

Package ‘MouseFM’

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Type Package

Title In-silico methods for genetic finemapping in inbred mice

Version 1.8.0

Description This package provides methods for genetic finemapping in inbred mice by taking advantage of their very high homozygosity rate (>95%).

Encoding UTF-8

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BugReports <https://github.com/matmu/MouseFM/issues>

Depends R (>= 4.0.0)

License GPL-3

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Author Matthias Munz [aut, cre] (<<https://orcid.org/0000-0002-4728-3357>>), Inken Wohlers [aut] (<<https://orcid.org/0000-0003-4004-0464>>), Hauke Busch [aut] (<<https://orcid.org/0000-0003-4763-4521>>)

Maintainer Matthias Munz <matthias.munz@gmx.de>

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annotate_consequences *Annotate with consequences*

Description

Request variant consequences from Variant Effect Predictor (VEP) via Ensembl Rest Service. Not recommended for large queries.

Usage

```
annotate_consequences(geno, species)
```

Arguments

| | |
|---------|---|
| geno | Data frame or GenomicRanges::GRanges object including columns rsid, ref, alt. |
| species | Species name, e.g. mouse (GRCm38) or human (GRCh38). |

Value

Data frame.

Examples

```
geno = finemap("chr1",
  start = 5000000, end = 6000000,
  strain1 = c("C57BL_6J"), strain2 = c("AKR_J", "A_J", "BALB_cJ")
)

df = annotate_consequences(geno[seq_len(10), ], "mouse")

geno.granges = finemap("chr1",
  start = 5000000, end = 6000000,
  strain1 = c("C57BL_6J"), strain2 = c("AKR_J", "A_J", "BALB_cJ"),
  return_obj = "granges"
)

df2 = annotate_consequences(geno.granges[seq_len(10), ], "mouse")
```

annotate_mouse_genes *Annotate with genes*

Description

Request mouse genes from Ensembl Biomart.

Usage

```
annotate_mouse_genes(geno, flanking = NULL)
```

Arguments

| | |
|----------|---|
| geno | Data frame or GenomicRanges::GRanges object including columns chr, pos. |
| flanking | Size of flanking sequence to be included. |

Value

Data frame.

Examples

```
geno = finemap("chr1",
  start = 5000000, end = 6000000,
  strain1 = c("C57BL_6J"), strain2 = c("AKR_J", "A_J", "BALB_cJ")
)

genes = annotate_mouse_genes(geno, 50000)
```

avail_chromosomes *Available chromosomes*

Description

Available mouse chromosomes.

Usage

```
avail_chromosomes()
```

Value

Data frame

Examples

```
avail_chromosomes()
```

avail_consequences *Available consequences*

Description

Available consequence and impact types.

Usage

```
avail_consequences()
```

Value

Data frame.

Examples

```
avail_consequences()$consequence  
unique(avail_consequences()$impact)
```

| | |
|---------------|--------------------------|
| avail_strains | <i>Available strains</i> |
|---------------|--------------------------|

Description

There are 37 strains available.

Usage

```
avail_strains()
```

Value

Data frame.

Examples

```
avail_strains()
```

| | |
|------------|--|
| df2GRanges | <i>Data frame to GenomicRanges::GRanges object</i> |
|------------|--|

Description

Wrapper for `GenomicRanges::makeGRangesFromDataFrame()`.

Usage

```
df2GRanges(  
  geno,  
  chr_name = "chr",  
  start_name = "pos",  
  end_name = "pos",  
  strand_name = NULL,  
  ref_version = ref_genome(),  
  seq_lengths = NULL,  
  is_circular = FALSE  
)
```

Arguments

| | |
|--------------------------|--|
| <code>geno</code> | Data frame. |
| <code>chr_name</code> | Name of chromosome column. Default is 'chr'. |
| <code>start_name</code> | Name of start position column. Default is 'pos'. |
| <code>end_name</code> | Name of end position column. Default is 'pos'. |
| <code>strand_name</code> | Name of end position column. Default is NULL. |
| <code>ref_version</code> | Reference genome version. Default is 'ref_genome()'. |
| <code>seq_lengths</code> | List of sequence lengths with sequence name as key. Default is NULL. |
| <code>is_circular</code> | Whether genome is circular. Default is FALSE. |

Value

GenomicRanges::GRanges object.

