2018) for a review of CRISPR modalities).

It utilizes the crisprBase package to enable gRNA design for any CRISPR nuclease and base editor via the CrisprNuclease and BaseEditor classes, respectively. Nucleases that are commonly used in the field are provided, including DNA-targeting nucleases (e.g. SpCas9, AsCas12a) and RNA-targeting nucleases (e.g. CasRx (RfxCas13d)).

crisprDesign is fully developed to work with the genome of any organism, and can also be used to design gRNAs targeting custom DNA sequences.

Finally, more specialized gRNA design functionalities are also available, including design for optical pooled screening (OPS), paired gRNA design, and gRNA filtering and ranking functionalities.

This vignette is meant to be an overview of the main features included in the package, using toy examples for the sake of time (the vignette has to compile within a few minutes, as required by Bioconductor). For detailed and comprehensive tutorials, please visit our crisprVerse tutorials page.

### 2 Installation

crisprDesign can be installed from from the Bioconductor devel branch using the following commands in a fresh R session:

```
if (!require("BiocManager", quietly = TRUE))
    install.packages("BiocManager")

BiocManager::install(version="devel")
BiocManager::install("crisprDesign")
```

Users interested in contributing to crisprDesign might want to look at the following CRISPR-related package dependencies:

- crisprBase: core CRISPR functions and S4 objects
- crisprBowtie: aligns gRNA spacers to genomes using the ungapped aligner bowtie
- crisprBwa: aligns gRNA spacers to genomes using the ungapped aligner BWA

- crisprScore: implements state-of-the-art on- and off-target scoring algorithms
- crisprViz: gRNA visualization using genomic tracks

You can contribute to the package by submitting pull requests to our GitHub repo.

## 3 Terminology

CRISPR nucleases are examples of RNA-guided endonucleases. They require two binding components for cleavage. First, the nuclease needs to recognize a constant nucleotide motif in the target DNA called the protospacer adjacent motif (PAM) sequence. Second, the gRNA, which guides the nuclease to the target sequence, needs to bind to a complementary sequence adjacent to the PAM sequence, called the **protospacer** sequence. The latter can be thought of as a variable binding motif that can be specified by designing corresponding gRNA sequences.

The **spacer** sequence is used in the gRNA construct to guide the CRISPR nuclease to the target **protospacer** sequence in the host genome.

For DNA-targeting nucleases, the nucleotide sequence of the spacer and protospacer are identical. For RNA-targeting nucleases, they are the reverse complement of each other.

While a gRNA spacer sequence may not always uniquely target the host genome (i.e. it may map to multiple protospacers in the host genome), we can, for a given reference genome, uniquely identify a protospacer sequence with a combination of 3 attributes:

- chr: chromosome name
- strand: forward (+) or reverse (-)
- pam\_site: genomic coordinate of the first nucleotide of the nuclease-specific PAM sequence (e.g. for SpCas9, the "N" in the NGG PAM sequence; for AsCas12a, the first "T" of the TTTV PAM sequence)

For CRISPRko, we use an additional genomic coordinate, called cut\_site, to represent where the double-stranded break (DSB) occurs. For SpCas9, the cut site (blunt-ended dsDNA break) is located 4nt upstream of the pam\_site (PAM-proximal editing). For AsCas12a, the 5nt 5' overhang dsDNA break will cause a cut 19nt after the PAM sequence on the targeted strand, and 23nt after the PAM sequence on the opposite strand (PAM-distal editing).

# 4 CRISPRko design

We will illustrate the main functionalities of crisprDesign by performing a common task: designing gRNAs to knock out a coding gene. In our example, we will design gRNAs for the wildtype SpCas9 nuclease, with spacers having a length of 20nt.

library(crisprDesign)

## 4.1 Nuclease specification

The crisprBase package provides functionalities to create objects that store information about CRISPR nucleases, and functions to interact with those objects (see the crisprBase vignette). It also provides commonly-used CRISPR nucleases. Let's look at the SpCas9 nuclease object:

```
library(crisprBase)
data(SpCas9, package="crisprBase")
SpCas9
```

```
## Class: CrisprNuclease
## Name: SpCas9
## Target type: DNA
```

## Metadata: list of length 1

The three motifs (NGG, NAG and NGA) represent the recognized PAM sequences by SpCas9, and the weights indicate a recognition score. The canonical PAM sequence NGG is fully recognized (weight of 1), while the two non-canonical PAM sequences NAG and NGA are much less tolerated.

The spacer sequence is located on the 5-prime end with respect to the PAM sequence, and the default spacer sequence length is 20 nucleotides. If necessary, we can change the spacer length using the function crisprBase::spacerLength. Let's see what the protospacer construct looks like by using prototypeSequence:

```
prototypeSequence(SpCas9)
```

```
## [1] "5'--SSSSSSSSSSSSSSSSS[NGG]--3'"
```

## 4.2 Target DNA specification

PAMs: NGG, NAG, NGA

##

As an example, we will design gRNAs that knockout the human gene IQSEC3 by finding all protospacer sequences located in the coding region (CDS) of IQSEC3.

To do so, we need to create a GRanges object that defines the genomic coordinates of the CDS of IQSEC3 in a reference genome.

The toy dataset grListExample object in crisprDesign contains gene coordinates in hg38 for exons of all human IQSEC3 isoforms, and was obtained by converting an Ensembl TxDb object into a GRangesList object using the TxDb2GRangesList convenience function in crisprDesign.

```
data(grListExample, package="crisprDesign")
```

The queryTxObject function allows us to query such objects for a specific gene and feature. Here, we obtain a GRanges object containing the CDS coordinates of IQSEC3:

We will only consider the first exon to speed up design:

```
gr <- gr[1]
```

### 4.3 Designing spacer sequences

findSpacers is the main function to obtain a list of all possible spacer sequences targeting protospacers located in the target DNA sequence(s). If a GRanges object is provided as input, a BSgenome object (object containing sequences of a reference genome) will need to be provided as well:

```
## GuideSet object with 123 ranges and 5 metadata columns:
##
                                                        protospacer
                 segnames
                             ranges strand |
                                                                                 pam
##
                    <Rle> <IRanges>
                                      <Rle> |
                                                     <DNAStringSet> <DNAStringSet>
##
                                          - | CGCGCACCGGATTCTCCAGC
                    chr12
                              66893
                                                                                 AGG
       spacer_1
##
       spacer 2
                    chr12
                              66896
                                          + | GGGCGGCATGGAGAGCCTGC
                                                                                 TGG
##
       spacer 3
                              66905
                                          + | GGAGAGCCTGCTGGAGAATC
                                                                                 CGG
                    chr12
##
                              66906
                                          - | AGGTAGAGCACGGCGCGCAC
       spacer 4
                    chr12
                                                                                 CGG
                                          - | GAGCTCCTTGAGGTAGAGCA
##
       spacer_5
                    chr12
                              66916
                                                                                 CGG
##
                      . . .
             . . .
                                                                                 . . .
                                          + | CACAAATCCCCCTCCGCCCT
##
     spacer_119
                    chr12
                              67407
                                                                                 CGG
##
     spacer_120
                    chr12
                              67412
                                          + | ATCCCCTCCGCCTCGGCA
                                                                                 AGG
                                                                                 GGG
##
     spacer_121
                              67413
                                          + | TCCCCCTCCGCCAA
                    chr12
##
     spacer_122
                    chr12
                              67421
                                          - | CTCACTCAGGTCTCCTGCTC
                                                                                 AGG
##
     spacer_123
                              67426
                                          + | TCGGCAAGGGCGTCCTGAGC
                    chr12
                                                                                 AGG
##
                  pam_site cut_site
                                           region
##
                 <numeric> <numeric> <character>
##
                     66893
                                66896
                                         region_1
       spacer_1
##
       spacer 2
                     66896
                                66893
                                         region 1
##
                     66905
       spacer_3
                                66902
                                         region_1
##
       spacer 4
                     66906
                                66909
                                         region 1
##
       spacer_5
                     66916
                                66919
                                         region_1
##
                       . . .
                                  . . .
##
     spacer_119
                     67407
                                67404
                                         region_1
     spacer 120
##
                     67412
                                67409
                                         region 1
##
     spacer_121
                     67413
                                67410
                                         region_1
##
     spacer_122
                     67421
                                67424
                                         region_1
##
     spacer_123
                     67426
                                67423
                                         region_1
##
##
     seqinfo: 640 sequences (1 circular) from hg38 genome
##
     crisprNuclease: SpCas9
```

This returns a GuideSet object that stores genomic coordinates for all spacer sequences found in the regions provided by gr. The GuideSet object is an extension of a GenomicRanges object that stores additional information about gRNAs.

For the subsequent sections, we will only work with a random subset of 20 spacer sequences:

```
set.seed(10)
guideSet <- guideSet[sample(seq_along((guideSet)),20)]</pre>
```

Several accessor functions are provided to extract information about the spacer sequences:

### spacers(guideSet)

```
## DNAStringSet object of length 20:
##
        width seq
                                                                  names
##
    [1]
           20 CCGAGTTGCTGCGCTGCTGC
                                                                  spacer 107
   [2]
           20 GCTCTGCTGGTTCTGCACGA
                                                                  spacer_9
##
    [3]
           20 CGGCCGCCGCGTCAGCACCA
##
                                                                  spacer 74
##
   [4]
           20 GCCCTTGCCGAGGGCGGAGG
                                                                  spacer 112
   [5]
##
           20 GGCCCCGCTGGGGCTGCTCC
                                                                  spacer_76
##
   . . .
          . . . . . .
## [16]
           20 TCCCCCTCCGCCAA
                                                                  spacer_121
## [17]
           20 CGGCAGCGGGGCCGATGACG
                                                                  spacer_34
## [18]
           20 GACGAGCCCGGGCGGAGGCT
                                                                  spacer_24
## [19]
           20 CTCGTCGATACGCTCTCGCT
                                                                  spacer_13
```

```
## [20]
           20 CAGTCGCCCCACAAGCATCT
                                                                   spacer_95
protospacers(guideSet)
## DNAStringSet object of length 20:
##
        width seq
                                                                   names
##
    [1]
           20 CCGAGTTGCTGCGCTGCTGC
                                                                   spacer_107
##
    [2]
           20 GCTCTGCTGGTTCTGCACGA
                                                                   spacer_9
##
    [3]
           20 CGGCCGCCGCGTCAGCACCA
                                                                   spacer_74
##
   [4]
           20 GCCCTTGCCGAGGGCGGAGG
                                                                   spacer_112
##
    [5]
           20 GGCCCCGCTGGGGCTGCTCC
                                                                   spacer_76
##
## [16]
           20 TCCCCCTCCGCCCTCGGCAA
                                                                   spacer_121
## [17]
           20 CGGCAGCGGGGCCGATGACG
                                                                   spacer_34
##
   [18]
           20 GACGAGCCCGGGCGGAGGCT
                                                                   spacer_24
## [19]
           20 CTCGTCGATACGCTCTCGCT
                                                                   spacer_13
## [20]
           20 CAGTCGCCCCACAAGCATCT
                                                                   spacer_95
pams(guideSet)
## DNAStringSet object of length 20:
##
        width seq
                                                                   names
            3 CGG
##
    [1]
                                                                   spacer_107
    [2]
            3 TGG
##
                                                                   spacer_9
##
    [3]
            3 CGG
                                                                   spacer_74
                                                                   spacer_112
##
    [4]
            3 GGG
##
    [5]
            3 AGG
                                                                   spacer_76
##
   . . .
   [16]
            3 GGG
                                                                   spacer_121
##
##
  [17]
            3 GGG
                                                                   spacer_34
            3 GGG
  Г18]
                                                                   spacer 24
            3 GGG
  [19]
                                                                   spacer_13
##
## [20]
            3 GGG
                                                                   spacer_95
head(pamSites(guideSet))
## spacer_107
                 spacer 9
                           spacer_74 spacer_112
                                                   spacer_76
                                                               spacer_55
##
        67371
                    66943
                                67233
                                           67396
                                                       67244
                                                                   67153
head(cutSites(guideSet))
## spacer_107
                 spacer_9
                           spacer_74 spacer_112 spacer_76
                                                               spacer_55
        67368
                                67230
                                           67399
                    66946
                                                       67247
                                                                   67156
```

The genomic locations stored in the IRanges represent the PAM site locations in the reference genome.

#### 4.4 Sequence features characterization

There are specific spacer sequence features, independent of the genomic context of the protospacer sequence, that can reduce or even eliminate gRNA activity:

- **Poly-T stretches**: four or more consecutive T nucleotides in the spacer sequence may act as a transcriptional termination signal for the U6 promoter.
- **Self-complementarity**: complementary sites with the gRNA backbone can compete with the targeted genomic sequence.
- Percent GC: gRNAs with GC content between 20% and 80% are preferred.

