Package 'topr'

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Description A collection of functions for visualizing, exploring and annotating genetic association results. Association results from multiple traits can be viewed simultaneously along with gene annotation, over the entire genome (Manhattan plot) or in the more detailed regional view.
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annotate_with_nearest_gene

Get the nearest gene for one or more snps

Description

annotate_with_nearest_gene() Annotate the variant/snp with their nearest gene Required parameters is a dataframe of SNPs (with the columns CHROM and POS)

CD_FINNGEN 3

Usage

```
annotate_with_nearest_gene(
  variants,
  protein_coding_only = FALSE,
  build = 38,
  .chr_map = NULL
)
```

Arguments

variants a dataframe of variant positions (CHROM and POS)

protein_coding_only

Logical, if set to TRUE only annotate with protein coding genes (the default value is FALSE)

build A number representing the genome build. Set to 37 to change to build (GRCh37).

The defent is build 29 (CDCh29)

The default is build 38 (GRCh38).

. chr_map An internally used list which maps chromosome names to numbers.

Value

the input dataframe with Gene_Symbol as an additional column

Examples

```
## Not run:
variants <-get_lead_snps(CD_UKBB)
annotate_with_nearest_gene(variants)
## End(Not run)</pre>
```

CD_FINNGEN

Finngen r7 Crohn's disease (K11_CROHNS)

Description

Dataset retrieved from the Finngen database (version 7) including 3147 crohn's cases (K50) and 296,100 controls. The dataset has been filtered on variants with P <1e-03. FinnGen data are publicly available and were downloaded from https://finngen.fi.

Usage

CD_FINNGEN

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Format

A data frame with 32,303 rows and 8 variables:

CHROM Chromosome, written as for example chr1 or 1

POS genetic position of the variant

REF the reference allele

ALT the alternative allele

P P-value from Plink run, additive model, regression model GLM_FIRTH

BETA Variant effect

ID Variant identifier, e.g. rsid

AF Allele frequency

Source

Crohn's K50 (K11_CROHNS), only including variants with P<1e-03

CD_UKBB

UKBB Crohns disease (ICD 10 code K50)

Description

Dataset retrieved from the UK biobank consisting of 2,799 crohn's cases (K50) and 484,515 controls. The dataset has been filtered on variants with P <1e-03.

Usage

CD_UKBB

Format

A data frame with 21,717 rows and 8 variables:

CHROM Chromosome, written as for example chr1 or 1

POS genetic position of the variant

REF the reference allele

ALT the alternative allele

ID Variant identifier, e.g. rsid

P P-value from Plink run, additive model, regression model GLM_FIRTH

OR Odds Ratio

AF Allele frequency

Source

Crohn's UKBB ICD10 code K50, only including variants with P<1e-03

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create_snpset

Create a dataframe that can be used as input for making effect plots

Description

```
create_snpset()
```

This method is deprecated and will be removed in future versions. use get_snpset instead.

Usage

```
create_snpset(
   df1,
   df2,
   thresh = 1e-08,
   protein_coding_only = TRUE,
   region_size = 1e+06,
   verbose = F
)
```

Arguments

df1 The dataframe to extract the top snps from (with p-value below thresh)

df2 The dataframe in which to search for overlapping SNPs from dataframe1

thresh Numeric, the p-value threshold used for extracting the top snps from dataset 1

protein_coding_only

Logical, set this variable to TRUE to only use protein_coding genes for the annotation

region_size Integer, the size of the interval which to extract the top snps from

verbose Logical, (default: FALSE). Assign to TRUE to get information on which alleles

Value

Dataframe containing the top hit

Examples

```
## Not run:
create_snpset(CD_UKBB,CD_FINNGEN, thresh=1e-09)
## End(Not run)
```

are matched and which are not.

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create_snpset_code

Show the code/functions used to create a snpset

Description

This method is deprecated and will be removed in future versions. use get_snpset_code instead. create_snpset_code()

Usage

```
create_snpset_code()
```

Value

Dataframe containing the top hit

Examples

```
## Not run:
create_snpset_code()
## End(Not run)
```

 ${\tt effectplot}$

Create a plot comparing variant effects in two datasets

Description

```
effectplot()
```

Usage

```
effectplot(
   df,
   pheno_x = "x_pheno",
   pheno_y = "y_pheno",
   annotate_with = "Gene_Symbol",
   thresh = 5e-08,
   ci_thresh = 1,
   gene_label_thresh = 5e-08,
   color = get_topr_colors()[1],
   scale = 1,
   build = 38,
   label_fontface = "italic",
   label_family = "",
```

effectplot 7

```
nudge_y = 0.001,
 nudge_x = 0.001,
  size = 2,
  segment.size = 0.2,
  segment.linetype = "solid",
  segment.color = "transparent",
  angle = 0,
  title = NULL,
  axis_text_size = 10,
  axis_title_size = 12,
  title_text_size = 13,
  subtitle_text_size = 11,
  gene_label_size = 3.2,
  snpset_thresh = 5e-08,
  snpset_region_size = 1e+06,
 max.overlaps = 10,
  annotate = 0,
  label_color = NULL
)
```

