# Reference Manual: Sasquatch R-Tool



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**Requires:** ggplot2, RColorBrewer

Recommends: Biostrings, TFBSTools, phapply

# Contents

ontents	1
alcSFR	1
ompareSequences	2
ecodeKmer	4
issectSequence	4
etCount	5
etFootprint	5
etFootprintStrand	7
etPossibleMutations	7
etSFR	8
repProfile	9
SilicoMutation	10
SilicoMutationPlot	12
[akeInSilicoMutationTrackHub	13
lotOverlap	15
lotOverlapKmers	16
$lot Single \ldots \ldots$	18
lotSingleKmer	19
$lot Single Strands \ldots \ldots$	20
reLoadProfiles	21
reLoadVocab	21
runeProfile	22
uerv.Iaspar	22

QueryJasparBatch	23
QueryLongSequence	24
RefVarBatch	26
SmoothProfile	27
SobelBorders	28
Sobeln	28

# CalcSFR.

# Description

Calculate the Shoulder-to-Footprint-Ratio of a given footprint profile and estimated shoulder positions and ranges.

# Usage

```
> CalcSFR(profile,
+ upstream.shoulder.position,
+ upstream.shoulder.position,
+ upstream.shoulder.range,
+ upstream.shoulder.range)
```

# Arguments

```
    profile input cut profile (should be smoothed)
    us.mid center bp position of the upstream shoulder (as estimated from SobelBorders)
    ds.mid center bp position of the downstream shoulder
    range.us range in bp of the upstream shoulder (as estimated from SobelBorders)
    range in bp of the downstream shoulder
```

#### Value

Returns the SFR as single numeric value.

```
> frag.type <- "DNase"
> data.dir <- "./my_sasq_database/human/DNase/"
> tissue <- "blood_tissue"
> kmer <- "CACGTG"
> # get the average profile for a single kmer
> profile<- GetFootprint(
+ kmer=kmer,
+ tissue=tissue,
+ data.dir=data.dir,
+ frag.type=frag.type,
+ smooth=TRUE
+ )
> # estimate the shoulders from the profile
> sh <- SobelBorders(profile, kl=nchar(kmer))
> #calculate the SFR
> CalcSFR(profile, sh$us, sh$ds, sh$range.us, sh$range.ds)
```

# CompareSequences

# Description

Wrapper function to compare two input sequences on a kmer basis. Both sequence are split up and compared pairwise, calculating the damage. Total damage is calculated either as summed up or as highest pairwise damage, as chosen. Overlay plots for none, all or only the highest scoring pair are created as specified.

(If preload = TRUE and if vocab.flag = FALSE, preload.profiles has to be submitted. If vocab.flag = TRUE with preload = TRUE then preload.vocab has to be submitted. If preload =TRUE and plots should be generated, preload.profiles have to be provided.)

```
> CompareSequences(sequence1,
                   sequence2,
+
                   kl,
                   damage.mode="exhaustive",
                   tissue,
                   data.dir,
                   pnorm.tag,
                   vocab.flag=FALSE,
                   vocab.file=paste0(data.dir,"/",tissue,
                               "/vocabulary_",tissue,".txt"),
                   frag.type,
                   plots="highest",
                   smooth=TRUE,
                   ylim=c(0,0.01),
                   xlim=c(-125,125),
                   plot.shoulders=FALSE,
                   preload,
                   preload.vocab="",
                   preload.profiles="")
```

vocab.file

sequence 1 input sequence 1, character string (usually length of kmer or a window over a

variant)

sequence 2 input sequence 2, character string, required to have same length as sequence 1

kl length of the kmer windows, to split up the sequences

damage.mode {"exhaustive", "local"} mode to calculate the total damage for the comparison.

"exhaustive" sum over all pair-wise comparison. "local" extract the highest

pair-wise damage score. [default="exhaustive"]

tissue name of tissue directory of interest

data.dir path to directory containing preprocessed data

pnorm.tag tag indicating which background propensities were used for normalisation vocab.flag {FALSE, TRUE} flag if a precalculated vocabulary file is present and should

be used [default=FALSE]

Speeds up the analysis but precalculation is time consuming, therefore has to be done manually after initial processing of new tissue data. Recommended if

tissue is frequently used for analysis. path to precalculated vocabulary file

[default=paste0(data.dir,"",tissue,"vocabulary\_",tissue,".txt")]

frag.type {"DNase", "ATAC"} fragmentation type.

plots {FALSE, "highest", "all"} indicate if and what overlay plots to retrieve [de-

fault="highest"]

FALSE no plots, highest only the highest scoring pair, "all" plots for all pairs

smooth {FALSE, TRUE} smooth the profile [default=TRUE]

ylim y-limit for plots [default c(0,0.01)] xlim x-limit for plots [default c(-125,125)]

plot.shoulders {FALSE, TRUE} print the estimated shoulders with the overlay plots [de-

fault=FALSE]

preload flag [FALSE/TRUE] if to use preloaded average profiles [default=FALSE]

preload.vocab preloaded vocabulary data frame from PreLoadVocab()

preload.profiles two dataframe list with the preloaded profiles from PreLoadProfiles()

# Value

Returns a list containing two data frames and a plot or list of plots, depending on how the plotting was specified.

...\$summary: a data frame listing the reference and variant sequence with highest scoring kmer pair ...\$df: data frame containing the pair-wise results per row (kmer and SFR from reference and variant and the damage)

...\$plot: Overlay plot(s), either single ggplot2 plot or list of plots depending on mode selected for "plots"

```
frag.type="DNase",
   plots="highest"
> #With preload
> vocabulary <- PreLoadVocab(data.dir, tissue)</pre>
> comp <- CompareSequences(</pre>
    sequence1="CAGTTTCATGAGG",
    sequence2="CAGTTTTATGAGG",
   k1=7,
   data.dir=data.dir,
   pnorm.tag=pnorm.tag,
   damage.mode="exhaustive",
   tissue=tissue,
   vocab.flag=TRUE,
   frag.type="DNase",
   plots="highest",
   preload=TRUE,
   preload.vocab=vocabulary)
```

# DecodeKmer

# Description

Takes a kmer and dissects all ambivalent FASTA characters to create a character vector of all matching kmers.