Use the function addSequenceFeatures to adds these spacer sequence characteristics to the GuideSet object:

guideSet <- addSequenceFeatures(guideSet)
head(guideSet)</pre>

```
GuideSet object with 6 ranges and 11 metadata columns:
##
                 segnames
                              ranges strand |
                                                         protospacer
                                                                                 pam
##
                    <Rle> <IRanges>
                                      <Rle> |
                                                      <DNAStringSet> <DNAStringSet>
##
     spacer_107
                    chr12
                               67371
                                           + |
                                              CCGAGTTGCTGCGCTGCTGC
                                                                                 CGG
                                                                                 TGG
##
       spacer_9
                    chr12
                               66943
                                              GCTCTGCTGGTTCTGCACGA
##
      spacer 74
                               67233
                                           + | CGGCCGCCGCGTCAGCACCA
                                                                                 CGG
                    chr12
##
     spacer 112
                    chr12
                               67396
                                            | GCCCTTGCCGAGGGCGGAGG
                                                                                 GGG
##
      spacer_76
                    chr12
                               67244
                                           - | GGCCCCGCTGGGGCTGCTCC
                                                                                 AGG
##
      spacer 55
                               67153
                                           - | CTGGTCCTGGAGAGGTTCCT
                                                                                 GGG
                    chr12
##
                                            region percentGC
                  pam_site cut_site
                                                                             polyC
                                                                  polyA
##
                           <numeric> <character> <numeric> <logical> <logical>
                 <numeric>
##
     spacer_107
                     67371
                                                           70
                                                                  FALSE
                                                                             FALSE
                                67368
                                          region_1
##
       spacer 9
                     66943
                                66946
                                          region_1
                                                           60
                                                                  FALSE
                                                                             FALSE
##
      spacer_74
                     67233
                                67230
                                          region_1
                                                           80
                                                                  FALSE
                                                                             FALSE
     spacer_112
                                          region_1
##
                     67396
                                67399
                                                           80
                                                                  FALSE
                                                                             FALSE
##
      spacer_76
                     67244
                                67247
                                          region_1
                                                           85
                                                                  FALSE
                                                                              TRUE
##
      spacer_55
                     67153
                                67156
                                          region_1
                                                           60
                                                                  FALSE
                                                                             FALSE
##
                     polyG
                                polyT startingGGGGG
##
                 <logical> <logical>
                                           <logical>
##
     spacer_107
                     FALSE
                                               FALSE
                                FALSE
##
       spacer_9
                     FALSE
                                FALSE
                                               FALSE
      spacer_74
##
                     FALSE
                                FALSE
                                               FALSE
##
     spacer 112
                     FALSE
                                FALSE
                                               FALSE
##
      spacer 76
                      TRUE
                                FALSE
                                               FALSE
##
                     FALSE
                                FALSE
                                               FALSE
      spacer_55
##
##
     seqinfo: 640 sequences (1 circular) from hg38 genome
##
     crisprNuclease: SpCas9
```

#### 4.5 Off-target search

In order to select gRNAs that are most specific to our target of interest, it is important to avoid gRNAs that target additional loci in the genome with either perfect sequence complementarity (multiple on-targets), or imperfect complementarity through tolerated mismatches (off-targets).

For instance, both the SpCas9 and AsCas12a nucleases can be tolerant to mismatches between the gRNA spacer sequence (RNA) and the protospacer sequence (DNA), thereby making it critical to characterize off-targets to minimize the introduction of double-stranded breaks (DSBs) beyond our intended target.

The addSpacerAlignments function appends a list of putative on- and off-targets to a GuideSet object using one of three methods. The first method uses the fast aligner bowtie (Langmead et al. 2009) via the crisprBowtie package to map spacer sequences to a specified reference genome. This can be done by specifying aligner="bowtie" in addSpacerAlignments.

The second method uses the fast aligner BWA via the crisprBwa package to map spacer sequences to a specified reference genome. This can be done by specifying aligner="bwa" in addSpacerAlignments. Note that this is not available for Windows machines.

The third method uses the package Biostrings to search for similar sequences in a set of DNA coordinates sequences, usually provided through a BSGenome object. This can be done by specifying aligner="biostrings" in addSpacerAlignments. This is extremely slow, but can be useful when searching for off-targets in custom short DNA sequences.

We can control the alignment parameters and output using several function arguments. n\_mismatches sets the maximum number of permitted gRNA:DNA mismatches (up to 3 mismatches). n\_max\_alignments specifies the maximum number of alignments for a given gRNA spacer sequence (1000 by default). The n\_max\_alignments parameter may be overruled by setting all\_Possible\_alignments=TRUE, which returns all possible alignments. canonical=TRUE filters out protospacer sequences that do not have a canonical PAM sequence.

Finally, the txObject argument in addSpacerAlignmentsused allows users to provide a TxDb object, or a TxDb object converted in a GRangesList using the TxDb2GRangesList function, to annotate genomic alignments with a gene model annotation. This is useful to understand whether or not off-targets are located in the CDS of another gene, for instance.

For the sake of time here, we will search only for on- and off-targets located in the beginning of human chr12 where IQSEC3 is located. We note note that users should always perform a genome-wide search as shown in the [CRISPRko design tutorial](https://github.com/crisprVerse/Tutorials/tree/master/Design\_CRISPRko\_Cas9].

We will use the bowtie method, with a maximum of 1 mismatch. First, we need to build a bowtie index sequence using the fasta file provided in crisprDesign. We use the RBowtie package to build the index:

For genome-wide off-target search, users will need to create a bowtie index on the whole genome. This is explained in this tutorial.

Finally, we also need to specify a BSgenome object storing DNA sequences of the human reference genome:

```
library(BSgenome.Hsapiens.UCSC.hg38)
bsgenome <- BSgenome.Hsapiens.UCSC.hg38
```

We are now ready to search for on- and off-targets:

## Loading required namespace: crisprBwa

Let's look at what was added to the GuideSet:

## guideSet

## GuideSet object with 20 ranges and 16 metadata columns:

```
##
                segnames
                             ranges strand |
                                                       protospacer
                                                                               pam
                   <Rle> <IRanges> <Rle> |
##
                                                    <DNAStringSet> <DNAStringSet>
##
     spacer_107
                   chr12
                              67371
                                         + | CCGAGTTGCTGCGCTGCTGC
                                                                               CGG
##
                              66943
                                         - | GCTCTGCTGGTTCTGCACGA
                                                                               TGG
       spacer_9
                   chr12
##
      spacer_74
                   chr12
                              67233
                                         + | CGGCCGCCGCGTCAGCACCA
                                                                               CGG
                                          - | GCCCTTGCCGAGGGCGGAGG
                                                                               GGG
##
     spacer_112
                   chr12
                              67396
##
      spacer_76
                              67244
                                         - | GGCCCCGCTGGGGCTGCTCC
                                                                               AGG
                   chr12
```

```
##
                       . . .
                                  . . .
                                          . . . .
                                                                                   . . .
##
                               67413
                                           + | TCCCCCTCCGCCAA
     spacer_121
                                                                                   GGG
                     chr12
##
      spacer 34
                     chr12
                                67093
                                            - | CGGCAGCGGGGCCGATGACG
                                                                                   GGG
                               67069
##
      spacer_24
                                            - | GACGAGCCCGGGCGGAGGCT
                                                                                   GGG
                     chr12
##
      spacer_13
                     chr12
                                66976
                                            - | CTCGTCGATACGCTCTCGCT
                                                                                   GGG
##
      spacer 95
                                            + | CAGTCGCCCCACAAGCATCT
                                                                                   GGG
                     chr12
                                67308
                                                                               polyC
##
                  pam_site cut_site
                                             region percentGC
                                                                    polyA
##
                  <numeric> <numeric> <character> <numeric> <logical> <logical>
##
     spacer_107
                     67371
                                 67368
                                          region 1
                                                            70
                                                                    FALSE
                                                                               FALSE
##
       spacer_9
                     66943
                                 66946
                                          region_1
                                                            60
                                                                    FALSE
                                                                               FALSE
##
      spacer_74
                     67233
                                 67230
                                          region_1
                                                            80
                                                                    FALSE
                                                                               FALSE
##
     spacer_112
                     67396
                                                                    FALSE
                                                                               FALSE
                                 67399
                                          region_1
                                                            80
##
      spacer_76
                     67244
                                 67247
                                          region_1
                                                            85
                                                                    FALSE
                                                                                TRUE
##
             . . .
                                   . . .
                                                . . .
                                                           . . .
                                                                      . . .
                                                                                 . . .
##
                     67413
                                 67410
                                                            75
                                                                    FALSE
                                                                                TRUE
     spacer_121
                                          region_1
##
      spacer_34
                     67093
                                 67096
                                          region_1
                                                            80
                                                                    FALSE
                                                                               FALSE
##
      spacer_24
                     67069
                                 67072
                                          region_1
                                                            80
                                                                    FALSE
                                                                               FALSE
      spacer_13
                                          region_1
##
                     66976
                                 66979
                                                            60
                                                                    FALSE
                                                                               FALSE
##
                                                                    FALSE
                                                                                TRUE
      spacer_95
                     67308
                                 67305
                                          region_1
                                                            60
##
                     polyG
                                 polyT startingGGGGG
                                                              n0
                                                                         n1
                                                                                  n0 c
##
                 <logical> <logical>
                                            <logical> <numeric> <numeric>
                                                                             <numeric>
##
                     FALSE
                                 FALSE
                                                FALSE
                                                               1
                                                                          0
     spacer_107
##
                                                                          0
       spacer 9
                     FALSE
                                 FALSE
                                                FALSE
                                                               1
                                                                                     1
                                 FALSE
                                                FALSE
                                                                          0
##
      spacer_74
                     FALSE
                                                               1
                                                                                     1
                                                                          0
##
     spacer_112
                     FALSE
                                 FALSE
                                                FALSE
                                                               1
                                                                                     1
##
      spacer_76
                       TRUE
                                 FALSE
                                                FALSE
                                                               1
                                                                          0
                                                                                     1
##
                        . . .
                                   . . .
                                                  . . .
                                                             . . .
##
     spacer_121
                     FALSE
                                 FALSE
                                                FALSE
                                                               1
                                                                          0
                                                                                     1
##
                       TRUE
                                                FALSE
                                                                          0
      spacer_34
                                 FALSE
                                                               1
                                                                                     1
##
      spacer_24
                     FALSE
                                 FALSE
                                                FALSE
                                                               1
                                                                          0
                                                                                     1
##
      spacer_13
                     FALSE
                                 FALSE
                                                FALSE
                                                               1
                                                                          0
                                                                                     1
##
      spacer_95
                     FALSE
                                 FALSE
                                                FALSE
                                                               1
                                                                           0
                                                                                     1
                      n1_c
##
                                alignments
##
                 <numeric> <GRangesList>
##
     spacer_107
                          0 chr12:67371:+
##
                          0 chr12:66943:-
       spacer_9
##
      spacer 74
                          0 chr12:67233:+
##
     spacer_112
                          0 chr12:67396:-
##
                          0 chr12:67244:-
      spacer_76
##
##
     spacer 121
                          0 chr12:67413:+
##
      spacer 34
                          0 chr12:67093:-
##
      spacer_24
                          0 chr12:67069:-
##
      spacer_13
                          0 chr12:66976:-
##
      spacer_95
                          0 chr12:67308:+
##
##
     seqinfo: 640 sequences (1 circular) from hg38 genome
##
     crisprNuclease: SpCas9
```

A few columns were added to the GuideSet object to summarize the number of on- and off-targets for each spacer sequence, taking into account genomic context:

- n0, n1, n2, n3: specify number of alignments with 0, 1, 2 and 3 mismatches, respectively.
- n0\_c, n1\_c, n2\_c, n3\_c: specify number of alignments in a coding region, with 0, 1, 2 and 3 mismatches, respectively.

• n0\_p, n1\_p, n2\_p, n3\_p: specify number of alignments in a promoter region of a coding gene, with 0, 1, 2 and 3 mismatches, respectively.

To look at the individual on- and off-targets and their context, use the alignments function to retrieve a table of all genomic alignments stored in the GuideSet object:

## alignments(guideSet)

##	GRanges object	ct with 20 ra	anges and	14 me	tadata co	lumns:		
##		seqnames	ranges s				acer	
##		<rle> <i< th=""><th>Ranges&gt;</th><th><rle></rle></th><th>  &lt;</th><th>DNAString</th><th>Set&gt;</th><th></th></i<></rle>	Ranges>	<rle></rle>	<	DNAString	Set>	
##	spacer_107	chr12	67371	+	CCGAGTT	CCTCCCCTC	CTGC	
##	spacer_9	chr12	66943	_	GCTCTGC	TGGTTCTGC	ACGA	
##	spacer_74	chr12	67233	+	CGGCCGC	CGCGTCAGC	ACCA	
##	spacer_112	chr12	67396	_	GCCCTTC	CCGAGGGCG	GAGG	
##	spacer_76	chr12	67244	_	GGCCCCC	CTGGGGCTG	CTCC	
##								
##	spacer_121	chr12	67413	+	TCCCCCT	CCGCCCTCG	GCAA	
##	spacer_34	chr12	67093	_	CGGCAGC	GGGGCCGAT	GACG	
##	spacer_24	chr12	67069	_	GACGAGO	CCGGGCGGA	GGCT	
##	spacer_13	chr12	66976	_	CTCGTCG	SATACGCTCT	CGCT	
##	spacer_95	chr12	67308	+	CAGTCGC	CCCACAAGC	ATCT	
##		pro	otospacer		pan	n pam_sit	e n_mismatche	98
##		<dnas< th=""><th>tringSet&gt;</th><th><dnas< th=""><th>tringSet&gt;</th><th><pre><numeric< pre=""></numeric<></pre></th><th>&gt; <integer< th=""><th>_&gt;</th></integer<></th></dnas<></th></dnas<>	tringSet>	<dnas< th=""><th>tringSet&gt;</th><th><pre><numeric< pre=""></numeric<></pre></th><th>&gt; <integer< th=""><th>_&gt;</th></integer<></th></dnas<>	tringSet>	<pre><numeric< pre=""></numeric<></pre>	> <integer< th=""><th>_&gt;</th></integer<>	_>
##	spacer_107	CCGAGTTGCTG	CGCTGCTGC		CGC	6737	1	0
##	spacer_9	GCTCTGCTGGT	TCTGCACGA		TGG	6694	3	0
##	spacer_74	CGGCCGCGCG'	TCAGCACCA		CGC	6723	3	0
##	spacer_112	GCCCTTGCCGA	GGGCGGAGG		GGG	6739	6	0
##	spacer_76	GGCCCCGCTGG	GGCTGCTCC		AGG	6724	4	0
##								
##	-	TCCCCCTCCGC			GGC	6741	3	0
##	-	CGGCAGCGGGG			GGC			0
##	-	GACGAGCCCGG			GGC			0
##	-	CTCGTCGATAC			GGC		6	0
##	spacer_95	CAGTCGCCCCA	CAAGCATCT		GGC		8	0
##			cut_site		cds	fiveUTRs	threeUTRs	
##		<logical> &lt;</logical>						
##	spacer_107	TRUE	67368		QSEC3	<na></na>	<na></na>	
##	spacer_9	TRUE	66946		QSEC3	<na></na>	<na></na>	
##	spacer_74	TRUE	67230		QSEC3	<na></na>	<na></na>	
##	spacer_112	TRUE	67399		QSEC3	<na></na>	<na></na>	
##	spacer_76	TRUE	67247	10	QSEC3	<na></na>	<na></na>	
##	• • •				•••		• • •	
##	spacer_121	TRUE	67410		QSEC3	<na></na>	<na></na>	
##	spacer_34	TRUE	67096		QSEC3	<na></na>	<na></na>	
##	spacer_24	TRUE	67072		QSEC3	<na></na>	<na></na>	
##	spacer_13	TRUE	66979		QSEC3	<na></na>	<na></na>	
##	spacer_95	TRUE	67305		QSEC3	<na></na>	<na></na>	
##		exons	intr		_	•	ic_distance	
##	100	<pre><character></character></pre>					<integer></integer>	
##	spacer_107	IQSEC3		NA>	<na></na>		<na></na>	
##	spacer_9	IQSEC3		NA>	<na></na>		<na></na>	
##	spacer_74	IQSEC3		NA>	<na></na>		<na></na>	
##	spacer_112	IQSEC3		NA>	<na></na>		<na></na>	
##	spacer_76	IQSEC3	<	NA>	<na></na>	•	<na></na>	

```
##
                            . . .
                                          . . .
                                                         . . .
                                                                                 . . .
                        IQSEC3
##
     spacer_121
                                         < NA >
                                                       < NA >
                                                                               < NA >
       spacer 34
##
                        IQSEC3
                                         <NA>
                                                        <NA>
                                                                               <NA>
       spacer_24
                        IQSEC3
                                                                               <NA>
##
                                         <NA>
                                                       <NA>
##
       spacer_13
                         IQSEC3
                                         <NA>
                                                        <NA>
                                                                               <NA>
       spacer 95
                        IQSEC3
                                         <NA>
                                                                               <NA>
##
                                                       <NA>
##
##
     seqinfo: 25 sequences (1 circular) from hg38 genome
```

The functions on Targets and off Targets will return on-target alignments (no mismatch) and off-target alignment (with at least one mismatch), respectively. See ?addSpacerAlignments for more details about the different options.

#### 4.5.1 Iterative spacer alignments

gRNAs that align to hundreds of different locations are highly unspecific and undesirable. This can also cause addSpacerAlignments to be slow. To mitigate this, we provide addSpacerAlignmentsIterative, an iterative version of addSpacerAlignments that curtails alignment searches for gRNAs having more hits than the user-defined threshold (see ?addSpacerAlignmentsIterative).

#### 4.5.2 Faster alignment by removing repeat elements

To remove protospacer sequences located in repeats or low-complexity DNA sequences (regions identified by RepeatMasker), which are usually not of interest due to their low specificity, we provide the convenience function removeRepeats:

## 4.6 Off-target scoring

After retrieving a list of putative off-targets and on-targets for a given spacer sequence, we can use addOffTargetScores to predict the likelihood of the nuclease to cut at the off-targets based on mismatch tolerance. Currently, only off-target scoring for the SpCas9 nuclease are available (MIT and CFD algorithms):

```
guideSet <- addOffTargetScores(guideSet)
guideSet</pre>
```

```
## GuideSet object with 17 ranges and 19 metadata columns:
```

##		seqnames	ranges	strand		prote	ospacer	pam
##		<rle></rle>	<iranges></iranges>	<rle>  </rle>		<dnastr< th=""><th>ingSet&gt; ·</th><th><pre><dnastringset></dnastringset></pre></th></dnastr<>	ingSet> ·	<pre><dnastringset></dnastringset></pre>
##	spacer_107	chr12	67371	+	CCGA	AGTTGCTGCG	CTGCTGC	CGG
##	spacer_9	chr12	66943	-	GCTC	CTGCTGGTTC	TGCACGA	TGG
##	spacer_74	chr12	67233	+	CGGC	CCGCCGCGTC	AGCACCA	CGG
##	spacer_112	chr12	67396	-	GCCC	CTTGCCGAGG	GCGGAGG	GGG
##	spacer_76	chr12	67244	-	GGCC	CCCGCTGGGG	CTGCTCC	AGG
##								
##	spacer_71	chr12	67218	-	TGTC	CCGTGGTGCT	GACGCGG	CGG
##	spacer_121	chr12	67413	+	TCCC	CCCTCCGCCC	TCGGCAA	GGG
##	spacer_24	chr12	67069	-	GACC	GAGCCCGGGC	GGAGGCT	GGG
##	spacer_13	chr12	66976	-	CTCC	GTCGATACGC	TCTCGCT	GGG
##	spacer_95	chr12	67308	+	CAGT	CGCCCCACA.	AGCATCT	GGG
##		pam_site	cut_site	e re	gion	${\tt percentGC}$	pol	yA polyC
##		<numeric></numeric>	<numeric></numeric>	<pre><charac< pre=""></charac<></pre>	ter>	<numeric></numeric>	<logical< td=""><td>l&gt; <logical></logical></td></logical<>	l> <logical></logical>
##	spacer_107	67371	67368	3 regi	on_1	70	FAL	SE FALSE

```
##
       spacer 9
                      66943
                                 66946
                                           region_1
                                                              60
                                                                      FALSE
                                                                                 FALSE
##
                                                              80
                                                                      FALSE
                                                                                FALSE
      spacer_74
                      67233
                                 67230
                                           region_1
                                                                                FALSE
##
     spacer 112
                      67396
                                 67399
                                           region 1
                                                              80
                                                                      FALSE
##
      spacer_76
                      67244
                                 67247
                                                             85
                                                                      FALSE
                                                                                  TRUE
                                           region_1
##
                                                                                   . . .
             . . .
                        . . .
                                    . . .
                                                                        . . .
                                                             . . .
##
      spacer 71
                                 67221
                                                             70
                                                                      FALSE
                                                                                 FALSE
                      67218
                                           region 1
##
     spacer 121
                      67413
                                 67410
                                           region 1
                                                              75
                                                                     FALSE
                                                                                  TRUE
##
      spacer_24
                      67069
                                 67072
                                           region_1
                                                              80
                                                                      FALSE
                                                                                 FALSE
##
      spacer_13
                      66976
                                 66979
                                           region_1
                                                              60
                                                                      FALSE
                                                                                 FALSE
                                                                                  TRUE
##
      spacer_95
                      67308
                                 67305
                                           region_1
                                                              60
                                                                      FALSE
##
                      polyG
                                 polyT startingGGGGG
                                                                n0
                                                                                    n0_c
                                                                           n1
##
                  <logical> <logical>
                                             <logical> <numeric>
                                                                   <numeric>
                                                                              <numeric>
##
     spacer_107
                      FALSE
                                 FALSE
                                                 FALSE
                                                                 1
                                                                            0
                                                 FALSE
                                                                            0
##
       spacer_9
                      FALSE
                                 FALSE
                                                                 1
                                                                                       1
##
      spacer_74
                                                 FALSE
                                                                            0
                      FALSE
                                 FALSE
                                                                 1
                                                                                       1
##
     spacer_112
                      FALSE
                                 FALSE
                                                 FALSE
                                                                 1
                                                                            0
                                                                                       1
                                                                            0
##
      spacer_76
                       TRUE
                                 FALSE
                                                 FALSE
                                                                 1
                                                                                       1
##
                         . . .
                                    . . .
                                                   . . .
                                                               . . .
                                                                          . . .
##
      spacer_71
                                                                            0
                      FALSE
                                 FALSE
                                                 FALSE
                                                                 1
                                                                                       1
##
     spacer 121
                      FALSE
                                 FALSE
                                                 FALSE
                                                                 1
                                                                            0
                                                                                       1
##
      spacer_24
                      FALSE
                                 FALSE
                                                 FALSE
                                                                 1
                                                                            0
                                                                                       1
##
      spacer 13
                      FALSE
                                 FALSE
                                                 FALSE
                                                                 1
                                                                            0
                                                                                       1
##
      spacer_95
                      FALSE
                                 FALSE
                                                 FALSE
                                                                            0
                                                                 1
                                                                                       1
##
                       n1 c
                                alignments inRepeats score cfd score mit
##
                  <numeric> <GRangesList> <logical> <numeric> <numeric>
##
     spacer_107
                           0 chr12:67371:+
                                                 FALSE
                                                                 1
                                                                            1
##
       spacer_9
                           0 chr12:66943:-
                                                 FALSE
                                                                 1
                                                                            1
##
      spacer_74
                           0 chr12:67233:+
                                                 FALSE
                                                                 1
                                                                            1
##
     spacer_112
                                                 FALSE
                                                                 1
                           0 chr12:67396:-
                                                                            1
##
      spacer_76
                           0 chr12:67244:-
                                                 FALSE
                                                                 1
                                                                            1
##
                                                   . . .
                                                               . . .
##
      spacer_71
                          0 chr12:67218:-
                                                 FALSE
                                                                 1
                                                                            1
##
     spacer_121
                           0 chr12:67413:+
                                                 FALSE
                                                                 1
                                                                            1
##
      spacer_24
                           0 chr12:67069:-
                                                 FALSE
                                                                 1
                                                                            1
##
      spacer 13
                           0 chr12:66976:-
                                                 FALSE
                                                                 1
                                                                            1
##
      spacer 95
                           0 chr12:67308:+
                                                 FALSE
                                                                            1
                                                                 1
##
##
     seqinfo: 640 sequences (1 circular) from hg38 genome
     crisprNuclease: SpCas9
```

Note that this will only work after calling addSpacerAlignments, as it requires a list of off-targets for each gRNA entry.

#### 4.7 On-target scoring

<Rle> <IRanges>

##

addOnTargetScores adds scores from all on-target efficiency algorithms available in the R package crisprScore and appends them to the GuideSet. By default, scores for all available methods for a given nuclease will be computed. Here, for the sake of time, let's add only the CRISPRater score:

```
guideSet <- addOnTargetScores(guideSet, methods="crisprater")
head(guideSet)

## GuideSet object with 6 ranges and 20 metadata columns:
## seqnames ranges strand | protospacer pam</pre>
```

<DNAStringSet> <DNAStringSet>

<Rle> |

```
##
     spacer 107
                    chr12
                               67371
                                           + | CCGAGTTGCTGCGCTGCTGC
                                                                                  CGG
                                             | GCTCTGCTGGTTCTGCACGA
##
                                                                                  TGG
       spacer_9
                    chr12
                               66943
##
      spacer 74
                    chr12
                               67233
                                             | CGGCCGCCGCGTCAGCACCA
                                                                                  CGG
     spacer_112
                                                                                  GGG
##
                               67396
                                           - | GCCCTTGCCGAGGGCGGAGG
                    chr12
##
      spacer_76
                    chr12
                               67244
                                             | GGCCCCGCTGGGGCTGCTCC
                                                                                  AGG
      spacer 55
                                           - | CTGGTCCTGGAGAGGTTCCT
##
                    chr12
                               67153
                                                                                  GGG
##
                  pam site cut site
                                            region percentGC
                                                                              polyC
                                                                   polyA
##
                 <numeric> <numeric> <character> <numeric> <logical> <logical>
##
     spacer_107
                     67371
                                67368
                                          region 1
                                                           70
                                                                   FALSE
                                                                              FALSE
##
       spacer_9
                     66943
                                66946
                                          region_1
                                                           60
                                                                   FALSE
                                                                              FALSE
##
      spacer_74
                     67233
                                67230
                                          region_1
                                                           80
                                                                   FALSE
                                                                              FALSE
##
     spacer_112
                     67396
                                                           80
                                                                   FALSE
                                                                              FALSE
                                67399
                                          region_1
##
      spacer_76
                     67244
                                67247
                                          region_1
                                                           85
                                                                   FALSE
                                                                               TRUE
                     67153
##
      spacer_55
                                67156
                                          region_1
                                                           60
                                                                   FALSE
                                                                              FALSE
##
                                polyT startingGGGGG
                                                                                 n0_c
                     polyG
                                                             n0
                                                                        n1
##
                 <logical> <logical>
                                           <logical> <numeric>
                                                                            <numeric>
                                                                 <numeric>
##
     spacer_107
                     FALSE
                                FALSE
                                               FALSE
                                                                         0
                                                               1
                                                                                    1
##
       spacer 9
                     FALSE
                                FALSE
                                               FALSE
                                                               1
                                                                          0
                                                                                    1
##
      spacer 74
                     FALSE
                                FALSE
                                               FALSE
                                                                         0
                                                                                    1
                                                               1
                                                                         0
##
     spacer 112
                     FALSE
                                FALSE
                                               FALSE
                                                               1
                                                                                    1
##
      spacer_76
                      TRUE
                                FALSE
                                               FALSE
                                                               1
                                                                         0
                                                                                    1
##
      spacer 55
                     FALSE
                                FALSE
                                               FALSE
                                                               1
                                                                          0
                                                                                    1
##
                      n1_c
                               alignments inRepeats score_cfd score_mit
                 <numeric> <GRangesList> <logical> <numeric> <numeric>
##
##
     spacer 107
                          0 chr12:67371:+
                                               FALSE
                                                               1
##
       spacer 9
                          0 chr12:66943:-
                                               FALSE
                                                               1
                                                                         1
##
      spacer_74
                          0 chr12:67233:+
                                               FALSE
                                                               1
                                                                          1
     spacer_112
                          0 chr12:67396:-
                                                               1
##
                                               FALSE
                                                                          1
##
      spacer_76
                          0 chr12:67244:-
                                               FALSE
                                                               1
                                                                          1
##
      spacer_55
                          0 chr12:67153:-
                                               FALSE
                                                               1
                                                                          1
##
                 score_crisprater
##
                         <numeric>
##
     spacer_107
                          0.782780
##
       spacer_9
                          0.834319
##
      spacer 74
                          0.764870
##
     spacer 112
                          0.795745
##
      spacer 76
                          0.755493
##
      spacer_55
                          0.711902
##
##
     seqinfo: 640 sequences (1 circular) from hg38 genome
     crisprNuclease: SpCas9
```

See the crisprScore vignette for a full description of the different scores.

## 4.8 Restriction enzymes

Restriction enzymes are usually involved in the gRNA library synthesis process. Removing gRNAs that contain specific restriction sites is often necessary. We provide the function addRestrictionEnzymes to indicate whether or not gRNAs contain restriction sites for a user-defined set of enzymes:

```
guideSet <- addRestrictionEnzymes(guideSet)</pre>
```

When no enzymes are specified, the function adds annotation for the following default enzymes: EcoRI, KpnI, BsmBI, BsaI, BbsI, PacI, ISceI and MluI. The function also has two additional arguments, flanking5 and flanking3, to specify nucleotide sequences flanking the spacer sequence (5' and 3', respectively) in the

lentiviral cassette that will be used for gRNA delivery. The function will effectively search for restriction sites in the full sequence [flanking5] [spacer] [flanking3].

The enzymeAnnotation function can be used to retrieve the added annotation:

head(enzymeAnnotation(guideSet))

```
## DataFrame with 6 rows and 7 columns
##
                   EcoRI
                               KpnI
                                         BsmBI
                                                     BsaI
                                                                BbsT
                                                                           PacI
##
               <logical> <logical> <logical> <logical> <logical> <logical> <logical>
                   FALSE
                              FALSE
## spacer_107
                                         FALSE
                                                    FALSE
                                                               FALSE
                                                                          FALSE
## spacer 9
                   FALSE
                              FALSE
                                         FALSE
                                                    FALSE
                                                               FALSE
                                                                          FALSE
## spacer_74
                   FALSE
                              FALSE
                                         FALSE
                                                    FALSE
                                                               FALSE
                                                                         FALSE
## spacer_112
                   FALSE
                              FALSE
                                         FALSE
                                                    FALSE
                                                               FALSE
                                                                         FALSE
## spacer_76
                   FALSE
                              FALSE
                                         FALSE
                                                    FALSE
                                                               FALSE
                                                                          FALSE
## spacer_55
                   FALSE
                              FALSE
                                         FALSE
                                                    FALSE
                                                               FALSE
                                                                          FALSE
##
                    MluI
##
               <logical>
## spacer_107
                   FALSE
## spacer_9
                   FALSE
## spacer_74
                   FALSE
## spacer_112
                   FALSE
## spacer_76
                   FALSE
## spacer_55
                   FALSE
```

#### 4.9 Gene annotation

The function addGeneAnnotation adds transcript- and gene-level contextual information to gRNAs from a TxDb-like object:

The gene annotation can be retrieved using the function geneAnnotation:

geneAnnotation(guideSet)

```
## DataFrame with 17 rows and 23 columns
##
                    chr anchor_site
                                      strand gene_symbol
                                                                   gene_id
##
                          <integer> <factor> <character>
              <factor>
                                                               <character>
## spacer_107
                 chr12
                              67368
                                            +
                                                   IQSEC3 ENSG00000120645
## spacer_9
                 chr12
                              66946
                                                   IQSEC3 ENSG00000120645
## spacer_74
                              67230
                                            +
                                                   IQSEC3 ENSG00000120645
                 chr12
## spacer_112
                  chr12
                              67399
                                                   IQSEC3 ENSG00000120645
                                                   IQSEC3 ENSG00000120645
## spacer_76
                  chr12
                              67247
                    . . .
## spacer_71
                 chr12
                              67221
                                                   IQSEC3 ENSG00000120645
## spacer 121
                 chr12
                              67410
                                                   IQSEC3 ENSG00000120645
## spacer_24
                 chr12
                              67072
                                                   IQSEC3 ENSG00000120645
## spacer 13
                 chr12
                              66979
                                                   IQSEC3 ENSG00000120645
                              67305
                                                   IQSEC3 ENSG00000120645
## spacer_95
                  chr12
                                    protein_id
                                                  cut_cds cut_fiveUTRs cut_threeUTRs
##
                         tx id
##
                                    <character> <logical>
                                                              <logical>
                                                                             <logical>
                   <character>
## spacer 107 ENST00000538872 ENSP00000437554
                                                     TRUE
                                                                  FALSE
                                                                                FALSE
## spacer 9
              ENST00000538872 ENSP00000437554
                                                     TRUE
                                                                  FALSE
                                                                                FALSE
## spacer_74
              ENST00000538872 ENSP00000437554
                                                     TRUE
                                                                  FALSE
                                                                                FALSE
## spacer_112 ENST00000538872 ENSP00000437554
                                                     TRUE
                                                                  FALSE
                                                                                FALSE
```

	-	ENST00000538872	ENSP0000	0437554		FALSE	FALSE	
	71		ENGDOOO	0407554	···			
		ENST00000538872				FALSE	FALSE	
	-	ENST00000538872				FALSE	FALSE	
		ENST00000538872				FALSE	FALSE	
		ENST00000538872				FALSE	FALSE	
	spacer_95	ENST00000538872				FALSE	FALSE	
##		cut_introns per				_		
##		<logical> <n< td=""><td></td><td><numeric></numeric></td><td><numeric< td=""><td><pre>&gt; <numeric></numeric></pre></td><td></td></numeric<></td></n<></logical>		<numeric></numeric>	<numeric< td=""><td><pre>&gt; <numeric></numeric></pre></td><td></td></numeric<>	<pre>&gt; <numeric></numeric></pre>		
	spacer_107	FALSE	13.7	162		1 8.5		
	spacer_9	FALSE	1.8	22		0 2.5		
	spacer_74		9.8	116		0 6.5		
##	spacer_112	FALSE	14.6	173		1 8.9		
##	spacer_76	FALSE	10.3	122		0 6.8		
##								
##	spacer_71	FALSE	9.6	113		0 6.4		
##	spacer_121	FALSE	14.9	176		1 9.1		
##	spacer_24	FALSE	5.4	64		0 4.3		
##	spacer_13	FALSE	2.7	33		0 3.0		
##	spacer_95	FALSE	11.9	141		1 7.6		
##		nIsoforms total	Isoforms	percentIsoform	ns isCommonE	xon nCoding	Isoforms	
##		<integer> &lt;</integer>	numeric>	<numerio< td=""><td>c&gt; <logic< td=""><td>al&gt; &lt;</td><td>integer&gt;</td></logic<></td></numerio<>	c> <logic< td=""><td>al&gt; &lt;</td><td>integer&gt;</td></logic<>	al> <	integer>	
##	spacer_107	1	2	Ę	50 FA	LSE	1	
	spacer_9	1	2	Ę	50 FA	LSE	1	
	spacer_74	1	2	Ę		LSE	1	
	spacer_112	1	2	Ę		LSE	1	
	spacer_76	1	2			LSE	1	
	spacer_71	1	2			LSE	1	
	spacer_121	1	2			LSE	1	
	spacer_24	1	2			LSE	1	
	spacer_13	1	2			LSE	1	
	spacer_95	1	2			LSE	1	
##	phacer_co	=	_					
##		totalCodingIsoforms percentCodingIsoforms isCommonCodingExon <numeric> <numeric> <logical></logical></numeric></numeric>						
	spacer_107	VII ulii C	2	VII dillici	50	FALSE		
	spacer_9			50	FALSE			
	spacer_74			50	FALSE			
				50	FALSE			
	spacer_112			50	FALSE			
	spacer_76		2					
			2		· · ·	EALCE		
	spacer_71			50 FALSE				
	spacer_121			50 50	FALSE			
	spacer_24		2		50			
	spacer_13		2		50	FALSE		
##	spacer_95		2		50	FALSE		

It contains a lot of information that contextualizes the genomic location of the protospacer sequences.

The ID columns (tx\_id, gene\_id, protein\_id, exon\_id) give Ensembl IDs. The exon\_rank gives the order of the exon for the transcript, for example "2" indicates it is the second exon (from the 5' end) in the mature transcript.

The columns cut\_cds, cut\_fiveUTRs, cut\_threeUTRs and cut\_introns indicate whether the guide sequence overlaps with CDS, 5' UTR, 3' UTR, or an intron, respectively.

percentCDS gives the location of the cut\_site within the transcript as a percent from the 5' end to the 3' end. aminoAcidIndex gives the number of the specific amino acid in the protein where the cut is predicted to occur. downstreamATG shows how many in-frame ATGs are downstream of the cut\_site (and upstream from the defined percent transcript cutoff, met\_cutoff), indicating a potential alternative translation initiation site that may preserve protein function.

For more information about the other columns, type ?addGeneAnnotation.

#### 4.10 TSS annotation

Similarly, one might want to know which protospacer sequences are located within promoter regions of known genes:

```
## DataFrame with 10 rows and 11 columns
##
                   chr anchor_site
                                                        tx_id
                                                                       gene_id
##
                         <integer> <factor>
              <factor>
                                                  <character>
                                                                   <character>
## spacer_9
                 chr12
                             66946
                                           - ENST00000538872 ENSG00000120645
## spacer_74
                 chr12
                             67230
                                           + ENST00000538872 ENSG00000120645
## spacer 76
                                           - ENST00000538872 ENSG00000120645
                 chr12
                             67247
## spacer 55
                                           - ENST00000538872 ENSG00000120645
                 chr12
                             67156
## spacer 72
                 chr12
                             67224
                                           - ENST00000538872 ENSG00000120645
## spacer_54
                                           + ENST00000538872 ENSG00000120645
                 chr12
                             67145
## spacer_15
                 chr12
                             66995
                                           + ENST00000538872 ENSG00000120645
## spacer 71
                                           - ENST00000538872 ENSG00000120645
                 chr12
                             67221
                                           - ENST00000538872 ENSG00000120645
## spacer 24
                 chr12
                             67072
   spacer_13
                             66979
                                           - ENST00000538872 ENSG00000120645
##
                 chr12
             gene_symbol
                             promoter
                                            tss id
                                                    tss_strand
                                                                   tss_pos dist_to_tss
              <character>
                                       <character> <character> <integer>
##
                          <character>
                                                                             <numeric>
## spacer_9
                   IQSEC3
                                    P1
                                         IQSEC3_P1
                                                                     66767
                                                                                    179
## spacer_74
                   IQSEC3
                                    P1
                                         IQSEC3 P1
                                                                     66767
                                                                                    463
## spacer_76
                   IQSEC3
                                    P1
                                         IQSEC3 P1
                                                                     66767
                                                                                    480
## spacer_55
                   IQSEC3
                                    P1
                                         IQSEC3_P1
                                                                     66767
                                                                                    389
## spacer_72
                   IQSEC3
                                    Ρ1
                                         IQSEC3_P1
                                                                                    457
                                                                     66767
## spacer_54
                   IQSEC3
                                    P1
                                         IQSEC3_P1
                                                                     66767
                                                                                    378
## spacer_15
                   IQSEC3
                                    P1
                                         IQSEC3_P1
                                                                     66767
                                                                                    228
## spacer 71
                                         IQSEC3 P1
                   IQSEC3
                                    P1
                                                                     66767
                                                                                    454
## spacer 24
                   IQSEC3
                                    P1
                                         IQSEC3 P1
                                                                                    305
                                                                     66767
## spacer 13
                   IQSEC3
                                         IQSEC3 P1
                                                                     66767
                                                                                    212
```

For more information, type ?addTssAnnotation.

#### 4.11 SNP information

Common single-nucleotide polymorphisms (SNPs) can change the on-target and off-target properties of gRNAs by altering the binding. The function addSNPAnnotation annotates gRNAs with respect to a reference database of SNPs (stored in a VCF file), specified by the vcf argument.

VCF files for common SNPs (dbSNPs) can be downloaded from NCBI on the dbSNP website. We include in this package an example VCF file for common SNPs located in the proximity of human gene IQSEC3. This was obtained using the dbSNP151 RefSNP database obtained by subsetting around IQSEC.

## DataFrame with 0 rows and 9 columns

The rs\_site\_rel gives the relative position of the SNP with respect to the pam\_site. allele\_ref and allele\_minor report the nucleotide of the reference and minor alleles, respectively. MAF\_1000G and MAF\_TOPMED report the minor allele frequency (MAF) in the 1000Genomes and TOPMED populations.

## 4.12 Filtering and ranking gRNAs

Once gRNAs are fully annotated, it is easy to filter out any unwanted gRNAs since GuideSet objects can be subsetted like regular vectors in R.

As an example, suppose that we only want to keep gRNAs that have percent GC between 20% and 80% and that do not contain a polyT stretch. This can be achieved using the following lines:

```
guideSet <- guideSet[guideSet$percentGC>=20]
guideSet <- guideSet[guideSet$percentGC<=80]
guideSet <- guideSet[!guideSet$polyT]</pre>
```

Similarly, it is easy to rank gRNAs based on a set of criteria using the regular order function.

For instance, let's sort gRNAs by the CRISPRater on-target score:

```
# Creating an ordering index based on the CRISPRater score:
# Using the negative values to make sure higher scores are ranked first:
o <- order(-guideSet$score_crisprater)
# Ordering the GuideSet:
guideSet <- guideSet[o]
head(guideSet)</pre>
```

## GuideSet object with 6 ranges and 25 metadata columns:

```
##
                 segnames
                              ranges strand |
                                                        protospacer
                                                                                 pam
##
                    <Rle> <IRanges>
                                      <Rle> |
                                                     <DNAStringSet> <DNAStringSet>
                    chr12
##
       spacer_9
                               66943
                                              GCTCTGCTGGTTCTGCACGA
                                                                                 TGG
##
                               67396
                                           - | GCCCTTGCCGAGGGCGGAGG
                                                                                 GGG
     spacer_112
                    chr12
##
     spacer_107
                    chr12
                               67371
                                          + | CCGAGTTGCTGCGCTGCTGC
                                                                                 CGG
##
                                                                                 CGG
      spacer_74
                               67233
                                          + | CGGCCGCCGCGTCAGCACCA
                    chr12
##
      spacer 76
                    chr12
                               67244
                                           - | GGCCCCGCTGGGGCTGCTCC
                                                                                 AGG
##
     spacer_121
                    chr12
                               67413
                                          + | TCCCCTCCGCCTCGGCAA
                                                                                 GGG
##
                  pam site cut site
                                           region percentGC
                                                                  polyA
                                                                             polyC
##
                 <numeric> <numeric> <character> <numeric> <logical> <logical>
##
                     66943
                                                                  FALSE
       spacer 9
                                66946
                                         region 1
                                                          60
                                                                             FALSE
##
     spacer_112
                     67396
                                67399
                                         region_1
                                                          80
                                                                  FALSE
                                                                             FALSE
##
     spacer 107
                     67371
                                67368
                                         region_1
                                                          70
                                                                  FALSE
                                                                             FALSE
##
                                                           80
                                                                             FALSE
      spacer_74
                     67233
                                67230
                                         region_1
                                                                  FALSE
##
      spacer_76
                     67244
                                67247
                                         region_1
                                                           85
                                                                  FALSE
                                                                              TRUE
                                                                              TRUE
##
     spacer_121
                     67413
                                                           75
                                                                  FALSE
                                67410
                                         region_1
##
                     polyG
                                polyT startingGGGGG
                                                            n0
                                                                       n1
                                                                                n0_c
##
                 <logical> <logical>
                                           <logical> <numeric> <numeric>
                                                                           <numeric>
##
                     FALSE
                                FALSE
                                               FALSE
                                                              1
                                                                        0
       spacer_9
                                                                                   1
##
     spacer_112
                     FALSE
                                FALSE
                                               FALSE
                                                              1
                                                                        0
                                                                                   1
                     FALSE
##
     spacer_107
                                FALSE
                                               FALSE
                                                              1
                                                                        0
                                                                                   1
```

```
##
      spacer_74
                     FALSE
                                FALSE
                                                FALSE
                                                                          0
                                                               1
##
                       TRUE
                                FALSE.
                                                FALSE
                                                                          0
                                                                                     1
      spacer_76
                                                               1
##
     spacer 121
                     FALSE
                                FALSE
                                                FALSE
                                                               1
                                                                          0
                                                                                     1
##
                      n1_c
                               alignments inRepeats score_cfd score_mit
##
                 <numeric> <GRangesList> <logical> <numeric> <numeric>
##
                          0 chr12:66943:-
                                                FALSE
                                                               1
       spacer 9
                          0 chr12:67396:-
                                               FALSE
                                                               1
##
     spacer 112
                                                                          1
##
     spacer_107
                          0 chr12:67371:+
                                               FALSE
                                                               1
                                                                          1
##
      spacer_74
                          0 chr12:67233:+
                                               FALSE
                                                               1
                                                                          1
##
      spacer_76
                          0 chr12:67244:-
                                                FALSE
                                                               1
                                                                          1
##
     spacer_121
                          0 chr12:67413:+
                                                FALSE
                                                               1
                                                                          1
##
                 score_crisprater
                                         enzymeAnnotation
                                                                  geneAnnotation
##
                         <numeric>
                                    <SplitDataFrameList> <SplitDataFrameList>
                          0.834319 FALSE:FALSE:FALSE:...
##
       spacer_9
                                                               chr12:66946:-:...
##
     spacer_112
                          0.795745 FALSE:FALSE:FALSE:...
                                                               chr12:67399:-:...
##
     spacer_107
                          0.782780 FALSE:FALSE:FALSE:...
                                                               chr12:67368:+:...
##
                          0.764870 FALSE:FALSE:FALSE:...
      spacer_74
                                                               chr12:67230:+:...
##
      spacer 76
                          0.755493 FALSE:FALSE:FALSE:...
                                                               chr12:67247:-:...
##
                          0.741315 FALSE:FALSE:FALSE:...
                                                               chr12:67410:+:...
     spacer_121
##
                         tssAnnotation
                                           hasSNP
                                                                    snps
##
                 <SplitDataFrameList> <logical> <SplitDataFrameList>
##
                    chr12:66946:-:...
                                            FALSE
       spacer 9
##
                                            FALSE
     spacer_112
                              : . . . , . . .
                                                                : . . . , . . .
                              :...,...
                                            FALSE
##
     spacer 107
##
      spacer 74
                    chr12:67230:+:...
                                            FALSE
##
      spacer_76
                    chr12:67247:-:...
                                            FALSE
                                                                : . . . , . . .
##
     spacer_121
                                            FALSE
                              :...,...
##
##
     seqinfo: 640 sequences (1 circular) from hg38 genome
##
     crisprNuclease: SpCas9
```

One can also sort gRNAs using several annotation columns. For instance, let's sort gRNAs using the CRISPRrater score, but also by prioritizing first gRNAs that have no 1-mismatch off-targets:

```
o <- order(guideSet$n1, -guideSet$score_crisprater)
# Ordering the GuideSet:
guideSet <- guideSet[o]
head(guideSet)</pre>
```

## GuideSet object with 6 ranges and 25 metadata columns: ## segnames ranges strand | protospacer pam ## <Rle> <IRanges> <Rle> | <DNAStringSet> <DNAStringSet> ## chr12 66943 - | GCTCTGCTGGTTCTGCACGA TGG spacer\_9 ## spacer\_112 67396 - | GCCCTTGCCGAGGGCGGAGG GGG chr12 spacer 107 67371 + | CCGAGTTGCTGCGCTGCTGC CGG ## chr12 + | CGGCCGCCGCGTCAGCACCA CGG ## spacer 74 chr12 67233 ## spacer\_76 chr12 67244 - | GGCCCCGCTGGGGCTGCTCC AGG ## spacer\_121 chr12 67413 + | TCCCCTCCGCCTCGGCAA GGG ## cut\_site region percentGC polyC pam\_site polyA ## <numeric> <numeric> <character> <numeric> <logical> <logical> ## 66943 spacer\_9 66946 region\_1 60 FALSE FALSE ## spacer\_112 67396 67399 region\_1 80 **FALSE FALSE** ## spacer\_107 67371 67368 region\_1 70 FALSE **FALSE** ## 67233 67230 80 FALSE FALSE spacer\_74 region\_1 ## spacer\_76 67244 67247 region\_1 85 FALSE TRUE

```
##
     spacer_121
                      67413
                                 67410
                                                             75
                                                                     FALSE
                                                                                 TRUE
                                           region 1
##
                                 polyT startingGGGGG
                                                               n0
                                                                                   n0 c
                      polyG
                                                                          n1
##
                  <logical> <logical>
                                            <ld><logical> <numeric>
                                                                  <numeric>
                                                                             <numeric>
##
                      FALSE
                                                                           0
       spacer_9
                                 FALSE
                                                 FALSE
                                                                1
##
     spacer_112
                      FALSE
                                 FALSE
                                                 FALSE
                                                                1
                                                                           0
                                                                                      1
                                                                           0
##
     spacer 107
                      FALSE
                                 FALSE
                                                 FALSE
                                                                1
                                                                                      1
##
      spacer 74
                      FALSE
                                 FALSE
                                                 FALSE
                                                                1
                                                                           0
                                                                                      1
##
      spacer 76
                       TRUE
                                 FALSE
                                                 FALSE
                                                                1
                                                                           0
                                                                                      1
##
     spacer_121
                      FALSE
                                 FALSE
                                                 FALSE
                                                                1
                                                                           0
                                                                                      1
##
                       n1_c
                                alignments inRepeats score_cfd score_mit
##
                  <numeric> <GRangesList> <logical>
                                                       <numeric>
##
       spacer_9
                          0 chr12:66943:-
                                                 FALSE
                                                                1
                                                                           1
##
     spacer_112
                          0 chr12:67396:-
                                                 FALSE
                                                                1
                                                                           1
##
     spacer_107
                          0 chr12:67371:+
                                                 FALSE
                                                                1
                                                                           1
##
      spacer_74
                          0 chr12:67233:+
                                                 FALSE
                                                                1
                                                                           1
##
      spacer_76
                          0 chr12:67244:-
                                                 FALSE
                                                                1
                                                                           1
                          0 chr12:67413:+
##
                                                 FALSE
                                                                1
                                                                           1
     spacer_121
##
                 score_crisprater
                                          enzymeAnnotation
                                                                   geneAnnotation
##
                                     <SplitDataFrameList> <SplitDataFrameList>
                         <numeric>
##
       spacer 9
                          0.834319 FALSE:FALSE:FALSE:...
                                                                chr12:66946:-:...
##
     spacer_112
                          0.795745 FALSE:FALSE:FALSE:...
                                                                chr12:67399:-:...
##
     spacer_107
                          0.782780 FALSE:FALSE:FALSE:...
                                                                chr12:67368:+:...
##
      spacer_74
                          0.764870 FALSE:FALSE:FALSE:...
                                                                chr12:67230:+:...
##
      spacer 76
                          0.755493 FALSE: FALSE: FALSE: . . .
                                                                chr12:67247:-:...
##
     spacer_121
                          0.741315 FALSE:FALSE:FALSE:...
                                                                chr12:67410:+:...
##
                         tssAnnotation
                                            hasSNP
                                                                      snps
##
                  <SplitDataFrameList> <logical> <SplitDataFrameList>
##
       spacer_9
                     chr12:66946:-:...
                                             FALSE
                                                                 : . . . . . . .
##
     spacer_112
                                             FALSE
                               : . . . , . . .
                                                                 : . . . , . . .
##
     spacer_107
                                             FALSE
                               :...,...
##
      spacer_74
                     chr12:67230:+:...
                                             FALSE
##
      spacer_76
                     chr12:67247:-:...
                                             FALSE
##
     spacer_121
                                             FALSE
                               :...,...
                                                                 : . . . , . . .
##
##
     seqinfo: 640 sequences (1 circular) from hg38 genome
##
     crisprNuclease: SpCas9
```

The rankSpacers function is a convenience function that implements our recommended rankings for the SpCas9, enAsCas12a and CasRx nucleases. For a detailed description of our recommended rankings, see the documentation of rankSpacers by typing ?rankSpacers.

If an Ensembl transcript ID is provided, the ranking function will also take into account the position of the gRNA within the target CDS of the transcript ID in the ranking procedure. Our recommendation is to specify the Ensembl canonical transcript as the representative transcript for the gene. In our example, ENST00000538872 is the canonical transcript for IQSEC3:

# 5 CRISPRa/CRISPRi design

For CRISPRa and CRISPRi applications, the CRISPR nuclease is engineered to lose its endonuclease activity, therefore should not introduce double-stranded breaks (DSBs). We will use the dead SpCas9 (dSpCas9) nuclease as an example here. Note that users don't have to distinguish between dSpCas9 and SpCas9 when

specifying the nuclease in crisprDesign and crisprBase as they do not differ in terms of the characteristics stored in the CrisprNuclease object.

CRISPRi: Fusing dSpCas9 with a Krüppel-associated box (KRAB) domain has been shown to be effective at repressing transcription in mammalian cells (Gilbert et al. 2013). The dSpCas9-KRAB fused protein is a commonly-used construct to conduct CRISPR inhibition (CRISPRi) experiments. To achieve optimal inhibition, gRNAs are usually designed targeting the region directly downstream of the gene transcription starting site (TSS).

CRISPRa: dSpCas9 can also be used to activate gene expression by coupling the dead nuclease with activation factors. The technology is termed CRISPR activation (CRISPRa), and several CRISPRa systems have been developed (see Kampmann (2018) for a review). For optimal activation, gRNAs are usually designed to target the region directly upstream of the gene TSS.

crisprDesign provides functionalities to be able to take into account design rules that are specific to CRISPRa and CRISPRi applications. The queryTss function allows to specify genomic coordinates of promoter regions. The addTssAnnotation annotates gRNAs for known TSSs, and includes a column named dist\_to\_tss that indicates the distance between the TSS position and the PAM site of the gRNA. For CRISPRi, we recommend targeting the 25-75bp region downstream of the TSS for optimal inhibition. For CRISPRa, we recommend targeting the region 75-150bp upstream of the TSS for optimal activation; see (Sanson et al. 2018) for more information.

For more information, please see the following two tutorials:

- CRISPR activation (CRISPRa) design
- CRISPR interference (CRISPRi) design

## 6 CRISPR base editing with BE4max

We illustrate the CRISPR base editing (CRISPRbe) functionalities of crisprDesign by designing and characterizing gRNAs targeting IQSEC3 using the cytidine base editor BE4max (Koblan et al. 2018).

We first load the BE4max BaseEditor object available in crisprBase:

```
data(BE4max, package="crisprBase")
BE4max
## Class: BaseEditor
##
    CRISPR Nuclease name: SpCas9
##
       Target type: DNA
##
       Metadata: list of length 2
##
       PAMs: NGG, NAG, NGA
##
       Weights: 1, 0.2593, 0.0694
##
       Spacer length: 20
##
       PAM side: 3prime
##
         Distance from PAM: 0
       ##
##
    Base editor name: BE4max
       Editing strand: original
##
       Maximum editing weight: C2T at position -15
##
```

The editing probabilities of the base editor BE4max are stored in a matrix where rows correspond to the different nucleotide substitutions, and columns correspond to the genomic coordinate relative to the PAM site. The editingWeights function from crisprBase allows to retrieve those probabilities. One can see that C to T editing is optimal around 15 nucleotides upstream of the PAM site for the BE4max base editor:

```
crisprBase::editingWeights(BE4max)["C2T",]
```

```
##
     -36
            -35
                  -34
                        -33
                               -32
                                     -31
                                            -30
                                                  -29
                                                         -28
                                                               -27
                                                                                  -24
## 0.007 0.007 0.008 0.018 0.010 0.020 0.014 0.012 0.023 0.013 0.024 0.022 0.034
##
     -23
            -22
                  -21
                        -20
                               -19
                                     -18
                                            -17
                                                  -16
                                                         -15
                                                               -14
                                                                     -13
## 0.022 0.021 0.035 0.058 0.162 0.318 0.632 0.903 1.000 0.870 0.620 0.314 0.163
     -10
            -9
                   -8
                         -7
                                -6
                                      -5
                                             -4
                                                   -3
                                                          -2
## 0.100 0.056 0.033 0.019 0.018 0.024 0.017 0.005 0.002 0.001
```

We obtain a GuideSet object using the first exon of the IQSEC3 gene and retain only the first 2 gRNAs for the sake of time:

The function addEditedAlleles finds, characterizes, and scores predicted edited alleles for each gRNA, for a chosen transcript. It requires a transcript-specific annotation that can be obtained using the function getTxInfoDataFrame. Here, we will perform the analysis using the main isoform of IQSEC3 (transcript id ENST00000538872).

We first get the transcript table for ENST00000538872,

```
## DataFrame with 6 rows and 10 columns
##
              chr
                         pos
                                      nuc
                                                     aa aa_number
                                                                         exon pos_plot
##
     <character> <numeric> <character> <character> <integer>
                                                                   <integer> <integer>
## 1
            chr12
                       66767
                                        Α
                                                                NA
                                                                            1
                                                     NA
                                                                                      31
                                        G
## 2
            chr12
                       66768
                                                     NA
                                                                NA
                                                                            1
                                                                                      32
## 3
                                        G
                                                                NA
                                                                                      33
            chr12
                       66769
                                                     NA
                                                                            1
                                        С
## 4
            chr12
                       66770
                                                     NA
                                                                NA
                                                                            1
                                                                                      34
## 5
            chr12
                                        Τ
                                                     NA
                                                                NA
                                                                                      35
                       66771
                                                                            1
## 6
                       66772
                                                     NA
                                                                NA
                                                                                      36
            chr12
##
      pos_mrna
                  pos_cds
                                 region
##
     <integer> <integer> <character>
## 1
              1
                        NA
                                   5UTR
## 2
              2
                        NA
                                   5UTR
## 3
                                   5UTR
              3
                        NA
## 4
              4
                        NA
                                   5UTR
## 5
              5
                        NA
                                   5UTR
                        NA
                                   5UTR
```

and then add the edited alleles annotation to the GuideSet:

```
## [addEditedAlleles] Obtaining edited alleles at each gRNA target site.
```

## [addEditedAlleles] Adding functional consequences to alleles.

The editingWindow argument specifies the window of editing that we are interested in. When not provided, it uses the default window provided in the BaseEditor object. Note that providing large windows can exponentially increase computing time as the number of possible alleles grows exponentially.Let's retrieve the edited alleles for the first gRNA:

```
alleles <- editedAlleles(gs)[[1]]
```

It is a DataFrame object that contains useful metadata information:

#### metadata(alleles)

```
## $wildtypeAllele
##
          spacer_1
##
   "CGCGCACCGGATT"
##
## $start
## [1] 66901
##
## $end
##
  [1] 66913
##
## $chr
##
   [1] "chr12"
##
## $strand
  [1] "-"
##
##
## $editingWindow
  [1] -20 -8
##
## $wildtypeAmino
## [1] "NNNPPPVVVRRRA"
```

The wildtypeAllele reports the unedited nucleotide sequence of the region specified by the editing window (with respect to the gRNA PAM site). It is always reported from the 5' to 3' direction on the strand corresponding to the gRNA strand. The start and end specify the corresponding coordinates on the transcript.

Let's look at the edited alleles:

#### head(alleles)

```
## DataFrame with 6 rows and 4 columns
##
                seq
                         score
                                   variant
                                                       aa
##
     <DNAStringSet> <numeric> <character>
                                             <character>
     CGCGTATTGGATT 0.2471509
##
  1
                                  missense NNNPPPIIIRRRA
##
  2
      CGCGTATCGGATT 0.1618439
                                  missense NNNPPPIIIRRRA
## 3
      CGTGTATTGGATT 0.1057792
                                  missense NNNPPPIIIHHHA
## 4
      CGTGTATCGGATT 0.0692683
                                  missense NNNPPPIIIHHHA
## 5
      CGCGTACTGGATT 0.0372147
                                    silent NNNPPPVVVRRRA
      CGCGCATTGGATT 0.0292859
                                  missense NNNPPPMMMRRRA
```

The DataFrame is ordered so that the top predicted alleles (based on the score column) are shown first. The score represents the likelihood of the edited allele to occur relative to all possible edited alleles, and is calculated using the editing weights stored in the BE4max object. The seq column represents the edited

nucleotide sequences. Similar to the wildtypeAllele above, they are always reported from the 5' to 3' direction on the strand corresponding to the gRNA strand. The variant column indicates the functional consequence of the editing event (silent, nonsense or missense mutation). In case an edited allele leads to multiple editing events, the most detrimental mutation (nonsense over missense, missense over silent) is reported. The aa column reports the result edited amino acid sequence.

Note that several gRNA-level aggregate scores have also been added to the GuideSet object when calling addEditedAlleles:

head(gs)

```
## GuideSet object with 2 ranges and 11 metadata columns:
##
              segnames
                           ranges strand |
                                                      protospacer
                                                                              pam
##
                  <Rle> <IRanges>
                                    <Rle>
                                                  <DNAStringSet> <DNAStringSet>
                                          | CGCGCACCGGATTCTCCAGC
##
     spacer_1
                  chr12
                            66893
                                                                              AGG
##
     spacer_2
                  chr12
                            66896
                                        + | GGGCGGCATGGAGAGCCTGC
                                                                              TGG
##
               pam_site
                         cut_site
                                         region
##
               <numeric> <numeric> <character>
##
     spacer_1
                   66893
                             66896
                                       region_1
##
                   66896
                             66893
     spacer_2
                                       region_1
##
##
##
     spacer_1 CGCGTATTGGATT:0.247151:missense:...,CGCGTATCGGATT:0.161844:missense:...,CGTGTATTGGATT:0.1
     spacer_2
                  GGGTGGTATGGAG:0.4644396:silent:...,GGGCGGTATGGAG:0.2976235:silent:...,GGGTGGCATGGAG:0.
##
##
              score missense score nonsense score silent maxVariant
                                                 <numeric> <character>
##
                    <numeric>
                                    <numeric>
                    0.9020188
                                                 0.0745221
##
     spacer_1
                                                               missense
                                            \cap
                                                 0.9514897
##
     spacer 2
                    0.0036734
                                                                 silent
##
              maxVariantScore
##
                     <numeric>
##
     spacer_1
                      0.902019
##
                      0.951490
     spacer_2
##
```

The score\_missense, score\_nonsense and score\_silent columns represent aggregated scores for each of the mutation type. They were obtained by summing adding up all scores for a given mutation type across the set of edited alleles for a given gRNA. The maxVariant column indicates the most likely to occur mutation type for a given gRNA, and is based on the maximum aggregated score, which is stored in maxVariantScore. For instance, for spacer\_1, the higher score is the score\_missense, and therefore maxVariant is set to missense.

For more information, please see the following tutorial:

• CRISPR base editing (CRISPRbe) design

crisprNuclease: SpCas9

## ##

### 7 CRISPR knockdown with Cas13d

It is also possible to design gRNAs for RNA-targeting nucleases using crisprDesign. In contrast to DNA-targeting nucleases, the target spacer is composed of mRNA sequences instead of DNA genomic sequences.

We illustrate the functionalities of crisprDesign for RNA-targeting nucleases by designing gRNAs targeting IQSEC3 using the CasRx (RfxCas13d) nuclease (Konermann et al. 2018).

We first load the CasRx CrisprNuclease object from crisprBase:

seqinfo: 640 sequences (1 circular) from hg38 genome

```
data(CasRx, package="crisprBase")
CasRx

## Class: CrisprNuclease
## Name: CasRx
```

## Target type: RNA
## Metadata: list of length 2
## PFS: N
## Weights: 1
## Spacer length: 23
## PFS side: 3prime
## Distance from PFS: 0

## Prototype protospacers: 5'--SSSSSSSSSSSSSSSSSS[N]--3'

The PFS sequence (the equivalent of a PAM sequence for RNA-targeting nucleases) for CasRx is N, meaning that there is no specific PFS sequences preferred by CasRx.

We will now design CasRx gRNAs for the transcript ENST00000538872 of IQSEC3.

Let's first extract all mRNA sequences for IQSEC3:

## [1] 2701 AAGCCCCTCCCCTTCTCTGGGCC...AAAGTTACTGCTAGCATGGGTAA ENST00000382841 ## [2] 7087 AGGCTGGGCCGGTGGGAGAGGGA...TTATATTGAAAGATGTCACTTGA ENST00000538872

We can use the usual function findSpacers to design gRNAs, and we only consider a random subset of 100 gRNAs for the sake of time:

## GuideSet object with 6 ranges and 5 metadata columns:

```
##
                 segnames
                              ranges strand |
                                                           protospacer
##
                    <Rle> <IRanges> <Rle> |
                                                        <DNAStringSet>
##
     spacer_1000 region_1
                                1023
                                          + | TTGACCTAAAGAATAAACAGATT
##
     spacer_1001 region_1
                                1024
                                          + | TGACCTAAAGAATAAACAGATTG
                                          + | GACCTAAAGAATAAACAGATTGA
##
     spacer_1002 region_1
                                1025
##
     spacer_1003 region_1
                                1026
                                          + | ACCTAAAGAATAAACAGATTGAA
     spacer_1004 region_1
##
                                1027
                                          + | CCTAAAGAATAAACAGATTGAAA
##
     spacer_1005 region_1
                                1028
                                          + | CTAAAGAATAAACAGATTGAAAT
##
                             pam pam_site cut_site
                                                           region
                 <DNAStringSet> <numeric> <numeric> <character>
##
##
     spacer 1000
                               G
                                      1023
                                                  NA
                                                         region 1
##
     spacer 1001
                               Α
                                      1024
                                                   NA
                                                         region_1
     spacer_1002
##
                                      1025
                                                   NA
                               Α
                                                         region_1
##
     spacer_1003
                               Α
                                      1026
                                                   NA
                                                         region_1
     spacer_1004
                               Т
##
                                      1027
                                                   NA
                                                         region_1
##
     spacer_1005
                               G
                                      1028
                                                   NA
                                                         region_1
```

```
## -----
## seqinfo: 1 sequence from custom genome
## crisprNuclease: CasRx
```

Note that all protospacer sequences are located on the original strand of the mRNA sequence. For RNA-targeting nucleases, the spacer and protospacer sequences are the reverse complement of each other:

#### head(spacers(gs))

```
## DNAStringSet object of length 6:
##
       width seq
                                                                 names
          23 AATCTGTTTATTCTTTAGGTCAA
## [1]
                                                                  spacer_1000
## [2]
          23 CAATCTGTTTATTCTTTAGGTCA
                                                                  spacer_1001
## [3]
          23 TCAATCTGTTTATTCTTTAGGTC
                                                                  spacer_1002
## [4]
          23 TTCAATCTGTTTATTCTTTAGGT
                                                                  spacer_1003
## [5]
          23 TTTCAATCTGTTTATTCTTTAGG
                                                                  spacer_1004
## [6]
          23 ATTTCAATCTGTTTATTCTTTAG
                                                                  spacer_1005
head(protospacers(gs))
  DNAStringSet object of length 6:
##
```

```
width seq
##
                                                                  names
## [1]
          23 TTGACCTAAAGAATAAACAGATT
                                                                  spacer_1000
## [2]
          23 TGACCTAAAGAATAAACAGATTG
                                                                  spacer_1001
## [3]
          23 GACCTAAAGAATAAACAGATTGA
                                                                  spacer_1002
## [4]
          23 ACCTAAAGAATAAACAGATTGAA
                                                                  spacer_1003
## [5]
          23 CCTAAAGAATAAACAGATTGAAA
                                                                  spacer_1004
## [6]
          23 CTAAAGAATAAACAGATTGAAAT
                                                                  spacer_1005
```

The addSpacerAlignments can be used to perform an off-target search across all mRNA sequences using the argument custom\_seq. Here, for the sake of time, we only perform an off-target search to the 2 isoforms of IQSEC3 specified by the mRNAs object:

## GuideSet object with 6 ranges and 10 metadata columns:

```
protospacer
##
                  segnames
                              ranges strand |
##
                     <Rle> <IRanges>
                                       <Rle> |
                                                         <DNAStringSet>
##
     spacer_1095 region_1
                                 1118
                                           + | CGCCAATACCAGCTCAGCAAGAA
##
     spacer_1096 region_1
                                           + | GCCAATACCAGCTCAGCAAGAAC
                                 1119
##
     spacer_1097 region_1
                                1120
                                           + | CCAATACCAGCTCAGCAAGAACT
                                           + | CAATACCAGCTCAGCAAGAACTT
##
     spacer_1098 region_1
                                 1121
##
     spacer_1099 region_1
                                 1122
                                           + | AATACCAGCTCAGCAAGAACTTC
##
     spacer_1100 region_1
                                 1123
                                           + | ATACCAGCTCAGCAAGAACTTCG
##
                             pam pam_site cut_site
                                                            region
##
                  <DNAStringSet> <numeric> <numeric> <character> <numeric>
##
     spacer 1095
                               C
                                                                            2
                                       1118
                                                    NA
                                                          region 1
##
     spacer 1096
                               Т
                                       1119
                                                    NA
                                                          region 1
                                                                            2
##
     spacer_1097
                               Т
                                       1120
                                                    NA
                                                          region_1
                                                                            2
##
     spacer 1098
                               C
                                       1121
                                                    NA
                                                          region 1
                                                                            2
                                                                            2
##
     spacer_1099
                               G
                                       1122
                                                    NA
                                                          region_1
     spacer_1100
##
                               Α
                                       1123
                                                    NA
                                                          region_1
```

```
##
                      n1_tx
                              n0_gene
                                         n1_gene
##
                  <numeric> <numeric> <numeric>
##
     spacer 1095
                          0
                                     1
                                               0
                                               0
##
     spacer_1096
                          0
                                     1
##
     spacer_1097
                          0
                                     1
                                               0
                          0
                                               0
##
     spacer 1098
                                     1
                          0
                                               0
##
     spacer 1099
                                     1
                                               0
##
     spacer_1100
                          0
                                     1
##
                                                      alignments
##
                                                   <GRangesList>
##
     spacer_1095 ENST00000382841:505:+,ENST00000538872:1118:+
     spacer_1096 ENST00000382841:506:+,ENST00000538872:1119:+
##
##
     spacer_1097 ENST00000382841:507:+,ENST00000538872:1120:+
     spacer_1098 ENST00000382841:508:+,ENST00000538872:1121:+
##
##
     spacer_1099 ENST00000382841:509:+,ENST00000538872:1122:+
##
     spacer_1100 ENST00000382841:510:+,ENST00000538872:1123:+
##
##
     seqinfo: 1 sequence from custom genome
##
     crisprNuclease: CasRx
```

The columns no\_gene and no\_tx report the number of on-targets at the gene- and transcript-level, respectively. For instance, spacer\_1095 maps to the two isoforms of IQSEC3 has no\_tx is equal to 2:

```
onTargets(gs["spacer_1095"])
```

```
GRanges object with 2 ranges and 9 metadata columns:
                                                                        spacer
##
                         seqnames
                                      ranges strand |
##
                                                                   <character>
                             <Rle> <IRanges>
                                              <Rle> |
##
     spacer 1095 ENST00000382841
                                         505
                                                   + | TTCTTGCTGAGCTGGTATTG..
     spacer_1095 ENST00000538872
                                                       TTCTTGCTGAGCTGGTATTG..
##
                                        1118
##
                              protospacer
                                                       pam
                                                            pam_site n_mismatches
##
                           <DNAStringSet> <DNAStringSet>
                                                           <numeric>
                                                                         <numeric>
##
     spacer_1095 CGCCAATACCAGCTCAGCAAGAA
                                                         C
                                                                  505
                                                                                  0
                                                         C
                                                                                  0
##
     spacer_1095 CGCCAATACCAGCTCAGCAAGAA
                                                                 1118
##
                  canonical
                             cut_site
                                                gene_id gene_symbol
##
                  <logical>
                            <numeric>
                                            <character> <character>
##
                       TRUE
                                    NA ENSG00000120645
                                                             IQSEC3
     spacer_1095
     spacer_1095
##
                       TRUE
                                    NA ENSG00000120645
                                                             IQSEC3
##
##
     seqinfo: 2 sequences from custom genome
```

Note that one can also use the **bowtie** aligner to perform an off-target search to a set of mRNA sequences. This requires building a transcriptome bowtie index first instead of building a genome index. See the **crisprBowtie** vignette for more detail.

For more information, please see the following tutorial:

• CRISPR knockdown (CRISPRkd) design with CasRxdesign

# 8 Design for optical pooled screening (OPS)

Optical pooled screening (OPS) combines image-based sequencing (in situ sequencing) of gRNAs and optical phenotyping on the same physical wells (Feldman et al. 2019). In such experiments, gRNA spacer sequences are partially sequenced from the 5 prime end. From a gRNA design perspective, additional gRNA design constraints are needed to ensure sufficient dissimilarity of the truncated spacer sequences. The length of the truncated sequences, which corresponds to the number of sequencing cycles, is fixed and chosen by the

experimentalist.

To illustrate the functionalities of crisprDesign for designing OPS libraries, we use the guideSetExample. We will design an OPS library with 8 cycles.

```
n_cycles=8
```

We add the 8nt OPS barcodes to the GuideSet using the addOpsBarcodes function:

```
## DNAStringSet object of length 6:
##
       width seq
                                                                   names
## [1]
           8 CGCGCACC
                                                                   spacer_1
## [2]
           8 GGGCGGCA
                                                                   spacer_2
## [3]
           8 GGAGAGCC
                                                                   spacer_3
## [4]
           8 AGGTAGAG
                                                                   spacer_4
## [5]
           8 GAGCTCCT
                                                                   spacer_5
           8 CGATGGCC
## [6]
                                                                   spacer_6
```

The function getBarcodeDistanceMatrix calculates the nucleotide distance between a set of query barcodes and a set of target barcodes. The type of distance (hamming or levenstein) can be specified using the dist\_method argument. The Hamming distance (default) only considers substitutions when calculating distances, while the Levenstein distance allows insertions and deletions.

When the argument binnarize is set to FALSE, the return object is a matrix of pairwise distances between query and target barcodes:

```
## 5 x 5 sparse Matrix of class "dgCMatrix"
##
             CGATGGCC GCGCGCCG GCTCTACC GCTCTGCT GGGTGTGG
## CGCGCACC
                              7
                                        5
                                                  7
                                                            7
                    4
                                        5
## GGGCGGCA
                    4
                              3
                                                  4
                                                            4
                    3
                              6
                                        5
                                                  5
                                                            6
## GGAGAGCC
## AGGTAGAG
                    5
                              6
                                        8
                                                  7
                                                            4
                    7
## GAGCTCCT
                              3
                                        4
                                                  3
```

When binnarize is set to TRUE (default), the matrix of distances is binnarized so that 1 indicates similar barcodes, and 0 indicates dissimilar barcodes. The min\_dist\_edit argument specifies the minimal distance between two barcodes to be considered dissimilar:

The designOpsLibrary allows users to perform a complete end-to-end library design; see ?designOpsLibrary for documentation.

For more information, please see the following tutorial:

• Design for OPS

## 9 Design of gRNA pairs with the PairedGuideSet object

The findSpacerPairs function in crisprDesign enables the design of pairs of gRNAs and works similar to findSpacers. As an example, we will design candidate pairs of gRNAs that target a small locus located on chr12 in the human genome:

```
library(GenomicRanges)
library(BSgenome.Hsapiens.UCSC.hg38)
library(crisprBase)
bsgenome <- BSgenome.Hsapiens.UCSC.hg38</pre>
```

We first specify the genomic locus:

```
gr <- GRanges(c("chr12"),
IRanges(start=22224014, end=22225007))
```

and find all pairs using the function findSpacerPairs:

```
pairs <- findSpacerPairs(gr, gr, bsgenome=bsgenome)</pre>
```

The first and second arguments of the function specify the which genomic region the first and second gRNA should target, respectively. In our case, we are targeting the same region with both gRNAs. The other arguments of the function are similar to the findSpacers function described below.

The output object is a PairedGuideSet, which can be thought of a list of two GuideSet:

pairs

```
## PairedGuideSet object with 2626 pairs and 4 metadata columns:
```

```
##
                        first
                                         second | pamOrientation pamDistance
##
                   <GuideSet>
                                     <GuideSet> |
                                                      <character>
                                                                      <numeric>
##
        [1] chr12:22224025:- chr12:22224033:+ |
                                                               out
                                                                              8
        [2] chr12:22224025:- chr12:22224055:- |
                                                                             30
##
                                                               rev
        [3] chr12:22224033:+ chr12:22224055:- |
##
                                                                             22
                                                                in
        [4] chr12:22224025:- chr12:22224056:- |
##
                                                                             31
                                                               rev
##
        [5] chr12:22224033:+ chr12:22224056:- |
                                                                             23
                                                                in
##
##
     [2622] chr12:22224937:- chr12:22224994:+ |
                                                                             57
                                                               out
     [2623] chr12:22224938:- chr12:22224994:+ |
                                                                             56
##
                                                               out
##
     [2624] chr12:22224944:- chr12:22224994:+ |
                                                                             50
                                                               out
     [2625] chr12:22224950:+ chr12:22224994:+ |
##
                                                               fwd
                                                                             44
##
     [2626] chr12:22224958:- chr12:22224994:+ |
                                                               out
                                                                             36
##
            spacerDistance cutLength
##
                  <integer> <numeric>
##
        [1]
                        -32
                                     2
##
                                    30
        [2]
                         11
##
        [3]
                         24
                                    28
```

```
##
          [4]
                               12
                                            31
##
          [5]
                               25
                                            29
##
                              . . .
                                           . . .
##
      [2622]
                               17
                                            51
##
      [2623]
                               16
                                            50
##
                                            44
      [2624]
                               10
##
      [2625]
                               25
                                            44
##
      [2626]
                               -4
                                            30
```

The first and second GuideSet store information about gRNAs at position 1 and position 2, respectively. They can be accessed using the first and second functions:

```
grnas1 <- first(pairs)
grnas2 <- second(pairs)
grnas1</pre>
```

```
## GuideSet object with 2626 ranges and 5 metadata columns:
                            ranges strand |
##
               segnames
                                                      protospacer
                                                                              pam
##
                  <Rle> <IRanges>
                                    <Rle> |
                                                   <DNAStringSet> <DNAStringSet>
##
      spacer_1
                  chr12 22224025
                                        - | ATTAGTACAACCTTTCTTTT
                                                                              AGG
##
      spacer_1
                  chr12 22224025
                                        - | ATTAGTACAACCTTTCTTTT
                                                                              AGG
                  chr12 22224033
                                                                              AGG
##
      spacer_2
                                        + | CTTTTGTTTTCCTAAAAGAA
##
      spacer_1
                         22224025
                                        - | ATTAGTACAACCTTTCTTTT
                                                                              AGG
                  chr12
                                                                              AGG
##
      spacer_2
                  chr12 22224033
                                        + | CTTTTGTTTTCCTAAAAGAA
##
           . . .
                    . . .
                                                                              . . .
##
     spacer_68
                  chr12 22224937
                                        - | GGCTGCCAGTCATTGGATCA
                                                                              GGG
##
                  chr12 22224938
                                          | AGGCTGCCAGTCATTGGATC
                                                                              AGG
     spacer_69
##
     spacer_70
                  chr12 22224944
                                        - | TTTATAAGGCTGCCAGTCAT
                                                                              TGG
                                        + | GTGAGCCCTGATCCAATGAC
                                                                              TGG
##
     spacer_71
                  chr12 22224950
##
                  chr12 22224958
                                        - | CACTGTTTTTTTTTTATA
                                                                              AGG
     spacer_72
##
                pam_site cut_site
                                         region
##
               <numeric> <numeric> <character>
##
                22224025 22224028
                                       region_1
      spacer_1
##
      spacer_1
                22224025
                          22224028
                                       region_1
##
      spacer_2 22224033
                          22224030
                                       region_1
##
      spacer_1
                22224025
                          22224028
                                       region_1
##
      spacer_2
                22224033
                          22224030
                                       region_1
##
           . . .
                      . . .
                                . . .
##
     spacer_68
               22224937
                          22224940
                                       region_1
##
     spacer 69
               22224938
                          22224941
                                       region 1
##
                22224944
     spacer_70
                          22224947
                                       region_1
##
     spacer 71
                22224950
                           22224947
                                       region_1
##
     spacer_72 22224958
                          22224961
                                       region_1
##
##
     seqinfo: 640 sequences (1 circular) from hg38 genome
     crisprNuclease: SpCas9
##
```

#### grnas2

## GuideSet object with 2626 ranges and 5 metadata columns:

```
##
               segnames
                            ranges strand |
                                                      protospacer
##
                  <Rle> <IRanges>
                                    <Rle>
                                                   <DNAStringSet> <DNAStringSet>
##
      spacer_2
                  chr12 22224033
                                        + | CTTTTGTTTTCCTAAAAGAA
                                                                              AGG
##
                                        - | TATTCTCATGCACTGCTAGT
                                                                              GGG
      spacer_3
                  chr12 22224055
##
      spacer_3
                  chr12 22224055
                                        - | TATTCTCATGCACTGCTAGT
                                                                              GGG
                                        - | ATATTCTCATGCACTGCTAG
                                                                              TGG
##
      spacer_4
                  chr12 22224056
```

```
##
                  chr12 22224056
                                       - | ATATTCTCATGCACTGCTAG
                                                                             TGG
      spacer_4
##
           . . .
                   . . .
                                                                             . . .
                  chr12 22224994
##
     spacer 73
                                       + | CAGTGACATAGATCATACAT
                                                                            AGG
                  chr12 22224994
     spacer_73
                                       + | CAGTGACATAGATCATACAT
                                                                            AGG
##
     spacer_73
##
                  chr12 22224994
                                       + | CAGTGACATAGATCATACAT
                                                                            AGG
                                                                            AGG
     spacer 73
                  chr12 22224994
                                       + | CAGTGACATAGATCATACAT
##
                                       + | CAGTGACATAGATCATACAT
##
     spacer 73
                  chr12 22224994
                                                                            AGG
##
                pam_site cut_site
                                        region
##
               <numeric> <numeric> <character>
##
      spacer_2
               22224033 22224030
                                      region_1
##
      spacer_3 22224055 22224058
                                      region_1
##
      spacer_3 22224055
                          22224058
                                      region_1
      spacer_4 22224056 22224059
##
                                      region_1
##
      spacer_4 22224056 22224059
                                      region_1
##
                     . . .
##
     spacer_73
               22224994
                          22224991
                                      region_1
##
     spacer_73 22224994
                          22224991
                                      region_1
##
     spacer 73 22224994
                          22224991
                                      region 1
                          22224991
##
     spacer_73 22224994
                                      region_1
##
     spacer 73 22224994
                          22224991
                                      region 1
##
     seqinfo: 640 sequences (1 circular) from hg38 genome
##
##
     crisprNuclease: SpCas9
```

The pamOrientation function returns the PAM orientation of the pairs:

```
head(pamOrientation(pairs))
```

```
## [1] "out" "rev" "in" "rev" "in" "rev"
```

and takes 4 different values: in (for PAM-in configuration) out (for PAM-out configuration), fwd (both gRNAs target the forward strand) and rev (both gRNAs target the reverse strand).

The function pamDistance returns the distance between the PAM sites of the two gRNAs. The function cutLength returns the distance between the cut sites of the two gRNAs. The function spacerDistance returns the distance between the two spacer sequences of the gRNAs.

For more information, please see the following tutorial:

• Paired gRNA design

# 10 Miscellaneous design use cases

#### 10.1 Design with custom sequences

crisprDesign also allows gRNA design for DNA sequences without genomic context (such as a synthesized DNA construct). See ?findSpacers for more information, and here's an example:

```
## GuideSet object with 6 ranges and 5 metadata columns:
##
              seqnames
                          ranges strand |
                                                   protospacer
                                                                           pam
##
                 <Rle> <IRanges> <Rle> |
                                                <DNAStringSet> <DNAStringSet>
##
     spacer_1
                  seq1
                             12
                                      - | CGCCGCCCCGCGCCCGGGTC
                                                                           GGG
                  seq1
                              13
                                      - | GCGCCGCCCCGCGCT
                                                                           CGG
##
     spacer 2
```

```
##
     spacer_3
                   seq1
                                23
                                        + | GCGGAGGCCCGACCCGGGCG
                                                                               CGG
##
                                24
                                        + | CGGAGGCCCGACCCGGGCGC
                                                                               GGG
     spacer_4
                   seq1
     spacer_5
##
                   seq1
                                25
                                        + | GGAGGCCCGACCCGGGCGCG
                                                                               GGG
                                28
                                        + | GGCCCGACCCGGGCGCGGGG
                                                                               CGG
##
     spacer_6
                   seq1
##
               pam_site cut_site
                                         region
               <numeric> <numeric> <character>
##
                      12
##
     spacer_1
                                15
                                           seq1
##
     spacer_2
                      13
                                 16
                                           seq1
##
     spacer_3
                      23
                                 20
                                           seq1
                      24
##
     spacer_4
                                 21
                                           seq1
##
     spacer_5
                      25
                                 22
                                           seq1
                      28
                                 25
##
     spacer_6
                                           seq1
##
##
     seqinfo: 2 sequences from custom genome
##
     crisprNuclease: SpCas9
```

## 10.2 Off-target search in custom sequences

One can also search for off-targets in a custom sequence as follows:

```
GRanges object with 1 range and 7 metadata columns:
##
                 seqnames
                              ranges strand |
                                                             spacer
##
                     <Rle> <IRanges>
                                      <Rle> |
                                                     <DNAStringSet>
##
     spacer_1 custom_seq1
                                  21
                                           + | AAGACCCGGGCGCGGGCGG
##
                        protospacer
                                                pam pam_site n_mismatches canonical
##
                     <DNAStringSet> <DNAStringSet> <numeric>
                                                                 <numeric> <logical>
     spacer_1 TTGACCCGGGCGCGGGGGGGG
                                                GGG
                                                                                 TRUE
##
##
               cut_site
##
              <numeric>
##
     spacer_1
                     18
##
##
     seqinfo: 1 sequence from custom genome
```

For more information, please see the following tutorial:

• Working with custom DNA sequences

### 11 Session Info

```
## R version 4.2.1 (2022-06-23)
## Platform: x86_64-apple-darwin17.0 (64-bit)
## Running under: macOS Catalina 10.15.7
##
## Matrix products: default
```

```
/Library/Frameworks/R.framework/Versions/4.2/Resources/lib/libRblas.0.dylib
## LAPACK: /Library/Frameworks/R.framework/Versions/4.2/Resources/lib/libRlapack.dylib
##
## locale:
## [1] en_US.UTF-8/en_US.UTF-8/en_US.UTF-8/C/en_US.UTF-8/en_US.UTF-8
##
## attached base packages:
                           graphics grDevices utils
## [1] stats4
                 stats
                                                         datasets methods
## [8] base
##
## other attached packages:
## [1] Rbowtie_1.37.0
                                          BSgenome.Hsapiens.UCSC.hg38_1.4.4
## [3] BSgenome_1.65.2
                                          rtracklayer_1.57.0
## [5] Biostrings_2.65.2
                                          XVector_0.37.0
## [7] GenomicRanges_1.49.1
                                          GenomeInfoDb_1.33.5
## [9] IRanges_2.31.2
                                          S4Vectors_0.35.1
## [11] BiocGenerics_0.43.1
                                          crisprDesign_0.99.134
## [13] crisprBase_1.1.5
## loaded via a namespace (and not attached):
##
     [1] bitops_1.0-7
                                       matrixStats_0.62.0
     [3] bit64_4.0.5
                                       filelock_1.0.2
##
##
                                       httr_1.4.4
     [5] progress_1.2.2
##
     [7] tools 4.2.1
                                       utf8 1.2.2
##
     [9] R6 2.5.1
                                       DBI 1.1.3
## [11] tidyselect_1.1.2
                                       prettyunits_1.1.1
## [13] bit_4.0.4
                                       curl_4.3.2
## [15] compiler_4.2.1
                                       crisprBowtie_1.1.1
## [17] cli_3.3.0
                                       Biobase_2.57.1
## [19] basilisk.utils_1.9.1
                                       crisprScoreData_1.1.3
## [21] xml2_1.3.3
                                       DelayedArray_0.23.1
## [23] randomForest_4.7-1.1
                                       readr_2.1.2
## [25] rappdirs_0.3.3
                                       stringr_1.4.1
## [27] digest_0.6.29
                                       Rsamtools_2.13.4
                                       crisprScore_1.1.14
## [29] rmarkdown 2.15.2
## [31] basilisk_1.9.3
                                       pkgconfig_2.0.3
## [33] htmltools 0.5.3
                                       MatrixGenerics 1.9.1
## [35] dbplyr_2.2.1
                                       fastmap_1.1.0
## [37] rlang_1.0.4
                                       rstudioapi_0.14
## [39] RSQLite_2.2.16
                                       shiny_1.7.2
## [41] BiocIO 1.7.1
                                       generics 0.1.3
## [43] jsonlite_1.8.0
                                       vroom_1.5.7
## [45] BiocParallel_1.31.12
                                       dplyr 1.0.9
## [47] VariantAnnotation_1.43.3
                                       RCurl_1.98-1.8
## [49] magrittr_2.0.3
                                       GenomeInfoDbData_1.2.8
## [51] Matrix_1.4-1
                                       Rcpp_1.0.9
## [53] fansi_1.0.3
                                       reticulate_1.25
## [55] lifecycle_1.0.1
                                       stringi_1.7.8
## [57] yaml_2.3.5
                                       SummarizedExperiment_1.27.1
## [59] zlibbioc_1.43.0
                                       BiocFileCache_2.5.0
## [61] AnnotationHub_3.5.0
                                       grid_4.2.1
## [63] blob 1.2.3
                                       promises_1.2.0.1
## [65] parallel_4.2.1
                                       ExperimentHub_2.5.0
## [67] crayon_1.5.1
                                       crisprBwa_1.1.3
```

```
##
    [69] dir.expiry 1.5.0
                                         lattice 0.20-45
##
    [71] GenomicFeatures 1.49.6
                                         hms_1.1.2
##
    [73] KEGGREST 1.37.3
                                         knitr 1.40
    [75] pillar_1.8.1
                                         rjson_0.2.21
##
##
    [77] codetools 0.2-18
                                         biomaRt 2.53.2
    [79] BiocVersion 3.16.0
                                         XML 3.99-0.10
##
    [81] glue 1.6.2
                                         evaluate 0.16
##
                                         httpuv 1.6.5
##
    [83] BiocManager_1.30.18
##
    [85] png 0.1-7
                                         vctrs 0.4.1
##
    [87] tzdb_0.3.0
                                         purrr_0.3.4
    [89] assertthat_0.2.1
                                         cachem_1.0.6
    [91] xfun_0.32
                                         mime_0.12
##
##
    [93] Rbwa_1.1.0
                                         xtable_1.8-4
    [95] restfulr_0.0.15
                                         later_1.3.0
##
    [97] tibble_3.1.8
                                         GenomicAlignments_1.33.1
##
##
    [99] AnnotationDbi_1.59.1
                                         memoise_2.0.1
## [101] interactiveDisplayBase_1.35.0 ellipsis_0.3.2
```

## References

Feldman, David, Avtar Singh, Jonathan L Schmid-Burgk, Rebecca J Carlson, Anja Mezger, Anthony J Garrity, Feng Zhang, and Paul C Blainey. 2019. "Optical Pooled Screens in Human Cells." *Cell* 179 (3): 787–99.

Gilbert, Luke A, Matthew H Larson, Leonardo Morsut, Zairan Liu, Gloria A Brar, Sandra E Torres, Noam Stern-Ginossar, et al. 2013. "CRISPR-Mediated Modular RNA-Guided Regulation of Transcription in Eukaryotes." *Cell* 154 (2): 442–51.

Kampmann, Martin. 2018. "CRISPRi and CRISPRa Screens in Mammalian Cells for Precision Biology and Medicine." ACS Chemical Biology 13 (2): 406–16.

Koblan, Luke W, Jordan L Doman, Christopher Wilson, Jonathan M Levy, Tristan Tay, Gregory A Newby, Juan Pablo Maianti, Aditya Raguram, and David R Liu. 2018. "Improving Cytidine and Adenine Base Editors by Expression Optimization and Ancestral Reconstruction." *Nature Biotechnology* 36 (9): 843–46.

Konermann, Silvana, Peter Lotfy, Nicholas J Brideau, Jennifer Oki, Maxim N Shokhirev, and Patrick D Hsu. 2018. "Transcriptome Engineering with RNA-Targeting Type VI-d CRISPR Effectors." *Cell* 173 (3): 665–76.

Langmead, Ben, Cole Trapnell, Mihai Pop, and Steven L. Salzberg. 2009. "Ultrafast and Memory-Efficient Alignment of Short DNA Sequences to the Human Genome." Genome Biology 10 (3): R25. https://doi.org/10.1186/gb-2009-10-3-r25.

Sanson, Kendall R, Ruth E Hanna, Mudra Hegde, Katherine F Donovan, Christine Strand, Meagan E Sullender, Emma W Vaimberg, et al. 2018. "Optimized Libraries for CRISPR-Cas9 Genetic Screens with Multiple Modalities." *Nature Communications* 9 (1): 1–15.