Examples

```

geno = finemap("chr1",
  start = 5000000, end = 6000000,
  strain1 = c("C57BL_6J"), strain2 = c("AKR_J", "A_J", "BALB_cJ")
)

geno$strand = "+"
seq_lengths = stats::setNames(
  as.list(avail_chromosomes()$length),
  avail_chromosomes()$chr
)
geno.granges = df2GRanges(geno,
  strand_name = "strand",
  seq_lengths = seq_lengths
)

```

fetch

Fetch

Description

Fetch homozygous genotypes for a specified chromosomal region in 37 inbred mouse strains.

Usage

```

fetch(
  chr,
  start = NULL,
  end = NULL,
  consequence = NULL,
  impact = NULL,
  return_obj = "dataframe"
)

```

Arguments

| | |
|-------------|---|
| chr | Vector of chromosome names. |
| start | Optional vector of chromosomal start positions of target regions (GRCm38). |
| end | Optional vector of chromosomal end positions of target regions (GRCm38). |
| consequence | Optional vector of consequence types. |
| impact | Optional vector of impact types. |
| return_obj | The user can choose to get the result to be returned as data frame ("dataframe") or as a GenomicRanges::GRanges ("granges") object. Default value is "dataframe". |

Value

Data frame or GenomicRanges::GRanges object containing result data.

Examples

```
geno = fetch("chr7", start = 5000000, end = 6000000)
comment(geno)
```

finemap

Finemapping of genetic regions

Description

Finemapping of genetic regions in 37 inbred mice by taking advantage of their very high homozygosity rate (>95 chromosomal regions (GRCm38), this method extracts homozygous SNVs for which the allele differs between two sets of strains (e.g. case vs controls) and outputs respective causal SNV/gene candidates.

Usage

```
finemap(  
  chr,  
  start = NULL,  
  end = NULL,  
  strain1,  
  strain2,  
  consequence = NULL,  
  impact = NULL,  
  thr1 = 0,  
  thr2 = 0,  
  return_obj = "dataframe"  
)
```

Arguments

| | |
|-------------|---|
| chr | Vector of chromosome names. |
| start | Optional vector of chromosomal start positions of target regions (GRCm38). |
| end | Optional vector of chromosomal end positions of target regions (GRCm38). |
| strain1 | First strain set with strains from avail_strains(). |
| strain2 | Second strain set with strains from avail_strains(). |
| consequence | Optional vector of consequence types. |
| impact | Optional vector of impact types. |
| thr1 | Number discordant strains in strain1. Between 0 and length(strain1)-1. 0 by default. |
| thr2 | Number discordant strains in strain2. Between 0 and length(strain2)-1. 0 by default. |
| return_obj | The user can choose to get the result to be returned as data frame ("dataframe") or as a GenomicRanges::GRanges ("granges") object. Default value is "dataframe". |

Value

Data frame or GenomicRanges::GRanges object containing result data.

Examples

```

geno = finemap("chr1",
  start = 5000000, end = 6000000,
  strain1 = c("C57BL_6J"), strain2 = c(
    "129S1_SvImJ", "129S5SvEvBrd",
    "AKR_J"
  )
)
comment(geno)

```

getURL

Get backend service url

Description

Get backend service URL. Default: <http://mousefm.genehopper.de/rest/finemap/>

Usage

```
getURL()
```

Value

URL string.

Examples

```
getURL()
```

| | |
|---------|---------------------------------|
| get_top | <i>Best strain combinations</i> |
|---------|---------------------------------|

Description

Get best strain combinations

Usage

```
get_top(red, n_top)
```

Arguments

| | |
|-------|--|
| red | Reduction factors data frame. |
| n_top | Number of combinations to be returned. |

Value

Data frame

Examples

```
l = prio("chr1",
  start = 5000000, end = 6000000,
  strain1 = "C57BL_6J", strain2 = "AKR_J"
)

get_top(l$reduction, 3)
```

| | |
|------------|--|
| GRanges2df | <i>GenomicRanges::GRanges object to data frame</i> |
|------------|--|

Description

Wrapper for as.data.frame().

Usage

```
GRanges2df(granges)
```

Arguments

| | |
|---------|-------------------------------|
| granges | GenomicRanges::GRanges object |
|---------|-------------------------------|

Value

Data frame.

Examples

```

geno.granges = finemap("chr1",
  start = 5000000, end = 6000000,
  strain1 = c("C57BL_6J"), strain2 = c("AKR_J", "A_J", "BALB_cJ"),
  return_obj = "granges"
)

geno = GRanges2df(geno.granges)

```

prio

Prioritization of inbred mouse strains for refining genetic regions

Description

This method allows to select strain combinations which best refine a specified genetic region (GRCm38). E.g. if a crossing experiment with two inbred mouse strains 'strain1' and 'strain2' resulted in a QTL, the outputted strain combinations can be used to refine the respective region in further crossing experiments.

Usage

```

prio(
  chr,
  start = NULL,
  end = NULL,
  strain1 = NULL,
  strain2 = NULL,
  consequence = NULL,
  impact = NULL,
  min_strain_benef = 0.1,
  max_set_size = 3,
  return_obj = "dataframe"
)

```

Arguments

| | |
|-------------|--|
| chr | Vector of chromosome names. |
| start | Optional vector of chromosomal start positions of target regions (GRCm38). |
| end | Optional vector of chromosomal end positions of target regions (GRCm38). |
| strain1 | First strain set with strains from avail_strains(). |
| strain2 | Second strain set with strains from avail_strains(). |
| consequence | Optional vector of consequence types. |

impact Optional vector of impact types.
 min_strain_benef Minimum reduction factor (min) of a single strain.
 max_set_size Maximum set of strains.
 return_obj The user can choose to get the result to be returned as data frame ("dataframe") or as a GenomicRanges::GRanges ("granges") object. Default value is "data frame".

Value

Data frame

Examples

```

res = prio("chr1",
  start = 5000000, end = 6000000, strain1 = "C57BL_6J",
  strain2 = "AKR_J"
)

comment(res$genotypes)

```

| | |
|------------|---------------------------------|
| ref_genome | <i>Reference genome version</i> |
|------------|---------------------------------|

Description

Returns version of reference genome used in package MouseFM.

Usage

```
ref_genome()
```

Value

Vector.

Examples

```
ref_genome()
```

| | |
|--------|--------------------------------|
| setURL | <i>Set backend service url</i> |
|--------|--------------------------------|

Description

Set backend service URL. Default: `http://mousefm.genehopper.de/rest/finemap/`

Usage

```
setURL(url)
```

Arguments

| | |
|-----|-------------------------|
| url | URL of backend service. |
|-----|-------------------------|

Value

No return value.

Examples

```
setURL("http://backendserver.com")
```

| | |
|-----------------------|------------------|
| vis_reduction_factors | <i>Visualize</i> |
|-----------------------|------------------|

Description

Visualize reduction factors

Usage

```
vis_reduction_factors(geno, red, n_top)
```

Arguments

| | |
|-------|---|
| geno | Genotype data frame or GenomicRanges::GRanges object. |
| red | Reduction factor data frame. |
| n_top | Number if combinations to be returned. |

Value

Data frame

Examples

```
l = prio(c("chr1", "chr2"),
        start = c(5000000, 5000000),
        end = c(6000000, 6000000), strain1 = c("C3H_HeH"), strain2 = "AKR_J"
        )

plots = vis_reduction_factors(l$genotypes, l$reduction, 2)

plots[[1]]
plots[[2]]
```

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