# Usage

> DecodeKmer(kmer)

# Arguments

kmer character string, allowing all FASTA standard and ambivalent characters

# Value

Returns a vector containing of all matching definite kmers as character strings.

# Examples

> DecodeKmer("WGATAA")

# DissectSequence

# Description

Split an input sequence into kmer windows.

### Usage

> DissectSequence(sequence, kl, list=FALSE)

sequence character string, allowing all FASTA standard and ambivalent characters

kl length in bp to split the sequence into

list {FALSE, TRUE} select if output should be a list or a vector [default=FALSE]

### Value

Returns a vector or list of all kmers.

# Examples

> DissectList("AGGGATACGTAGACGGTGTAA", kl=7, list=FALSE)

# **GetCount**

# Description

Wrapper function to get the count of the k-mer in DHS in the tissue of interest.

### Usage

```
> GetCount(kmer, tissue, data.dir, pnorm.tag, frag.type)
```

# Arguments

**kmer** input kmer, FASTA characters string 5 - 7 bp

tissue name of tissue directory of interest

data.dir path to directory containing preprocessed data

**pnorm.tag** tag indicating which background propensities were used for normalisation

frag.type {"DNase", "ATAC"} fragmentation type.

# Value

Returns the count as single numeric value.

# Examples

# **GetFootprint**

# Description

Wrapper to retrieve the merged & smoothed profile of kmer.

# Usage

```
> GetFootprint(kmer,
+ tissue,
+ data.dir,
+ pnorm.tag,
+ frag.type,
+ smooth=TRUE,
+ smooth.bandwidth=5,
+ preload=FALSE,
+ preload.profiles="")
```

### Arguments

**kmer** input kmer, FASTA characters string 5 - 7 bp

tissue name of tissue directory of interest

data.dir path to directory containing preprocessed data

pnorm.tag tag indicating which background propensities were used for normalisation

frag.type {"DNase", "ATAC"} fragmentation type.

smooth {FALSE, TRUE} smooth the profile [default=TRUE]

**smooth.bandwidth** bandwidth to smooth in bp [default=5]

preload flag: [FALSE/TRUE] if to use preloaded average profiles [default=FALSE] preload.profiles two dataframe list with the preloaded profiles from PreLoadProfiles()

### Value

Returns list containing the profile and counts. ...\$profile, ...\$count.

# Examples

```
> frag.type <- "DNase"
> data.dir <- "./my_sasq_database/human/DNase/"</pre>
> tissue <- "blood_tissue"</pre>
> pnorm.tag <- "h_ery_1"</pre>
> fp <- GetFootprint(kmer="CACGTG",
                      tissue=tissue,
                      pnorm.tag=pnorm.tag,
                      data.dir=data.dir,
                      frag.type=frag.type,
                      smooth=TRUE)
> # With preloading
> profiles <- PreLoadKmerProfiles(7, data.dir, tissue, pnorm.tag)
> fp <- GetFootprint(kmer="CACGTG",
                      tissue=tissue,
                      data.dir=data.dir,
                      pnorm.tag=pnorm.tag,
                      frag.type=frag.type,
                      smooth=TRUE,
                      preload=TRUE,
                      preload.profiles=profiles)
```

# GetFootprintStrand

# Description

Wrapper to retrieve strand-specific (smoothed) profiles of a kmer from tissue or background

### Usage

```
> GetFootprintStrand(kmer,
                      tissue,
                      data.dir,
                      pnorm.tag,
                      frag.type,
                      smooth=TRUE,
                      smooth.bandwidth=5,
                      background.flag=FALSE)
```

### Arguments

kmer input kmer, FASTA characters string 5 - 7 bp tissue name of tissue directory of interest path to directory containing preprocessed data data.dir tag indicating which background propensities were used for normalisation pnorm.tag {"DNase", "ATAC"} fragmentation type. frag.type {FALSE, TRUE} smooth the profile [default=TRUE] smooth smooth.bandwidth bandwidth to smooth in bp [default=5] background.flag select if to retrieve the profile from the genome-wide background [de-

fault=FALSE]

# Value

Returns list with strand-specific profiles and counts. ...\$profile.plus, ...\$profile.minus, ...\$count.plus, ...\$count.minus

# Examples

```
> frag.type <- "DNase"</pre>
> data.dir <- "./my_sasq_database/human/DNase/"</pre>
> tissue <- "blood_tissue"
> pnorm.tag <- "h_ery_1"
  GetFootprintStrand(kmer="CACGTG",
                      tissue=tissue,
                      data.dir=data.dir,
                      pnorm.tag=pnorm.tag,
                      frag.type=frag.type,
                      smooth=TRUE,
                      background.flag=FALSE)
```

# GetPossibleMutations

#### Description

Take a sequence input and split into kmers of length kl \* 2 - 1 (e.g. for kl 7 always takes the 13 surrounding bases) and separate into reference and variance. Take the parsed position as index for the first base to mutate. List all possible mutations filling the kl'th position in the variant column with all possible substitutions depending on the reference base. First position of the seugence to be mutated is therefore the kl'th positoin of the sequence string.

# Usage

```
> GetPossibleMutations(sequence, kl=7, chr=".", position=1)
```

### Arguments

```
    sequence character string, allowing all FASTA standard and ambivalent characters
    kl length in bp to split the sequence into
```

chr chromosome of the sequence to print bp position bp position of the sequence to print

#### Value

Returns a 6 column data frame with "chr" "pos" "ref.base" "var.base" "ref.seq" "var.seq"

# Examples

```
> GetPossibleMutations(sequence=c("AGGGATACGTAGACGGTGTAA"),
+ kl=7, chr="chrX", position=1345990)
```

# GetSFR

# Description

Wrapper function to get the SFR ratio. If indicated and available, use the present vocabulary file to directly grep the SFR. Else get the average profile, estimate the borders and calculate the SFR. Note that for using the vocabulary file only nonambivlent DNA chars are allowed. For the alternative ambivalent chars are decoded.

If preload = TRUE and if vocab.flag = FALSE, preload.profiles has to be submitted. If vocab.flag = TRUE with preload = TRUE then preload.vocab has to be submitted.

**kmer** input kmer, FASTA characters string 5 - 7 bp

tissue name of tissue directory of interest

data.dir path to directory containing preprocessed data

pnorm.tag tag indicating which background propensities were used for normalisation vocab.flag {FALSE, TRUE} flag if a precalculated vocabulary file is present and should

be used [default=FALSE]

Speeds up the analysis but precalculation is time consuming, therefore has to be done manually after initial processing of new tissue data. Recommended if

tissue is frequently used for analysis.

vocab.file path to precalculated vocabulary file

[default=paste0(data.dir,"",tissue,"vocabulary\_",tissue,".txt")]

frag.type {"DNase", "ATAC"} fragmentation type.

preload flag [FALSE/TRUE] if to use preloaded average profiles [default=FALSE]

preload.vocab preloaded vocabulary data frame from PreLoadVocab()

preload.profiles two dataframe list with the preloaded profiles from PreLoadProfiles()

### Value

Returns SFR as sinlge numeric value.

### Examples

```
> GetSFR(kmer="CACGTG",
         tissue="blood_tissue",
+
         pnorm.tag="h_ery_1",
         data.dir="./my_sasq_database/human/DNase/",
         vocab.flag=TRUE,
         frag.type="DNase")
> # With preload
> data.dir <- "./my_sasq_database/human/DNase/"</pre>
> tissue <- "blood_tissue"
> vocabulary <- PreLoadVocab(data.dir, tissue)
> GetSFR(kmer="CACGTG",
         tissue=tissue,
         data.dir=data.dir,
         pnorm.tag="h_ery_1",
         vocab.flag=TRUE,
         frag.type="DNase",
         preload=TRUE,
         preload.vocab=vocabulary)
```

# GrepProfile

### Description

Grep strand-specific profile of 250 bp surrounding kmer and normalise for total cuts in 250 bp window.

```
> GrepProfile(kmer, infile, preload=FALSE, preload.profiles.strand="")
```

```
kmer input kmer, FASTA characters string 5 - 7 bp
```

**infile** path to the input file

preload flag: [FALSE/TRUE] if to use preloaded average profiles [de-

fault=FALSE]

preload.profiles.strand dataframe of preloaded profiles from a single strand [e.g. profiles\$plus]

#### Value

Returns list of ..\$profile and ..\$count.

# Examples

# **InSilicoMutation**

# Description

Wrapper for maximum/absolute damage insilico mutation. Takes an input sequence, splits into data frame of desried kmer length matching window sizes for running comparison and compares reference sequence against all possible mutated sequences (single base pair substitutions). Reports according to report mode ("all", "max", "maxabs").

Note that preloading data is highly recommended for speeding up.

(If preload = TRUE and if vocab.flag = FALSE, preload.profiles has to be submitted. If vocab.flag = TRUE with preload = TRUE then preload.vocab has to be submitted.)

```
> InSilicoMutation(sequence,
+ kl=7,
+ chr=".",
+ position=1,
+ report="all",
+ damage.mode="exhaustive",
+ tissue=tissue,
+ data.dir="./my_sasq_database/human/DNase/",
+ pnorm.tag,
+ vocab.flag=TRUE,
```

```
+ vocab.file=paste0(data.dir,"/",tissue,"/vocabulary_",tissue,".txt"),
```

- + frag.type=frag.type,
- progress.bar=FALSE,
- + preload=FALSE,
- + preload.vocab="",
- + preload.profiles="")

sequence1 input sequence, character string

kl length of the kmer windows, to split up the sequence

**chr** chromosome of the sequence to print

position bp position of the sequence to print, first mutated base is the kl'th base. There-

fore input +- 6 bp positions of sequence surrounding your sequence of interest

and set postion to the kl'th index.

report {"all", "max", "maxabs"} Select which damage per position to report. [de-

fault="all"]

"all" = report all 3 possible substitutions per position. Reports three rows per

bp position.

"max" = only report substitution with highest positive damage. Report one

row per bp position, easy to convert to wig.

"maxabs" = only report substitution with highest absolute damage. Reports

one row per bp positon as well.

 $\mathbf{damage.mode} \qquad \{\text{"exhaustive"},\,\text{"local"}\} \; \text{mode to calculate the total damage for the comparison}.$ 

"exhaustive" sum over all pair-wise comparison. "local" extract the highest

pair-wise damage score. [default="exhaustive"]

tissue name of tissue directory of interest

data.dir path to directory containing preprocessed data

pnorm.tag tag indicating which background propensities were used for normalisation vocab.flag {FALSE, TRUE} flag if a precalculated vocabulary file is present and should

be used [default=FALSE]

Speeds up the analysis but precalculation is time consuming, therefore has to be done manually after initial processing of new tissue data. Recommended if

tissue is frequently used for analysis.  $\,$ 

vocab.file path to precalculated vocabulary file

[default=paste0(data.dir,"",tissue,"vocabulary\_",tissue,".txt")]

frag.type {"DNase", "ATAC"} fragmentation type.

progress.bar {FALSE, TRUE} Select if to display a progress.bar when running. (Requires

packages phapply if set to TRUE!) [default=FALSE]

preload flag [FALSE/TRUE] if to use preloaded average profiles [default=FALSE]

preload.vocab
preloaded vocabulary data frame from PreLoadVocab()

preload.profiles two dataframe list with the preloaded profiles from PreLoadProfiles()

#### Value

Seven columns dataframe c(chr, position, ref.base, var.base, ref.sequence, var.sequence, damage).

```
> # use (for example) the BSgenome package to extract reference genome sequence
```

- > library(BSgenome)
- > library(BSgenome.Hsapiens.UCSC.hg18)
- > genome <- BSgenome.Hsapiens.UCSC.hg18
- > #set the sequence coordinates of the desired genomic location
- > chr <- "chr16"
- > start.pos <- 145852

```
> end.pos <- start.pos + 30
> # Get the sequence sequence
> seq <- as.character(getSeq(genome, "chr16", start=start.pos-6, end=end.pos+6))
> data.dir <- "./my_sasq_database/human/DNase/"</pre>
> tissue <- "blood_tissue"</pre>
> vocabulary <- PreLoadVocab(data.dir, tissue)</pre>
> #perform the in silcio mutation
> df.insilico <- InSilicoMutation(sequence=seq,</pre>
                                     chr="chr16",
                                     position=start.pos,
                                     report="all",
                                     damage.mode="exhaustive",
                                     tissue=tissue,
                                     pnorm.tag="h_ery_1",
                                     data.dir=data.dir,
                                     vocab.flag=TRUE,
                                     frag.type=frag.type,
                                     progress.bar = TRUE,
                                     preload=TRUE,
                                     preload.vocab=vocabulary
> #display as InSilicoMutationplot
> rp <- InSilicoMutationPlot(df.insilico, ylim=c(-4,4))</pre>
```

# **InSilicoMutationPlot**

### Description

Make a InSilicoMutation plot from data frame as output from InSilicoMutation. Must have been run with report="all"

### Usage

```
> InSilicoMutationPlot(df, ylim=c(-2,2))
```

#### Arguments

```
df data frame as output from InSilicoMutation (report="all") ylim y-limits for plot[default=c(-2,2)]
```

#### Value

Returns ggplot2 plot object containing the InSilicoMutationplot.

```
> # use (for example) the BSgenome package to extract reference genome sequence
> library(BSgenome)
> library(BSgenome.Hsapiens.UCSC.hg18)
> genome <- BSgenome.Hsapiens.UCSC.hg18
> #set the sequence coordinates of the desired genomic location
> chr <- "chr16"
> start.pos <- 145852</pre>
```

```
> end.pos <- start.pos + 30
> # Get the sequence sequence
> seq <- as.character(getSeq(genome, "chr16", start=start.pos-6, end=end.pos+6))
> #perform the i silcio mutation
> df.insilico <- InSilicoMutation(sequence=seq,</pre>
                                    k1=7,
                                    chr="chr16",
                                    position=start.pos,
                                    report="all",
                                    damage.mode="exhaustive",
                                    tissue="blood_tissue",
                                    data.dir="./my_sasq_database/human/DNase/",
                                    vocab.flag=TRUE,
                                    frag.type=frag.type,
                                    progress.bar = TRUE
> #display as InSilicoMutationplot
> rp <- InSilicoMutationPlot(df.insilico, ylim=c(-4,4))
```

# ${\bf Make In Silico Mutation Track Hub}$

# Description

Wrapper to make a UCSC browser track hub from the insilico mutation data frame. Input is a data frame in the same format as the output data frame of the InSilicoMutation function. 7 columns: chr pos ref.base var.base ref.seq var.seq damage

input.df input 7 column data frame. chr pos ref.base var.base ref.seq var.seq

damage

id.tag id tag to name the hub directory and bw tracks.
store.tracks path to directory to store the bigwig tracks.
store.hub were to store the hub & visualization folder.

genome.build select genome build ("hg19", "hg18", "mm9", ...) [default="hg19"]
path.chr.sizes full.path to chrsizes file matching to the selected genome. Not pro-

vided in package.

short.label shortLabel for track hub.

long.label longLabel for track hub [default = short.label]

set.email email contact address to appear in trackHub. [default="none"] bedgraph.to.bigwig.path full path to UCSC bedGraphtoBigWig convertion tool. Not in-

cluded.

make.softlinks {FALSE, TRUE} set flag if to directly make softlinks in hub folder

[default=FALSE]

(e.g. set TRUE if running directly on a cluster so that the final

softlinks paths are already correct

[default = FALSE] if data files will be copied to a different direction

afterwards (e.g. when mounted and ran locally)

if FALSE: After creation copy data hub to desired location and create softlinks in the hub folder to the tracks in the track folder e.g. "In

-s store.tracks/\*.bw store.hub"

#### Value

Writes bigWig tracks into desired directory and creates a trackHub structure to migrate to public domain and import to UCSC.

```
> frag.type <- "DNase"
> data.dir <- "./my_sasq_database/human/DNase/"</pre>
> tissue <- "blood_tissue"</pre>
> # use (for example) the BSgenome package to extract reference genome sequence
> library(BSgenome)
> library(BSgenome.Hsapiens.UCSC.hg18)
> genome <- BSgenome. Hsapiens. UCSC. hg18
> #set the sequence coordinates of the desired genomic location
> chr <- "chr16"
> start.pos <- 145852
> end.pos <- start.pos + 30
> # Get the sequence sequence
> seq <- as.character(getSeq(genome, "chr16", start=start.pos-6, end=end.pos+6))
> #perform the i silcio mutation
> df.insilico <- InSilicoMutation(sequence=seq,</pre>
                                    k1=7,
                                    chr="chr16",
                                    position=start.pos,
                                    report="all",
                                    damage.mode="exhaustive",
                                    tissue="blood_tissue",
                                    data.dir=data.dir,
                                    vocab.flag=TRUE,
                                    frag.type=frag.type,
```

```
progress.bar = TRUE

prog
```