Arguments df

_	and P2) and effects (E1 and E2) from two datasets/phenotypes OR a list containing two datasets.
pheno_x	A string representing the name of the phenotype whose effect is plotted on the x axis
pheno_y	A string representing the name of the phenotype whose effect is plotted on the y axis
annotate_with	A string, The name of the column that contains the label for the datapoints (default value is Gene_Symbol)
thresh	A number. Threshold cutoff, datapoints with P2 below this threshold are shown as filled circles whereas datapoints with P2 above this threshold are shown as

as filled circles whereas datapoints with P2 above this threshold are shown as open circles

The input dataframe (snpset) containing one row per variant and P values (P1

ci_thresh A number.Show the confidence intervals if the P-value is below this threshold gene_label_thresh

Deprecated: A number, label datapoints with P2 below this threshold

color A string, default value is the first of the topr colors

scale A number, to change the size of the title and axes labels and ticks at the same

time (default: 1)

build A number representing the genome build or a data frame. Set to 37 to change to

build (GRCh37). The default is build 38 (GRCh38).

label_fontface A string or a vector of strings. Label font "plain", "bold", "italic", "bold.italic"

(ggrepel argument)

label_family A string or a vector of strings. Label font name (default ggrepel argument is "")

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A number to horizontally adjust the starting position of each gene label (this is nudge_y a ggrepel parameter) A number to vertically adjust the starting position of each gene label (this is a nudge_x ggrepel parameter) A number or a vector of numbers, setting the size of the plot points (default: size size=1.2line segment color (ggrepel argument) segment.size segment.linetype line segment solid, dashed, etc.(ggrepel argument) segment.color line segment thickness (ggrepel argument) A number, the angle of the text label angle title A string to set the plot title axis_text_size A number, size of the x and y axes tick labels (default: 12) axis_title_size A number, size of the x and y title labels (default: 12) title_text_size A number, size of the plot title (default: 13) subtitle_text_size A number setting the text size of the subtitle (default: 11) gene_label_size A number setting the size of the gene labels shown at the bottom of the plot snpset_thresh A number representing the threshold used to create the snpset used for plotting (Only applicable if the input dataframe is a list containing two datasets) snpset_region_size A number representing the region size to use when creating the snpset used for plotting (Only applicable if the input dataframe is a list containing two datasets) Exclude text labels that overlap too many things. Defaults to 10 (ggrepel argumax.overlaps annotate A number, label datapoints with p-value below below this number (in the second df) by their nearest gene A string or a vector of strings. To change the color of the gene or variant labels label_color

Value

ggplot object

Examples

```
## Not run:
effectplot(list(CD_UKBB, CD_FINNGEN))
## End(Not run)
```

effect_plot

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Create a plot comparing effects within two datasets

Description

```
effect_plot()
```

This method is deprecated and will be removed in future versions. use effectplot instead.

Usage

```
effect_plot(
  dat,
  pheno_x = "pheno_x",
  pheno_y = "pheno_",
  annotate_with = "Gene_Symbol",
  thresh = 1e-08,
  ci_thresh = 1,
  gene_label_thresh = 1e-08,
  color = get_topr_colors()[1],
  scale = 1
)
```

time (default = 1)

Arguments

	dat	The input dataframe (snpset) containing one row per variant and P values (P1 and P2) and effects (E1 and E2) from two datasets/phenotypes
	pheno_x	A string representing the name of the phenotype whose effect is plotted on the x axis
	pheno_y	A string representing the name of the phenotype whose effect is plotted on the y axis
	annotate_with	A string, The name of the column that contains the label for the datapoints (default value is Gene_Symbol)
	thresh	A number. Threshold cutoff, datapoints with P2 below this threshold are shown as filled circles whereas datapoints with P2 above this threshold are shown as open circles
	ci_thresh	A number. Show the confidence intervals if the P-value is below this threshold
gene_label_thresh		esh
		A string, label datapoints with P2 below this threshold
	color	A string, default value is the first of the topr colors
	scale	A number, to change the size of the title and axes labels and ticks at the same

Examples

```
## Not run:
effect_plot(dat)
## End(Not run)
```

```
flip\_to\_positive\_allele\_for\_dat1\\ Flip\ to\ the\ positive\ allele\ for\ dataset\ 1
```

Description

```
flip_to_positive_allele_for_dat1()
```

Usage

```
flip_to_positive_allele_for_dat1(df)
```

Arguments

df

A dataframe that is in the snpset format (like returned by the get_snpset() function)

Value

The input dataframe after flipping to the positive effect allele in dataframe 1

Examples

```
## Not run:
CD_UKBB_index_snps <- get_lead_snps(CD_UKBB)
snpset <- get_snpset(CD_UKBB_index_snps, CD_FINNGEN)
flip_to_positive_allele_for_dat1(snpset$matched)
## End(Not run)</pre>
```

get_best_snp_per_MB Get the index/lead variants

Description

 $get_best_snp_per_MB()$ Get the top variants within 1 MB windows of the genome with association p-values below the given threshold

This method is deprecated and will be removed in future versions. use get_lead_snps instead.

Usage

```
get_best_snp_per_MB(
   df,
   thresh = 5e-09,
   region_size = 1e+06,
   protein_coding_only = FALSE,
   chr = NULL,
   .checked = FALSE,
   verbose = FALSE
)
```

Arguments

df	Dataframe		
thresh	A number. P-value threshold, only extract variants with p-values below this threshold (5e-09 by default) $\frac{1}{2}$		
region_size	An integer (default = 20000000) (or a string represented as $200kb$ or $2MB$) indicating the window size for variant labeling. Increase this number for sparser annotation and decrease for denser annotation.		
<pre>protein_coding_only</pre>			
	Logical, set this variable to TRUE to only use protein_coding genes for annotation		
chr	String, get the top variants from one chromosome only, e.g. chr="chr1"		
. checked	Logical, if the input data has already been checked, this can be set to TRUE so it wont be checked again (FALSE by default)		
verbose	Logical, set to TRUE to get printed information on number of SNPs extracted		

Value

Dataframe of lead variants. Returns the best variant per MB (by default, change the region size with the region argument) with p-values below the input threshold (thresh=5e-09 by default)

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Examples

```
## Not run:
    get_best_snp_per_MB(CD_UKBB)
## End(Not run)
```

get_gene

Get the genetic position of a gene by gene name

Description

get_gene() Get the gene coordinates for a gene Required parameter is gene name

This method is deprecated and will be removed in future versions. use get_gene_coords instead.

Usage

```
get_gene(gene_name, chr = NULL, build = 38)
```

Arguments

gene_name A string representing a gene name (e.g. "FTO")

chr A string, search for the genes on this chromosome only, (e.g chr="chr1")

build A string, genome build, choose between builds 37 (GRCh37) and 38 (GRCh38)

(default is 38)

Value

Dataframe with the gene name and its genetic coordinates

Examples

```
## Not run:
get_gene("FTO")
## End(Not run)
```

```
get_genes_by_Gene_Symbol
```

Get the genetic position of a gene by its gene name

Description

get_genes_by_Gene_Symbol() Get genes by their gene symbol/name Required parameters is on gene name or a vector of gene names

Usage

```
get_genes_by_Gene_Symbol(genes, chr = NULL, build = 38)
```

Arguments

genes A string or vector of strings representing gene names, (e.g. "FTO") or (c("FTO","NOD2"))

chr A string, search for the genes on this chromosome only, (e.g chr="chr1")

build A string, genome build, choose between builds 37 (GRCh37) and 38 (GRCh38)

(default is 38)

Value

Dataframe of genes

Examples

```
## Not run:
   get_genes_by_Gene_Symbol(c("FTO","THADA"))
## End(Not run)
```

get_genes_in_region

Get SNPs/variants within region

Description

```
get_genes_in_region()
```

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Usage

```
get_genes_in_region(
  chr = chr,
  xmin = xmin,
  xmax = xmax,
  protein_coding_only = F,
  show_exons = F,
  show_genes = T,
  build = 38,
  region = NULL
)
```

Arguments

chr A string, chromosome (e.g. chr16)

xmin An integer representing genetic position

xmax An integer representing genetic position

protein_coding_only

A logical scalar, if TRUE, only protein coding genes are used for annotation

show_exons Deprecated: A logical scalar, show exons instead of genes (default show_exons=FALSE)

show_genes A logical scalar, show genes instead of exons (default show_genes=FALSE) build A number representing the genome build or a data frame. Set to 37 to change to

build (GRCh37). The default is build 38 (GRCh38).

region A string representing the genetic region (e.g chr16:50693587-50734041)

Value

the genes the requested region

Examples

```
## Not run:
get_genes_in_region(region="chr16:50593587-50834041")
## End(Not run)
```

get_gene_coords

Get the genetic position of a gene by gene name

Description

get_gene_coords() Get the gene coordinates for a gene Required parameter is gene name

Usage

```
get_gene_coords(gene_name, chr = NULL, build = 38)
```

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Arguments

gene_name A string representing a gene name (e.g. "FTO")

chr A string, search for the genes on this chromosome only, (e.g chr="chr1")

build A string, genome build, choose between builds 37 (GRCh37) and 38 (GRCh38)

(default is 38)

Value

Dataframe with the gene name and its genetic coordinates

Examples

```
## Not run:
get_gene_coords("FTO")
## End(Not run)
```

get_lead_snps

Get the index/lead variants

Description

get_lead_snps() Get the top variants within 1 MB windows of the genome with association p-values below the given threshold

Usage

```
get_lead_snps(
   df,
   thresh = 5e-08,
   region_size = 1e+06,
   protein_coding_only = FALSE,
   chr = NULL,
   .checked = FALSE,
   verbose = NULL,
   keep_chr = TRUE
)
```

Arguments

df Dataframe

thresh A number. P-value threshold, only extract variants with p-values below this

threshold (5e-08 by default)

region_size An integer (default = 20000000) (or a string represented as 200kb or 2MB)

indicating the window size for variant labeling. Increase this number for sparser

annotation and decrease for denser annotation.

protein_coding_only

Logical, set this variable to TRUE to only use protein_coding genes for annota-

tion

chr String, get the top variants from one chromosome only, e.g. chr="chr1"

. checked Logical, if the input data has already been checked, this can be set to TRUE so

it wont be checked again (FALSE by default)

verbose Logical, set to TRUE to get printed information on number of SNPs extracted keep_chr Logical, set to FALSE to remove the "chr" prefix before each chromosome if

present (TRUE by default)

Value

Dataframe of lead variants. Returns the best variant per MB (by default, change the region size with the region argument) with p-values below the input threshold (thresh=5e-08 by default)

Examples

```
## Not run:
get_lead_snps(CD_UKBB)
## End(Not run)
```

```
get_overlapping_snps_by_pos
```

Get variants that overlap between two datasets

Description

```
get_overlapping_snps_by_pos()
```

This method is deprecated and will be removed in future versions. use match_by_pos instead.

Usage

```
get_overlapping_snps_by_pos(df1, df2, verbose = F)
```

Arguments

df1 A dataframe of variants, has to contain CHROM and POS df2 A dataframe of variants, has to contain CHROM and POS

verbose A logical scalar (default: FALSE). Assign to TRUE to get information on which

alleles are matched and which are not.

Value

The input dataframe containing only those variants with matched alleles in the snpset

Examples

Description

get_lead_snps() Get the top variants within 1 MB windows of the genome with association p-values below the given threshold

Usage

```
get_sign_and_sugg_loci(
   df,
   genome_wide_thresh = 5e-08,
   suggestive_thresh = 1e-06,
   flank_size = 1e+06,
   region_size = 1e+06
)
```

Arguments

Value

List of genome wide and suggestive loci.

Examples

```
## Not run:
get_sign_and_sugg_loci(CD_UKBB)
## End(Not run)
```

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get_snpset

Create a dataframe that can be used as input for making effect plots

Description

```
get_snpset()

Usage

get_snpset(
    df1,
    df2,
    thresh = 5e-08,
    protein_coding_only = TRUE,
    region_size = 1e+06,
    verbose = NULL,
    show_full_output = FALSE,
    build = 38,
    format = "wide"
)
```

Arguments

df1 The dataframe to extract the top snps from (with p-value below thresh)
df2 The dataframe in which to search for overlapping SNPs from dataframe1

thresh A number. P-value threshold, only extract variants with p-values below this

threshold (5e-08 by default)

protein_coding_only

Logical, set this variable to TRUE to only use protein coding genes for annota-

tion

region_size An integer (default = 20000000) (or a string represented as 200kb or 2MB)

indicating the window size for variant labeling. Increase this number for sparser

annotation and decrease for denser annotation.

verbose Logical, (default: FALSE). Assign to TRUE to get information on which alleles

are matched and which are not.

show_full_output

A logical scalar (default:FALSE). Assign to TRUE to show the full output from

this function

build A string, genome build, choose between builds 37 (GRCh37) and 38 (GRCh38)

(default is 38)

format A string, representing either wide or long format (default : "wide"). By default

a snpset created from two dataframes is returned in a wide format.

Value

Dataframe of overlapping snps (snpset)

get_snpset_code 19

Examples

```
## Not run:
CD_UKBB_index_snps <-get_lead_snps(CD_UKBB)
get_snpset(CD_UKBB_index_snps, CD_FINNGEN)
## End(Not run)</pre>
```

get_snpset_code

Show the code/functions used to get a snpset

Description

```
get_snpset_code()
```

Usage

```
get_snpset_code()
```

Value

Dataframe containing the top hit

Examples

```
## Not run:
get_snpset_code()
## End(Not run)
```

```
get_snps_within_region
```

Get SNPs/variants within region

Description

```
get_snps_within_region()
```

Usage

```
get_snps_within_region(
   df,
   region,
   chr = NULL,
   xmin = NULL,
   xmax = NULL,
   keep_chr = NULL
)
```

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Arguments

df data frame of association results with the columns CHR and POS

region A string representing the genetic region (e.g chr16:50693587-50734041)

chr A string, chromosome (e.g. chr16)

xmin An integer, include variants with POS larger than xminxmax An integer, include variants with POS smaller than xmax

keep_chr Deprecated: Logical, set to FALSE to remove the "chr" prefix before each chro-

mosome if present (TRUE by default)

Value

the variants within the requested region

Examples

```
## Not run:
get_snps_within_region(CD_UKBB, "chr16:50593587-50834041")
## End(Not run)
```

get_topr_colors

Get the top hit from the dataframe

Description

get_topr_colors() Get the top hit from the dataframe All other input parameters are optional

Usage

```
get_topr_colors()
```

Value

Vector of colors used for plotting

Examples

```
## Not run:
get_topr_colors()
## End(Not run)
```

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Get the top hit from the dataframe

Description

get_top_snp() Get the top hit from the dataframe All other input parameters are optional

Usage

```
get_top_snp(df, chr = NULL)
```

Arguments

df Dataframe containing association results

chr String, get the top hit in the data frame for this chromosome. If chromosome is

not provided, the top hit from the entire dataset is returned.

Value

Dataframe containing the top hit

Examples

```
## Not run:
get_top_snp(CD_UKBB, chr="chr1")
## End(Not run)
```

locuszoom

Create a locuszoom-like plot

Description

locuszoom() displays the association results for a smaller region within one chromosome. Required parameter is at least one dataset (dataframe) containing the association data (with columns CHROM, POS, P in upper or lowercase)

Usage

```
locuszoom(
  df,
  annotate = NULL,
  ntop = 3,
  xmin = 0,
  size = 2,
  shape = 19,
  alpha = 1,
  label_size = 4,
  annotate_with = "ID",
  color = NULL,
  axis_text_size = 11,
  axis_title_size = 12,
  title_text_size = 13,
  show_genes = NULL,
  show_overview = FALSE,
  show_exons = FALSE,
  max\_genes = 200,
  sign_thresh = 5e-08,
  sign_thresh_color = "red",
  sign_thresh_label_size = 3.5,
  xmax = NULL,
  ymin = NULL,
  ymax = NULL,
  protein_coding_only = FALSE,
  region_size = 1e+06,
  gene_padding = 1e+05,
  angle = 0,
  legend_title_size = 12,
  legend_text_size = 12,
  nudge_x = 0.01,
  nudge_y = 0.01,
  rsids = NULL,
  variant = NULL,
  rsids_color = "gray40",
  legend_name = "Data:",
  legend_position = "right",
  chr = NULL,
  vline = NULL,
  show_gene_names = NULL,
  legend_labels = NULL,
  gene = NULL,
  title = NULL,
  label_color = "gray40",
  region = NULL,
  scale = 1,
  rsids_with_vline = NULL,
```

```
annotate_with_vline = NULL,
  sign_thresh_size = 0.5,
  unit_main = 7,
  unit_gene = 2,
  gene_color = NULL,
  segment.size = 0.2,
  segment.color = "black",
  segment.linetype = "solid",
  show_gene_legend = TRUE,
 max.overlaps = 10,
  extract_plots = FALSE,
  label_fontface = "plain",
  label_family = "",
  gene_label_fontface = "plain",
  gene_label_family = "",
  build = 38,
  verbose = NULL,
  show_legend = TRUE,
  label_alpha = 1,
  gene_label_size = NULL,
  vline_color = "grey",
  vline_linetype = "dashed",
  vline_alpha = 1,
 vline\_size = 0.5,
 log_trans_p = TRUE
)
```

Arguments

df	Dataframe or a list of dataframes (required columns are CHROM, POS, P), in upper- or lowercase) of association results.
annotate	A number (p-value). Display annotation for variants with p-values below this threshold
ntop	An integer, number of datasets (GWAS results) to show on the top plot
xmin, xmax	Integer, setting the chromosomal range to display on the x-axis
size	A number or a vector of numbers, setting the size of the plot points (default: size=1.2)
shape	A number of a vector of numbers setting the shape of the plotted points
alpha	A number or a vector of numbers setting the transparency of the plotted points
label_size	An number to set the size of the plot labels (default: label_size=3)
annotate_with	A string. Annotate the variants with either Gene_Symbol or ID (default: "Gene_Symbol")
color	A string or a vector of strings, for setting the color of the datapoints on the plot
<pre>axis_text_size</pre>	A number, size of the x and y axes tick labels (default: 12)
axis_title_size	
	A number, size of the x and y title labels (default: 12)

title_text_size

A number, size of the plot title (default: 13)

show_genes A logical scalar, show genes instead of exons (default show_genes=FALSE)

show_overview A logical scalar, shows/hides the overview plot (default= TRUE)

show_exons Deprecated: A logical scalar, show exons instead of genes (default show_exons=FALSE)

max_genes An integer, only label the genes if they are fewer than max_genes (default values

is 200).

sign_thresh A number or vector of numbers, setting the horizontal significance threshold

(default: sign_thresh=5e-8). Set to NULL to hide the significance threshold.

sign_thresh_color

A string or vector of strings to set the color/s of the significance threshold/s

sign_thresh_label_size

A number setting the text size of the label for the significance thresholds (default

text size is 3.5)

ymin, ymax Integer, min and max of the y-axis, (default values: ymin=0, ymax=max(-log10(df\$P)))

protein_coding_only

A logical scalar, if TRUE, only protein coding genes are used for annotation

region_size An integer (default = 20000000) (or a string represented as 200kb or 2MB)

indicating the window size for variant labeling. Increase this number for sparser

annotation and decrease for denser annotation.

gene_padding An integer representing size of the region around the gene, if the gene argument

was used (default = 100000)

angle A number, the angle of the text label

legend_title_size

A number, size of the legend title

legend_text_size

A number, size of the legend text

nudge_x A number to vertically adjust the starting position of each gene label (this is a

ggrepel parameter)

nudge_y A number to horizontally adjust the starting position of each gene label (this is

a ggrepel parameter)

rsids A string (rsid) or vector of strings to highlight on the plot, e.g. rsids=c("rs1234,

rs45898")

variant A string representing the variant to zoom in on. Can be either an rsid, or a

dataframe (with the columns CHROM,POS,P)

rsids_color A string, the color of the variants in variants_id (default color is red)

legend_name A string, use to change the name of the legend (default: None)

legend_position

A string, top, bottom, left or right

chr A string or integer, the chromosome to plot (i.e. chr15), only required if the

input dataframe contains results from more than one chromosome

vline A number or vector of numbers to add a vertical line to the plot at a specific

chromosomal position, e.g vline=204000066. Multiple values can be provided

in a vector, e.g vline=c(204000066,100500188)

show_gene_names

A logical scalar, if set to TRUE, gene names are shown even though they exceed

the max_genes count

legend_labels A string or vector of strings representing legend labels for the input datasets

gene A string representing the gene to zoom in on (e.g. gene=FTO)

title A string to set the plot title

label_color A string or a vector of strings. To change the color of the gene or variant labels

region A string representing a genetic region, e.g. chr1:67038906-67359979

scale A number, to change the size of the title and axes labels and ticks at the same

time (default : 1)

rsids_with_vline

A string (rsid) or vector of strings to highlight on the plot with their rsids and

vertical lines further highlighting their positions

annotate_with_vline

A number (p-value). Display annotation and vertical lines for variants with p-

values below this threshold

sign_thresh_size

A number, sets the size of the horizontal significance threshold line (default : 1)

unit_main the height unit of the main plot (default = 7)
unit_gene the height unit of the gene plot (default= 2)

gene_color A string representing a color, can be used to change the color of the genes/exons

on the geneplot

segment.size line segment color (ggrepel argument)

segment.color line segment thickness (ggrepel argument)

segment.linetype

line segment solid, dashed, etc.(ggrepel argument)

show_gene_legend

A logical scalar, set to FALSE to hide the gene legend (default value is TRUE)

max.overlaps Exclude text labels that overlap too many things. Defaults to 10 (ggrepel argu-

ment)

a list

label_fontface A string or a vector of strings. Label font "plain", "bold", "italic", "bold.italic"

(ggrepel argument)

label_family A string or a vector of strings. Label font name (default ggrepel argument is "")

gene_label_fontface

Gene label font "plain", "bold", "italic", "bold.italic" (ggrepel argument)

gene_label_family

Gene label font name (default ggrepel argument is "")

build A number representing the genome build or a data frame. Set to 37 to change to build (GRCh37). The default is build 38 (GRCh38).

verbose Logical, set to FALSE to get suppress printed information

show_legend A logical scalar, set to FALSE to hide the legend (default : TRUE)

fault: label_alpha=1)

gene_label_size

A number setting the size of the gene labels shown at the bottom of the plot

vline_color A string. The color of added vertical line/s (default: grey)
vline_linetype A string. The linetype of added vertical line/s (default: dashed)
vline_alpha A number. The alpha of added vertical line/s (default: 1)
vline_size A number. The size of added vertical line/s (default: 0.5)

log_trans_p A logical scalar (default: TRUE). By default the p-values in the input datasets

are log transformed using -log10. Set this argument to FALSE if the p-values in

the datasets have already been log transformed.

Value

plots using egg (https://cran.r-project.org/web/packages/egg/vignettes/Ecosystem.html)

Examples

```
## Not run:
locuszoom(R2_CD_UKBB)
## End(Not run)
```

manhattan

Create a Manhattan plot

Description

manhattan() displays association results for the entire genome on a Manhattan plot. Required parameter is at least one dataset (dataframe) containing the association data (with columns CHROM, POS, P in upper or lowercase)

All other input parameters are optional

Usage

```
manhattan(
   df,
   ntop = 4,
   title = "",
   annotate = NULL,
   color = NULL,
```

```
sign_{thresh} = 5e-08,
sign_thresh_color = "red",
sign_thresh_label_size = 3.5,
label_size = 3.5,
size = 0.8,
shape = 19,
alpha = 1,
highlight_genes_color = "darkred",
highlight_genes_ypos = 1.5,
axis_text_size = 12,
axis_title_size = 14,
title_text_size = 15,
legend_title_size = 13,
legend_text_size = 12,
protein_coding_only = TRUE,
angle = 0,
legend_labels = NULL,
chr = NULL,
annotate_with = "Gene_Symbol",
region_size = 2e+07,
legend_name = NULL,
legend_position = "bottom",
nudge_x = 0.1,
nudge_y = 0.7,
xmin = NULL,
xmax = NULL,
ymin = NULL,
ymax = NULL,
highlight_genes = NULL,
label_color = NULL,
legend_nrow = NULL,
gene_label_size = NULL,
gene_label_angle = 0,
scale = 1,
show_legend = TRUE,
sign_thresh_linetype = "dashed",
sign_thresh_size = 0.5,
rsids = NULL,
rsids_color = NULL,
rsids_with_vline = NULL,
annotate_with_vline = NULL,
shades_color = NULL,
shades_alpha = 0.5,
segment.size = 0.2,
segment.color = "black",
segment.linetype = "dashed",
max.overlaps = 10,
label_fontface = "plain",
```

```
label_family = "",
  gene_label_fontface = "plain",
  gene_label_family = "",
  build = 38,
  verbose = NULL,
  label_alpha = 1,
  shades_line_alpha = 1,
  vline = NULL,
  vline_color = "grey",
  vline_linetype = "dashed",
  vline_alpha = 1,
  vline_size = 0.5,
  region = NULL,
  theme_grey = FALSE,
  xaxis_label = "Chromosome",
  use_shades = FALSE,
  even_no_chr_lightness = 0.8,
  get_chr_lengths_from_data = TRUE,
  log_trans_p = TRUE,
  chr_ticknames = NULL,
  show_all_chrticks = FALSE,
  hide_chrticks_from_pos = 17,
  hide_chrticks_to_pos = NULL,
  hide_every_nth_chrtick = 2,
  downsample\_cutoff = 0.05,
  downsample_prop = 0.1
)
```

Arguments

df	Dataframe	e or a list o	f dataframes	(required columns are	CHROM, POS, P), in upper-
	1	\ C		1.	

or lowercase) of association results.

ntop An integer, number of datasets (GWAS results) to show on the top plot

title A string to set the plot title

annotate A number (p-value). Display annotation for variants with p-values below this

threshold

color A string or a vector of strings, for setting the color of the datapoints on the plot

sign_thresh A number or vector of numbers, setting the horizontal significance threshold

(default: sign_thresh=5e-8). Set to NULL to hide the significance threshold.

sign_thresh_color

A string or vector of strings to set the color/s of the significance threshold/s

sign_thresh_label_size

A number setting the text size of the label for the significance thresholds (default

text size is 3.5)

label_size An number to set the size of the plot labels (default: label_size=3)

size A number or a vector of numbers, setting the size of the plot points (default:

size=1.2)

shape A number of a vector of numbers setting the shape of the plotted points

alpha A number or a vector of numbers setting the transparency of the plotted points

highlight_genes_color

A string, color for the highlighted genes (default: darkred)

highlight_genes_ypos

An integer, controlling where on the y-axis the highlighted genes are placed

(default value is 1)

axis_text_size A number, size of the x and y axes tick labels (default: 12)

axis_title_size

A number, size of the x and y title labels (default: 12)

title_text_size

A number, size of the plot title (default: 13)

legend_title_size

A number, size of the legend title

legend_text_size

A number, size of the legend text

protein_coding_only

A logical scalar, if TRUE, only protein coding genes are used for annotation

angle A number, the angle of the text label

legend_labels A string or vector of strings representing legend labels for the input datasets

chr A string or integer, the chromosome to plot (i.e. chr15), only required if the

input dataframe contains results from more than one chromosome

annotate_with A string. Annotate the variants with either Gene_Symbol or ID (default: "Gene_Symbol")

region_size An integer (default = 20000000) (or a string represented as 200kb or 2MB)

indicating the window size for variant labeling. Increase this number for sparser

annotation and decrease for denser annotation.

legend_name A string, use to change the name of the legend (default: None)

legend_position

A string, top, bottom, left or right

nudge_x A number to vertically adjust the starting position of each gene label (this is a

ggrepel parameter)

nudge_y A number to horizontally adjust the starting position of each gene label (this is

a ggrepel parameter)

xmin, xmax Integer, setting the chromosomal range to display on the x-axis

ymin, ymax Integer, min and max of the y-axis, (default values: ymin=0, ymax=max(-log10(df\$P)))

highlight_genes

A string or vector of strings, gene or genes to highlight at the bottom of the plot

label_color A string or a vector of strings. To change the color of the gene or variant labels

legend_nrow An integer, sets the number of rows allowed for the legend labels

gene_label_size

A number setting the size of the gene labels shown at the bottom of the plot

gene_label_angle

A number setting the angle of the gene label shown at the bottom of the plot

(default: 0)

scale A number, to change the size of the title and axes labels and ticks at the same

time (default: 1)

show_legend A logical scalar, set to FALSE to hide the legend (default : TRUE)

sign_thresh_linetype

A string, the line-type of the horizontal significance threshold (default : dashed)

sign_thresh_size

A number, sets the size of the horizontal significance threshold line (default : 1)

rsids A string (rsid) or vector of strings to highlight on the plot, e.g. rsids=c("rs1234,

rs45898")

rsids_color A string, the color of the variants in variants_id (default color is red)

rsids_with_vline

A string (rsid) or vector of strings to highlight on the plot with their rsids and vertical lines further highlighting their positions

annotate_with_vline

A number (p-value). Display annotation and vertical lines for variants with p-

values below this threshold

the Manhattan plot

shades_alpha The transparency (alpha) of the rectangles (shades)

segment.size line segment color (ggrepel argument)

segment.color line segment thickness (ggrepel argument)

segment.linetype

line segment solid, dashed, etc.(ggrepel argument)

max.overlaps Exclude text labels that overlap too many things. Defaults to 10 (ggrepel argu-

ment)

label_fontface A string or a vector of strings. Label font "plain", "bold", "italic", "bold.italic"

(ggrepel argument)

label_family A string or a vector of strings. Label font name (default ggrepel argument is "")

gene_label_fontface

Gene label font "plain", "bold", "italic", "bold.italic" (ggrepel argument)

gene_label_family

Gene label font name (default ggrepel argument is "")

build A number representing the genome build or a data frame. Set to 37 to change to

build (GRCh37). The default is build 38 (GRCh38).

verbose A logical scalar (default: NULL). Set to FALSE to suppress printed messages

label_alpha An number or vector of numbers to set the transparency of the plot labels (de-

fault: label_alpha=1)

shades_line_alpha

The transparency (alpha) of the lines around the rectangles (shades)

vline A number or vector of numbers to add a vertical line to the plot at a specific

> chromosomal position, e.g vline="chr1:204000066". Multiple values can be provided in a vector, e.g vline=c("chr1:204000066", "chr5:100500188")

A string. The color of added vertical line/s (default: grey) vline_color

vline_linetype A string. The linetype of added vertical line/s (default : dashed)

A number. The alpha of added vertical line/s (default: 1) vline_alpha vline_size A number. The size of added vertical line/s (default : 0.5)

region A string representing a genetic region, e.g. chr1:67038906-67359979

A logical scalar (default: FALSE). Use gray rectangles (instead of white to distheme_grev

tinguish between chromosomes)

A string. The label for the x-axis (default: Chromosome) xaxis_label

use_shades A logical scalar (default: FALSE). Use shades/rectangles to distinguish between

chromosomes

even_no_chr_lightness

Lightness value for even numbered chromosomes. A number or vector of numbers between 0 and 1 (default: 0.8). If set to 0.5, the same color as shown for odd numbered chromosomes is displayed. A value below 0.5 will result in a darker color displayed for even numbered chromosomes, whereas a value above

0.5 results in a lighter color.

get_chr_lengths_from_data

A logical scalar (default: TRUE). If set to FALSE, use the inbuilt chromosome lengths (from hg38), instead of chromosome lengths based on the max position

for each chromosome in the input dataset/s.

A logical scalar (default: TRUE). By default the p-values in the input datasets log_trans_p

are log transformed using -log10. Set this argument to FALSE if the p-values in

the datasets have already been log transformed.

chr_ticknames A vector containing the chromosome names displayed on the x-axis. If NULL,

the following format is used: chr_ticknames <- c(1:16, ",18, ",20, ",22, 'X')

show_all_chrticks

A logical scalar (default : FALSE). Set to TRUE to show all the chromosome names on the ticks on the x-axis

hide_chrticks_from_pos

A number (default: 17). Hide every nth chromosome name on the x-axis FROM

this position (chromosome number)

hide_chrticks_to_pos

A number (default: NULL). Hide every nth chromosome name on the x-axis TO this position (chromosome number). When NULL this variable will be set to the

number of numeric chromosomes in the input dataset.

hide_every_nth_chrtick

A number (default: 2). Hide every nth chromosome tick on the x-axis (from the hide_chr_ticks_from_pos to the hide_chr_ticks_to_pos).

downsample_cutoff

A number (default: 0.05) used to downsample the input dataset prior to plotting. Sets the fraction of high p-value (default: P>0.05) markers to display on the plot.

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```
downsample_prop
```

A number (default: 0.1) used to downsample the input dataset prior to plotting. Only a proportion of the variants (10% by default) with P-values higher than the downsample_cutoff will be displayed on the plot.

Value

ggplot object

Examples

```
## Not run:
manhattan(CD_UKBB)
## End(Not run)
```

manhattanExtra

Create a Manhattan plot highlighting genome-wide significant and suggestive loci

Description

manhattanExtra() displays association results for the entire genome on a Manhattan plot, high-lighting genome-wide significant and suggestive loci. Required parameter is at least one dataset (dataframe) containing the association data (with columns CHROM, POS, P in upper or lowercase)

All other input parameters are optional

Usage

```
manhattanExtra(
  df,
  genome_wide_thresh = 5e-08,
  suggestive_thresh = 1e-06,
  flank_size = 1e+06,
  region_size = 1e+06,
  sign_thresh_color = NULL,
  sign_thresh_label_size = NULL,
  show_legend = TRUE,
  label_fontface = NULL,
  nudge_y = NULL,
  ymax = NULL,
  sign_thresh = NULL,
  label_color = NULL,
  color = NULL,
  legend_labels = NULL,
  annotate = NULL,
)
```

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Arguments

Dataframe, GWAS summary statistics genome_wide_thresh A number. P-value threshold for genome wide significant loci (5e-08 by default) suggestive_thresh A number. P-value threshold for suggestive loci (1e-06 by default) flank_size A number (default = 1e6). The size of the flanking region for the significant and suggestitve snps. A number (default = 1e6). The size of the region for gene annotation. Increase region_size this number for sparser annotation and decrease for denser annotation. sign_thresh_color A string or vector of strings to set the color/s of the significance threshold/s sign_thresh_label_size A number setting the text size of the label for the significance thresholds (default text size is 3.5) show_legend A logical scalar, set to FALSE to hide the legend (default : TRUE) label_fontface A string or a vector of strings. Label font "plain", "bold", "italic", "bold.italic" (ggrepel argument) nudge_y A number to horizontally adjust the starting position of each gene label (this is a ggrepel parameter) Integer, max of the y-axis, (default value: ymax=(max(-log10(df\$P))) + max(-log10(df\$P))) ymax * .2)) A number or vector of numbers, setting the horizontal significance threshold sign_thresh (default: sign_thresh=5e-8). Set to NULL to hide the significance threshold. label_color A string or a vector of strings. To change the color of the gene or variant labels color A string or a vector of strings, for setting the color of the datapoints on the plot legend_labels A string or vector of strings representing legend labels for the input datasets A number (p-value). Display annotation for variants with p-values below this annotate threshold Additional arguments passed to other plotting functions.

Value

ggplot object

Examples

```
## Not run:
manhattanExtra(df)
## End(Not run)
```

34 match_by_alleles

match	all	e	les

Match the variants in the snpset by their alleles

Description

```
match_alleles()
```

This method is deprecated and will be removed in future versions. use match_by_alleles instead.

Usage

```
match_alleles(df, verbose = F)
```

Arguments

df A dataframe that is in the snpset format (like returned by the get_snpset() func-

tion)

verbose A logical scalar (default: FALSE). Assign to TRUE to get information on which

alleles are matched and which are not.

Value

The input dataframe containing only those variants whith matched alleles in the snpset

Examples

```
## