# Package 'G4SNVHunter'

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Type Package

Title Evaluating SNV-Induced Disruption of G-Quadruplex Structures

**Version** 0.99.4

**Description** G-quadruplexes (G4s) are unique nucleic acid secondary structures predominantly found in guanine-rich regions and have been shown to be involved in various biological regulatory processes. G4SNVHunter is an R package designed to rapidly identify genomic sequences with G4-forming potential and accurately screen user-provided single nucleotide variants (also applicable to single nucleotide polymorphisms) that may destabilize these structures. This enables users to screen key variants for further experimental study, investigating how these variants may influence biological functions, such as gene regulation, by altering G4 formation.

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LazyData false

**Depends** R (>= 4.3.0)

**Imports** Biostrings, stats, GenomicRanges, IRanges, Rcpp, RcppRoll, data.table, GenomeInfoDb, S4Vectors, ggplot2, cowplot, progress, ggseqlogo, viridis, ggpointdensity

LinkingTo Rcpp

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VignetteBuilder knitr

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News NEWS.md

URL https://github.com/rongxinzh/G4SNVHunter

BugReports https://github.com/rongxinzh/G4SNVHunter/issues

Config/testthat/edition 3

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# **Description**

This function checks whether the user-provided SNVs (or SNPs) are single nucleotide substitutions.

# Usage

```
checkSNV(snv_gr = NULL, mode = "wra", ref_col = NULL, alt_col = NULL)
```

# Arguments

snv_gr	A GRanges object containing SNV (or SNP) data.
mode	A character string specifying the checks to be performed. w for checking if all widths (w) are 1, r for checking if all ref (r) values are A, T, C, or G, a for checking if all alt (a) values are A, T, C, or G.
ref_col	Column name for the ref bases in snv_gr. Default is NULL.
alt_col	Column name for the alt bases in snv_gr. Default is NULL.

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### Value

A logical value indicating whether the user-provided SNVs (or SNPs) passed all checks.

```
if (!requireNamespace("BiocManager", quietly = TRUE)) {
  install.packages("BiocManager")
}
if (!requireNamespace("BSgenome.Hsapiens.UCSC.hg19", quietly = TRUE)) {
  BiocManager::install("GenomicRanges")
}
library(GenomicRanges)
gr1 <- GRanges("chr1", IRanges(start = 100, width = 1))</pre>
# check width ('w')
checkSNV(gr1, mode = "w")
gr2 <- GRanges(</pre>
  seqnames = Rle("seq1"),
  ranges = IRanges(c(100, 200, 300), width = 1),
  ref = c("A", "C", "G"),
  alt = c("T", "T", "A")
# check width ('w'), ref ('r'), and alt ('a')
checkSNV(gr2, mode = "wra", ref_col = "ref", alt_col = "alt")
# check width ('w') and alt ('a')
checkSNV(gr2, mode = "wa", alt_col = "alt")
gr3 <- GRanges("chr1", IRanges(start = 100, width = 10))</pre>
# widths should be all one
checkSNV(gr3, mode = "w")
gr4 <- GRanges(</pre>
  seqnames = Rle("seq1"),
  ranges = IRanges(start = 100, width = 1),
  ref = "AG",
  alt = T
)
# ref should be all one
checkSNV(gr4, mode = "wr", ref_col = "ref")
```

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## **Description**

This function filters the SNV Impact GRanges object returned by the SNVImpactG4 function based on user-defined thresholds for the G4.info.score, mut.score, and score.diff parameters. This function filters SNVs that may significantly impact the formation of G4 structures using customizable filtering criteria. You are not required to specify all three threshold parameters. However, at least one threshold parameter must be provided.

# Usage

```
filterSNVImpact(
   gr,
   raw_score_threshold = NULL,
   mut_score_threshold = NULL,
   score_diff_threshold = NULL)
```

## **Arguments**

gr A GRanges object returned by the SNVImpactG4 function.

raw\_score\_threshold

A positive numeric value no greater than 4 used as the threshold for the absolute value of G4.info.score. G4s with an absolute G4Hunter score exceeding this threshold will be retained. If NULL, this threshold is not applied.

mut\_score\_threshold

A positive numeric value no greater than 4 used as the threshold for the absolute value of mut.score. Mutated G4s with an absolute G4Hunter score below this threshold will be retained. If NULL, this threshold is not applied.

score\_diff\_threshold

A negative numeric value no less than -4 used as the threshold for score.diff. G4s with a decrease in G4Hunter score greater than this threshold after variation will be retained. If NULL, this threshold is not applied.

#### Value

A filtered GRanges object, containing only the records that meet the specified threshold criteria.

# See Also

SNVImpactG4 for assessing the impact of SNVs on G4 formation.

```
if (!requireNamespace("BiocManager", quietly = TRUE)) {
   install.packages("BiocManager")
}

if (!requireNamespace("GenomicRanges", quietly = TRUE)) {
   BiocManager::install("GenomicRanges")
}
```

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```
if (!requireNamespace("BSgenome.Hsapiens.UCSC.hg19", quietly = TRUE)) {
  BiocManager::install("BSgenome.Hsapiens.UCSC.hg19")
}
library(GenomicRanges)
library(BSgenome.Hsapiens.UCSC.hg19)
hg19 <- BSgenome.Hsapiens.UCSC.hg19
chr21_seq <- DNAStringSet(hg19$chr21)</pre>
# Chromosome name is needed
names(chr21\_seq) \leftarrow "chr21"
G4 <- G4HunterDetect(chr21_seq)
data(snp_gr)
res_snp <- SNVImpactG4(G4, snp_gr, alt_col = "alt")</pre>
filtered_snv_eff <- filterSNVImpact(res_snp,</pre>
  mut_score_threshold = 1.2,
  score\_diff\_threshold = -0.2
print(filtered_snv_eff)
```

G4HunterDetect

Detect G4 Sequences Using G4Hunter Algorithm

## **Description**

This function detects G4 sequences for a given DNAStringSet object using the G4Hunter algorithm.

## Usage

```
G4HunterDetect(
  sequences = NULL,
  threshold = 1.5,
  window_size = 25,
  include_sequences = TRUE,
  strands = "b"
)
```

## Arguments

sequences A DNAStringSet object containing the sequences to be analyzed.

threshold A numeric value specifying the threshold for the G4Hunter score (absolute

value). Default is 1.5. We do not recommend setting the threshold below 1.2.

window\_size An integer specifying the window size (bp) for prediction. Default is 25. An-

other commonly used window size is 20. However, 25 is generally preferred

unless otherwise required.

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include\_sequences

A logical value (TRUE/FALSE) indicating whether to include the predicted G4 sequences in the output. Default is TRUE. Setting this parameter to FALSE can reduce the memory overhead of the final output, which may be useful for extremely large genomes. However, we strongly recommend retaining the sequence information in the output, as it is indispensable for subsequent analysis of the impact of variants on G4 formation.

strands

A character string ("b" for both strands and "p" for positive strand only) indicating which strand to be considered. Default is "b".

#### Value

A GRanges object containing the predicted G4 sequences. The GRanges object includes the following metadata columns:

score The actual G4Hunter score for the G4 sequence detected (after merging, pruning, etc.). max\_score The maximum G4Hunter score for the window covering this G4. sequence The G4 sequence after merging, pruning, etc.

If no G4 sequences were detected, it will return an empty GRanges object.

#### See Also

loadSequence for loading genome sequences into a DNAStringSet object.

```
if (!requireNamespace("BiocManager", quietly = TRUE)) {
 install.packages("BiocManager")
}
if (!requireNamespace("Biostrings", quietly = TRUE)) {
 BiocManager::install("Biostrings")
library(Biostrings)
sequences <- DNAStringSet(c(</pre>
  "AGTGAATGGGATGGGAGGAGGGACGGGGTAGTACAGCATAGCATG",
  "TAGGTAGCTACGACACCCTGCCCTACCCCTATCTA"
names(sequences) <- c("seq1", "seq2")</pre>
G4s <- G4HunterDetect(sequences, threshold = 1.5, window_size = 25)
print(G4s)
seq_path <- system.file("extdata", "seq.fa", package = "G4SNVHunter")</pre>
G4s <- G4HunterDetect(loadSequence(seq_path = seq_path))
print(G4s)
seq_path <- system.file("extdata", "seq.txt", package = "G4SNVHunter")</pre>
G4s <- G4HunterDetect(loadSequence(seq_path = seq_path))
print(G4s)
```

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G4HunterScore Calculate the G4Hunter Score for a Given Sequence
---

# Description

This function calculates the G4Hunter score for a given nucleotide sequence, which reflects the ability of that sequence to form a G4 structure.

# Usage

```
G4HunterScore(seg = NULL)
```

#### **Arguments**

seq

A single character string representing the nucleotide sequence. Must contain only the characters A, T, C, G, U, and N. The length of the sequence should not be too short (e.g., less than 10 bp).

#### Value

A numeric value representing the G4Hunter score for the provided sequence.

#### See Also

G4HunterDetect for detecting the G4 sequences in a given DNAStringSet object.

### **Examples**

```
sequence <- "GGGTAAGGGATGGGTCGGG"
score <- G4HunterScore(sequence)
print(score)
# A negative value indicates that the G4 sequence
# is located on the reverse strand
sequence <- "GGGTAAGGGATGGGTCGGG"
score <- G4HunterScore(sequence)
print(score)</pre>
```

loadSequence

Load Genome Sequences

## **Description**

This function loads genomic sequences from various sources, including a FASTA file, a text file with sequence identifiers and corresponding sequences, or a data frame object.

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#### Usage

```
loadSequence(genome_seq = NULL, seq_path = NULL)
```

#### **Arguments**

genome\_seq
seq\_path

A data frame containing sequence identifiers and corresponding sequences.

A character string specifying the file path to a FASTA file (suffixed with .fa, .fna or .fasta) or a text file with sequence identifiers and corresponding sequences (file headers should not be provided). Ignored if genome\_seq is pro-

vided.

#### Value

A DNAStringSet object containing the genome sequences.

# **Examples**

```
# File path for sequences in fasta format
fa_path <- system.file("extdata", "seq.fa", package = "G4SNVHunter")
seq <- loadSequence(seq_path = fa_path)
print(seq)

# Another example
# Load sequences from data.frame
seq_df <- data.frame(
    chr = c("seq1", "seq2"),
    sequence = c(
        paste0(rep("G", 100), collapse = ""),
        paste0(rep("A", 100), collapse = "")
)
)
seq <- loadSequence(genome_seq = seq_df)
print(seq)</pre>
```

plotG4Info

Plot Basic Statistics of G4s Detected by the G4HunterDetect Function

# **Description**

This function generates a series of plots to visualize basic statistics of G4 sequences predicted by the G4Hunter algorithm. The function produces the following plots:

- Distribution of max scores (absolute values).
- Distribution of max scores, split by strand.
- Distribution of scores (absolute values).
- Distribution of scores, split by strand.
- Distribution of sequence lengths (with lengths greater than 50 bp grouped into a single bin).
- Distribution of sequence lengths, split by strand (with lengths greater than 50 bp grouped into a single bin).

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### Usage

```
plotG4Info(
   G4,
   p_colors = c("#6B9ECC", "#D91117", "#0E619C", "#58AC7B", "#D91117", "#0E619C",
        "#F39C40", "#D91117", "#0E619C")
)
```

#### **Arguments**

G4

A GRanges object containing G4 sequences. The object must include at least the following metadata columns:

- score: Numeric vector of scores for G4 sequences.
- max\_score: Numeric vector of maximum scores for G4 sequences.
- sequence: Character vector of G4 sequence strings.

The GRanges object must also have a strand information. Whenever possible, it should come from the results returned by the G4HunterDetect function.

p\_colors

A vector of colors to be used for the plots. It should contain nine color values for the different plot elements.

### Value

A combined plot displays all the generated plots arranged in a grid layout.

# See Also

G4HunterDetect for detecting the G4 sequences in a given DNAStringSet object.

## **Examples**

```
seq_path <- system.file("extdata", "seq.txt", package = "G4SNVHunter")
G4 <- G4HunterDetect(loadSequence(seq_path = seq_path))
plotG4Info(G4)</pre>
```

plotImpactSeq

Visualize the variants in G4 sequence

# **Description**

This function plot sequence logos to visualize sequence variants caused by SNVs or SNPs, with the location of the variants highlighted by rectangles and arrows.

# Usage

```
plotImpactSeq(filtered_gr, ncol = 1)
```

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### Arguments

A GRanges object containing sequence data and G4Hunter scores. The object must have metadata columns named G4.info.score, mut.score, G4.info.sequence, and mut.G4.seq.

An integer specifying the number of columns in the output plot grid. Default is 1.

#### Value

A plot that displays the grid of sequence logos, showing the differences between the original and mutated sequences.

### See Also

SNVImpactG4 for evaluating the impact of SNVs on G4 formation, and filterSNVImpact for filtering G4s that are significantly affected by SNVs.

# **Examples**

plotSNVImpact

Plot the Impact of SNVs on G4Hunter Scores

### **Description**

This function generates two plots for visualizing the impact of SNVs on G4 formation: 1. A scatter plot with density shading comparing the original G4Hunter score and the mutant G4Hunter score. 2. A density plot showing the distribution of score changes between the original and mutant G4 sequences.

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### Usage

```
plotSNVImpact(gr, p_colors = c("#b22d2d", "#6ca4d6", "#2d69b0", "#1f77b4"))
```

### **Arguments**

gr

A GRanges object containing the G4Hunter scores and associated metadata. The object should include the following columns:

- G4. info. score: Numeric vector of the original G4Hunter scores.
- mut.score: Numeric vector of the G4Hunter scores after mutation.
- score.diff: Numeric vector of the differences between the original and mutant G4Hunter scores.

p\_colors

A vector of four colors used for plotting.

### Value

A combined plot (using plot\_grid) containing two subplots: - Density scatter plot comparing original vs mutant G4Hunter scores. - Density plot of the G4Hunter score differences.

#### See Also

SNVImpactG4 for evaluating the impact of SNVs on G4 formation, and filterSNVImpact for filtering G4s that are significantly affected by SNVs.

```
if (!requireNamespace("BiocManager", quietly = TRUE)) {
  install.packages("BiocManager")
if (!requireNamespace("GenomicRanges", quietly = TRUE)) {
  BiocManager::install("GenomicRanges")
if (!requireNamespace("BSgenome.Hsapiens.UCSC.hg19", quietly = TRUE)) {
  BiocManager::install("BSgenome.Hsapiens.UCSC.hg19")
}
library(GenomicRanges)
library(BSgenome.Hsapiens.UCSC.hg19)
hg19 <- BSgenome.Hsapiens.UCSC.hg19
chr21_seq <- DNAStringSet(hg19$chr21)</pre>
# Chromosome name is needed
names(chr21_seq) <- "chr21"</pre>
G4 <- G4HunterDetect(chr21_seq)
# Load SNVs
data(snp_gr)
```

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```
res_snp <- SNVImpactG4(G4, snp_gr, alt_col = "alt")
plotSNVImpact(res_snp)</pre>
```

snp\_gr

Single Nucleotide Polymorphisms GRanges Object

# **Description**

This dataset contains a GRanges object storing single nucleotide polymorphisms (SNPs).

# Usage

```
data(snp_gr)
```

### **Format**

A GRanges object with 30,000 SNPs.

SNVImpactG4

Evaluate the Impact of SNVs on G4 Sequences

# Description

This function evaluates the impact of SNVs on G4 formation.

# Usage

```
SNVImpactG4(
  G4 = NULL,
   snvs = NULL,
  alt_col = "alt",
  mode = "s",
  sampleid_col = NULL,
  snvid_col = NULL)
```

# **Arguments**

G4	A GRanges object representing the G4 regions. This object must include metadata columns for G4 sequence and G4Hunter scores. Whenever possible, it should come from the results returned by the G4HunterDetect function.
snvs	A GRanges object representing the SNVs. This object must include metadata columns for SNV alternative alleles and, optionally, SNV IDs.
alt_col	A character string specifying the name of the column in snvs that contains the alternative alleles for the SNVs. The default is "alt".

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mode A character string indicating the mode of operation. Set to "s" to evaluate the

impact of individual SNVs on G4 regions one at a time. Set to "m" to assess the combined impact of multiple SNVs that overlap the same G4 region within a sample. If using "m" mode, you must specify the sampleid\_col and snvid\_col

parameters.

sampleid\_col A character string specifying the name of the column in snvs that contains the

sample IDs. This parameter is required when mode is "m", and is ignored if mode

is "s".

snvid\_col A character string specifying the column name in snvs that contains SNV IDs.

Required when mode is "m". Ignored if mode is "s".

#### Value

A GRanges object depending on the mode parameter:

**Mode "s":** For mode = "s", the returned GRanges object includes the following metadata columns:

- segnames Identifiers for SNVs.
- ranges Position of the SNVs.
- strand Strand of the SNVs.
- SNV.info.\* Metadata columns related to each SNV.
- G4. info.\* Metadata columns from the original G4 object.
- mut. G4. seq The mutated G4 sequence after applying the SNV change.
- mut.G4.anno.seq The mutated G4 sequence, with the mutated bases annotated using square brackets.
- mut. score The G4Hunter score of the mutated sequence.
- score.diff The difference between the mutated G4Hunter score and the original G4Hunter score. The value is calculated as the absolute value of the mutated G4Hunter score minus the absolute value of the original G4Hunter score.

**Mode "m":** For mode = "m", the returned GRanges object includes the following metadata columns:

- segnames Identifiers for G4 sequences.
- ranges Position of the G4 sequences (start and end).
- strand Strand of the G4 sequences.
- G4. info. \* Metadata columns from the original G4 object.
- snv. ids Concatenated SNV IDs for all SNVs affecting the G4 region.
- sample.ids A semicolon-separated list of sample IDs overlapping each G4 region.
- mut.G4.seq The mutated G4 sequence after applying the combined SNV changes.
- mut.G4.anno.seq The mutated G4 sequence, with the mutated bases annotated using square brackets.
- mut.score The G4Hunter score of the mutated sequence.
- score. diff The difference between the mutated G4Hunter score and the original G4Hunter score. This value is calculated as the absolute value of the mutated G4Hunter score minus the absolute value of the original G4Hunter score.

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#### See Also

G4HunterDetect for detecting the G4 sequences in a given DNAStringSet object. G4HunterScore for calculating the G4Hunter scores for a given sequence. filterSNVImpact for filtering out SNVs with significant impact.

```
if (!requireNamespace("BiocManager", quietly = TRUE)) {
  install.packages("BiocManager")
}
if (!requireNamespace("GenomicRanges", quietly = TRUE)) {
  BiocManager::install("GenomicRanges")
}
if (!requireNamespace("BSgenome.Hsapiens.UCSC.hg19", quietly = TRUE)) {
  BiocManager::install("BSgenome.Hsapiens.UCSC.hg19")
}
library(GenomicRanges)
library(BSgenome.Hsapiens.UCSC.hg19)
# Load sequence for chromosome 21 (hg19)
hg19 <- BSgenome.Hsapiens.UCSC.hg19
chr21_seq <- DNAStringSet(hg19$chr21)</pre>
# Chromosome name is needed
names(chr21_seq) <- "chr21"</pre>
# Detect G4s in human chromosome 21
G4 <- G4HunterDetect(chr21_seg)
# 's' mode
# Load SNPs
data(snp_gr)
# Obtain SNPs that overlap with G4 regions and assess their impact on G4.
# In variant-centric mode ('s'), evaluating each SNP individually.
res_snp <- SNVImpactG4(G4, snp_gr, alt_col = "alt")</pre>
print(res_snp)
# 'm' mode
# Load SNVs
data(snv_gr)
# Obtain SNVs that overlap with G4 regions and assess their impact on G4.
# In sample-centric mode ('m'), evaluate the combined impact of SNVs on G4s.
# Grouped by the sample IDs specified in 'sampleid_col'.
res_snv <- SNVImpactG4(G4, snv_gr,</pre>
  alt_col = "alt",
  mode = "m", sampleid_col = "sampleid", snvid_col = "snv_id"
print(res_snv)
```

snv\_gr

snv\_gr

Single Nucleotide Variant GRanges Object

# Description

This dataset contains a GRanges object storing single nucleotide variants (SNVs).

# Usage

```
data(snv_gr)
```

# **Format**

A GRanges object with 50,000 SNVs.

# **Index**

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