# PlotOverlap

# Description

Plot two average profiles overlapping or on top of each other given two input profiles. Plot the shoulders if plot.shoulders is set to TRUE, then use shoulders list as provided or determine.

```
profile1
                  input profile1, retrieved from the GetFootprint() function as list entry ...$profile
profile2
                  input profile2, retrieved from the GetFootprint() function as list entry ...$profile
kmer1
                  input k-mer 1 (reference)
kmer2
                  input k-mer 2 (variant)
count1
                  count of k-mer1 occurence
                  count of k-mer2 occurence
count2
ymode
                  mode how to plot the overlapping profiles ("merged" or as "separate" profiles
                  above each other) [default=separate]
ylim
                  ylim to fix for plot [default c(0, 0.01)]
                  xlim to fix for plot [default c(-125, 125)]
xlim
                  FALSE, TRUE if to plot the estimated shoulders with the profiles [de-
plot.shoulders
                  fault=FALSE]
                  note that it is only plotted if the separate profile option was selected to keep the
                  plots tidy
shoulders1
                  list object of estiamted shoulder postion and ranges for profile 1 [default=FALSE]
shoulders2
                  list object of estiamted shoulder postion and ranges for profile 2 [default=FALSE]
```

### Value

Returns ggplot2 plot object of the overlay plot.

# Examples

```
> frag.type <- "DNase"
> data.dir <- "./my_sasq_database/human/DNase/"</pre>
> tissue <- "blood_tissue"
> kmer1 <- "WGATAA" #note FASTA ambigous code is supported
 kmer2 <- "WGATTA"
  fp1 <- GetFootprint(kmer=kmer1,</pre>
                       tissue=tissue,
                       data.dir=data.dir,
                       frag.type=frag.type,
                       smooth=TRUE)
 fp2 <- GetFootprint(kmer=kmer2,</pre>
                       tissue=tissue,
                       data.dir=data.dir,
                       frag.type=frag.type,
                       smooth=TRUE)
> # make an overlap plot
> PlotOverlap(
    fp1$profile,
    fp2$profile,
    kmer1,
    kmer2,
    fp1$count,
    fp2$count,
    ymode="separate"
```

# **PlotOverlapKmers**

# Description

Wrapper function to produce an overlay plot from two kmers and a tissue input only. (For plotting: If preload = TRUE preload.profiles has to be submitted.)

# Usage

# Arguments

${ m kmer1}$	input k-mer 1 (reference)		
kmer2	input k-mer 2 (variant)		
${f tissue 1}$	name of tissue 1 directory of interest		
${f tissue 2}$	2 name of tissue 2 directory of interest		
${ m data.dir}$	r path to directory containing preprocessed data		
${f pnorm.tag}$	m.tag tag indicating which background propensities were used for normalisation		
frag.type	{"DNase", "ATAC"} fragmentation type.		
$\mathbf{smooth}$	{FALSE TRUE} if to smooth the profiles. [default=TRUE]		
$\mathbf{y}\mathbf{m}\mathbf{o}\mathbf{d}\mathbf{e}$	mode how to plot the overlapping profiles ("merged" or as "separate" profiles		
	above each other) [default=separate]		
$\mathbf{ylim}$	ylim to fix for plot [default $c(0, 0.01)$ )]		
xlim	xlim to fix for plot [default $c(-125, 125)$ ]		
plot.shoulders	{FALSE, TRUE} if to plot the estimated shoulders with the profiles [de-		
	fault=FALSE]		
	note that it is only plotted if the separate profile option was selected to keep		
	the plots tidy		
preload	flag [FALSE/TRUE] if to use preloaded average profiles [default=FALSE]		
preload.profiles	two dataframe list with the preloaded profiles from PreLoadProfiles()		

# Value

Returns ggplot2 plot object of the overlay plot.

```
> frag.type <- "DNase"
> data.dir <- "./my_sasq_database/human/DNase/"
> tissue <- "blood_tissue"
> PlotOverlapKmers(
+ kmer1="CACGTG",
+ kmer2="CACGTT",
+ tissue1=tissue,
+ issue2=tissue,
```

```
+ data.dir=data.dir,
+ pnorm.tag="h_ery_1",
+ frag.type="DNase",
+ smooth=TRUE,
+ plot.shoulders=FALSE
+ )
```

# **PlotSingle**

# Description

Plot the average profile plot given an input profile. Plot the shoulders if plot.shoulders is set to TRUE and use shoulders list provided or determine.

### Usage

```
> PlotSingle(profile,
+ kl=7,
+ plot.shoulders=FALSE,
+ shoulders=FALSE,
+ ylim=c(0,0.01),
+ xlim=c(-125,125),
+ color="black")
```

# Arguments

profile	input profile				
kl	length of the kmer windows, to split up the sequences				
$\mathbf{ylim}$	ylim to fix for plot $[default=c(0, 0.01))]$				
xlim	xlim to fix for plot $[default=c(-125, 125)]$				
plot.shoulders	{FALSE, TRUE} if to plot the estimated shoulders with the profiles [de-				
	fault=FALSE]				
shoulders	list object of estiamted shoulder postion and ranges for profile 1 [default=FALSE]				
color	Select a color for the profile [default="black"]				

# Value

Returns single ggplot2 plot object of the profile.

```
plot.shoulders=TRUE,
shoulders=sh
)
```

# PlotSingleKmer

# Description

Wrapper function to produce a plot from kmer and tissue input only. (For plotting: If preload = TRUE preload.profiles has to be submitted.)

# Usage

```
> PlotSingleKmer(kmer,
+ tissue,
+ data.dir,
+ frag.type,
+ pnorm.tag,
+ smooth=TRUE,
+ smooth.bandwidth=5,
+ plot.shoulders=FALSE,
+ ylim=c(0,0.01),
+ xlim=c(-125,125),
- color="black",
- preload=FALSE,
+ preload.profiles)
```

### Arguments

kmerinput k-mertissuename of tissue 1 directory of interestdata.dirpath to directory containing preprocessed datapnorm.tagtag indicating which background propensities were used for normalisationfrag.type{"DNase", "ATAC"} fragmentation type.smooth{FALSE TRUE} if to smooth the profiles. [default=TRUE]smooth.bandwidthbandwidth to smooth in bp [default=5]

plot.shoulders {FALSE, TRUE} if to plot the estimated shoulders with the profiles [de-

fault=FALSE]

ylim ylim to fix for plot [default c(0, 0.01))]
xlim xlim to fix for plot [default c(-125, 125)]
color Select a color for the profile [default="black"]

preload flag [FALSE/TRUE] if to use preloaded average profiles [default=FALSE] preload.profiles two dataframe list with the preloaded profiles from PreLoadProfiles()

### Value

Returns single ggplot2 plot object of the profile.

```
> PlotSingleKmer(kmer="CACGTG",
+ tissue="blood_tissue",
```

```
+ data.dir="./my_sasq_database/human/DNase/",
+ frag.type="DNase")
```

# PlotSingleStrands

# Description

Wrapper function to produce strand specific plots from kmer and tissue input only.

# Usage

```
> PlotSingleStrands(kmer,
+ tissue,
+ data.dir,
+ pnorm.tag,
+ frag.type,
+ smooth=TRUE,
+ smooth.bandwidth=5,
+ background.flag=FALSE,
+ ylim=c(0,0.01),
+ xlim=c(-125,125))
```

# Arguments

**kmer** input k-mer

tissue name of tissue 1 directory of interest

data.dir path to directory containing preprocessed data

pnorm.tag tag indicating which background propensities were used for normalisation

frag.type {"DNase", "ATAC"} fragmentation type.

smooth {FALSE TRUE} if to smooth the profiles. [default=TRUE]

smooth.bandwidth

bandwidth to smooth in bp [default=5]

background.flag FALSE TRUE Select if to visualize the genome-wide background cut pro-

files or from tissue data. Will adjust the required and queried data structure

accordingly.

Note that if set to TRUE, a different data directory, that pointing to the storange of the deproteinized, genome-wide background has to be provied.

[default=FALSE]

ylim ylim to fix for plot [default c(0, 0.01))] xlim xlim to fix for plot [default c(-125, 125)]

### Value

Returns list of strand-specific profile plots: for plus ...\$plot.plus and minus strand ...\$plot.minus.

```
> PlotSingleStrands(kmer="WGATAA",
+ tissue = "blood_tissue",
+ data.dir = "./my_sasq_database/human/DNase/",
+ pnorm.tag = "h_ery_1",
+ frag.type = "DNase")
```

# **PreLoadProfiles**

# Description

Preload the entire kmer based average profiles for a given tissue and kmer size.

# Usage

```
> PreLoadKmerProfiles(kl, data.dir, tissue, pnorm.tag)
```

### Arguments

kl length of the kmer windows, to split up the sequences

tissue name of tissue 1 directory of interest

data.dir path to directory containing preprocessed data

pnorm.tag tag indicating which background propensities were used for normalisation

### Value

List of legnth two ...\$plus and ...\$minus storing the occurrences and cut profiles per k-mer. Each list entry is a data frame with the following dimension:

Column[1] = kmer, Column[2] = kmer occurrence, Column[3 - (300+kl+2)] = 300 bp + kl kmer surrounding profile. Rows = number of possible kmers of length kl

Note that we store the 300 kmer surrounding bases but only query and process the surrounding 250 bp window. We kept the stroing at 300 bp to be quickly able to expand our scripts. All functions process the described 250 surrounding window from this start files.

### Examples

```
> kl <- 7
> data.dir <- "./my_sasq_database/human/DNase/"
> tissue <- "blood_tissue"
> pnorm.tag <- "h_ery_1"
> profiles <- PreLoadProfiles(data.dir, tissue)</pre>
```

# PreLoadVocab

### Description

Preload the entire tissue specific vocabulary file with preprocessed SFRs for every possible k-mer.

### Usage

```
> PreLoadVocab(data.dir, tissue)
```

# Arguments

```
tissue name of tissue 1 directory of interest
```

data.dir path to directory containing preprocessed data

#### Value

Vocabulary file loaded as two column data frame.

### Examples

```
> data.dir <- "./my_sasq_database/human/DNase/"
> tissue <- "blood_tissue"
> vocabulary <- PreLoadVocab(data.dir, tissue)</pre>
```

# **PruneProfile**

# Description

Prune an retrieved (250 bp) average profile equally from both directions given the profile and the desired window size around the kmer which to retrieve. Requires an even number as length to prune. Length of the output profile will always be (desired.length + kmer.length).

# Usage

```
> PruneProfile(profile, desired.length)
```

# Arguments

```
profile input profiledesired.length length of to prune to (will be desried.length + kmer.length)
```

### Value

Returns pruned profile as numeric vector.

# Examples

# QueryJaspar

# Description

Take a sequence input and query it against a set of Jaspar 2014 PWMs. (as provided or saved from the JASPAR 2014 R package). Requires "Biostrings" and "TFBSTools" packages.

#### Usage

> QueryJaspar(sequence, threshold=0.8, pwm.data)

sequence input sequence

threshold relative percentage score threshold above which to report matches [default=0.8] pwm.data a stored pwm.RData object as provided with the Sasquatch distriution or as retrieved

and saved from JASPAR2014 R package

#### Value

Returns character string listing the PWM matches above the relative percentage score threshold.

### Examples

```
> #requries Biostrings and TFBSTools R packages
> library(Biostrings)
> library(TFBSTools)
> #load human.pwm object
> load("./my_sasq_database/jaspar/jaspar2014.human.9606.all.versions")
> # Single JASPAR query
> QueryJaspar(sequence="AGATAATAG", threshold=0.8, pwm.data=human.pwm)
```

# QueryJasparBatch

# Description

Take a data frame from the RefVarBatch query as input and query it against the set of selected Jaspar2014 PWMs using a selected match.threshold. Select an absolute footprinting damage above which to query jaspar.

#### Usage

> QueryJasparBatch(df, damage.threshold=0, match.threshold=0.8, pwm.data)

# Arguments

df 9 column data frame as output from RefVarBatch()

damage.threshold absolute predicted footprinting damage above which a sequence should be

selected for the query. [default=0, query all]

match.threshold relative percentage score threshold above which to report matches [de-

fault=0.8

**pwm.data** a stored pwm.RData object as provided with the Sasquatch distriution or

as retrieved and saved from JASPAR2014 R package

### Value

Returns data frame with additional column for jaspar query results.

```
> #make a reference vs. variant batch query
 comp.df <- RefVarBatch(ref.var.df=tdf,</pre>
                        k1=7,
                        damage.mode="exhaustive",
                        tissue="blood_tissue",
                        data.dir="./my_sasq_database/human/DNase/",
                        vocab.flag=TRUE,
                        frag.type="DNase")
> #requries Biostrings and TFBSTools R packages
> library(Biostrings)
> library(TFBSTools)
> #load human.pwm object
> load("./my_sasq_database/jaspar/jaspar2014.human.9606.all.versions")
> #query refvar data frame against jaspar pwms
> comp.df.jaspar <- QueryJasparBatch(df=comp.df,</pre>
                                      damage.threshold=0.3,
                                      match.threshold=0.8,
                                      pwm.data=human.pwm)
```

# QueryLongSequence

# Description

Wrapper function to split a longer sequence into kmers of length kl and return kmer, SFR and plots if specified. (If preload = TRUE and if vocab.flag = FALSE, preload.profiles has to be submitted. If vocab.flag = TRUE with preload = TRUE then preload.vocab has to be submitted.)

```
> QueryLongSequence(sequence,
                    kl.
                    tissue,
                    data.dir,
                    pnorm.tag,
                    vocab.flag=FALSE,
                     vocab.file=paste0(data.dir,"/",tissue,
                                       "/vocabulary_",tissue,".txt"),
                    frag.type,
                    plots=FALSE,
                    smooth=TRUE,
                    plot.shoulders=TRUE,
                    ylim=c(0,0.01),
                    xlim=c(-125,125),
                    preload=FALSE,
                    preload.vocab="",
                    preload.profiles="")
```

sequence input sequence, character string length of the kmer windows, to split up the sequence  $\mathbf{kl}$ tissue name of tissue directory of interest data.dir path to directory containing preprocessed data pnorm.tag tag indicating which background propensities were used for normalisation vocab.flag {FALSE, TRUE} flag if a precalculated vocabulary file is present and should be used [default=FALSE] Speeds up the analysis but precalculation is time consuming, therefore has to be done manually after initial processing of new tissue data. Recommended if tissue is frequently used for analysis. vocab.file path to precalculated vocabulary file [default=paste0(data.dir,"",tissue,"vocabulary\_",tissue,".txt")] frag.type {"DNase", "ATAC"} fragmentation type. plots {FALSE, TRUE} indicate if to plot a profile per kmer [default=FALSE] {FALSE, TRUE} smooth the profile [default=TRUE]  $\mathbf{smooth}$ {FALSE, TRUE} print the estimated shoulders with the plots [default=TRUE] plot.shoulders vlim v-limit for plots [default c(0.0.01)] xlim x-limit for plots [default c(-125,125)] preload flag [FALSE/TRUE] if to use preloaded average profiles [default=FALSE] preload.vocab preloaded vocabulary data frame from PreLoadVocab() preload.profiles two dataframe list with the preloaded profiles from PreLoadProfiles()

#### Value

Returns ...\$df a data frame listing the splitted kmers with the respective SFR. ...\$plots if specified list of profile plots with one plot per splitted k-mer.

```
> QueryLongSequence(sequence="ATAGATAATCGCT",
                     tissue="blood_tissue",
                     pnorm.tag="h_ery_1",
                     data.dir="./my_sasq_database/human/DNase/",
                     vocab.flag=FALSE,
                     vocab.file=paste0(data.dir, "/", tissue,
                                        "/vocabulary_",tissue,".txt"),
                     frag.type="DNase",
                     plots=FALSE)
> # With preload
> data.dir <- "./my_sasq_database/human/DNase/"</pre>
 tissue <- "blood_tissue"
 vocabulary <- PreLoadVocab(data.dir, tissue)</pre>
  QueryLongSequence(sequence="ATAGATAATCGCT",
                     kl,
                     tissue=tissue,
                     pnorm.tag="h_ery_1",
                     data.dir=data.dir,
                     vocab.flag=TRUE,
                     vocab.file=paste0(data.dir, "/", tissue,
                                        "/vocabulary_",tissue,".txt"),
                     frag.type="DNase",
                     plots=FALSE,
```

```
+ preload=TRUE,
```

+ preload.vocab=vocabulary)

### RefVarBatch

# Description

Wrapper function to analyse multiple Ref-Var-Sequence pairs. Split each sequence into kmers of length kl, get their SFRs (ideally for speed from vocab.file or calculate per instance. Calculate the damage associated with each kmer pair and from that the local or the exhaustive summed up damage of entire sequence pair.

(If preload = TRUE and if vocab.flag = FALSE, preload.profiles has to be submitted. If vocab.flag = TRUE with preload = TRUE then preload.vocab has to be submitted.)

### Usage

```
> RefVarBatch(ref.var.df,
+ kl,
+ damage.mode="exhaustive",
+ tissue,
+ data.dir,
+ pnorm.tag,
+ vocab.flag=FALSE,
+ vocab.file=pasteO(data.dir,"/",tissue,"/vocabulary_",tissue,".txt"),
+ frag.type,
+ preload=FALSE,
+ preload.profiles="",
+ preload.vocab="")
```

# Arguments

C 1C	,1 1 1,	C 1 1	C 1	•	/ · 1 C
ret.var.dt	three column data	tromo lieting id	rotoroneo and	verience cognones	lid rotoronco
i ci. vai .ui	tinee column data	mame noung id	reference and	. variance secuence	i id reference

variant)

kl length of the kmer windows, to split up the sequences

damage.mode {"exhaustive", "local"} mode to calculate the total damage for the comparison.

"exhaustive" sum over all pair-wise comparison. "local" extract the highest

pair-wise damage score. [default="exhaustive"]

tissue name of tissue directory of interest

data.dir path to directory containing preprocessed data

pnorm.tag tag indicating which background propensities were used for normalisation vocab.flag {FALSE, TRUE} flag if a precalculated vocabulary file is present and should

be used [default=FALSE]

Speeds up the analysis but precalculation is time consuming, therefore has to be done manually after initial processing of new tissue data. Recommended if

tissue is frequently used for analysis.

vocab.file path to precalculated vocabulary file

 $[{\tt default=paste0}({\tt data.dir,"",tissue,"vocabulary\_",tissue,".txt"})]$ 

frag.type {"DNase", "ATAC"} fragmentation type.

preload flag [FALSE/TRUE] if to use preloaded average profiles [default=FALSE]

preload.vocab preloaded vocabulary data frame from PreLoadVocab()

preload.profiles two dataframe list with the preloaded profiles from PreLoadProfiles()

### Value

Returns data frame listing Ref and Var sequence with highest scoring kmer pair and according SFRs and calculated (exhaustive or local) total damage.

# Examples

```
> #ty example data frame
> tdf <- data.frame(</pre>
          id=c("1", "2", "3"),
          ref=c("ATAGATAATCGCT", "ATAGATAATCGCT", "ATATATTCTCGCT"),
          var=c("ATAGATCATCGCT", "ATAGATTATCGCT", "ATAGATGATCGCT")
> # Make a reference vs. variant batch query
  comp.df <- RefVarBatch(ref.var.df=tdf,</pre>
                        k1=7,
                        damage.mode="exhaustive",
                        tissue="blood_tissue",
                        data.dir="./my_sasq_database/human/DNase/",
                        pnorm.tag="h_ery_1",
                        vocab.flag=TRUE,
                        frag.type="DNase")
> # With Preload
> data.dir <- "./my_sasq_database/human/DNase/"</pre>
> tissue <- "blood_tissue"
> vocabulary <- PreLoadVocab(data.dir, tissue)</pre>
  comp.df <- RefVarBatch(ref.var.df=tdf,</pre>
                        k1=7,
                        damage.mode="exhaustive",
                        tissue="blood_tissue",
                        data.dir="./my_sasq_database/human/DNase/",
                        pnorm.tag="h_ery_1",
                        vocab.flag=TRUE,
                        frag.type="DNase",
                        preload=TRUE,
                        preload.vocab=vocabulary)
```

# **SmoothProfile**

### Description

Helper function to smooth a profile given the specified badnwidth and a gaussian, normal kernel.

### Usage

```
> SmoothProfile(profile, bandwidth=5)
```

# Arguments

```
profile input profile (numeric vector)
smooth.bandwidth bandwidth to smooth in bp [default=5]
```

#### Value

Returns smoothed profile as numeric vector.

# Examples

# **SobelBorders**

# Description

Estimate the footprint shoulders by based on zero crossings of the 1D 1st derivative approximation of a smoothed profile and get the optimal shoulder range that maximizies the SFRatio of the footprint. (Approximations for speeding up the process are: Estimate the optimal positions based on 6 bp wide shoulders. Then opimize the shoulder width/range with allowed ranges in  $\{4,6,8,10\}$ )

# Usage

```
> SobelBorders(profile, kl)
```

# Arguments

```
    profile input profile (numeric vector) has to be smoothed for proper function!
    kl {5,6,7} kmer length input
```

### Value

Returns list object containing the shoulder centric positions and their ranges.

- ..\$us upstream shoulder center
- ..\$ds downstream shoulder center
- ..\$us.range width/range of upstream shoulder
- ..\$ds.range width/range of downstream shoulder
- ..\$flag flag {TRUE FALSE} if shoulders could be estimated

### Examples

# Sobeln

# Description

Helper function to calculate 1D 1st derivative approximation of profile by 1D sobel filtering.

# Usage

```
> Sobeln(profile)
```

# Arguments

```
profile input profile (numeric vector) should be smoothed!
```

# Value

Returns 1D 1st derivative approimation of the profile as numeric vector.

```
> fp <- GetFootprint(kmer="CACGTG",
+ tissue="blood_tissue",
+ data.dir="./my_sasq_database/human/DNase/",
+ frag.type="DNase",
+ smooth=TRUE)
> # estimate the shoulders from the profile
> # (use smoothed profile or smooth within call!)
> approx.1st.deriv.prof<- Sobeln(fp$profile)</pre>
```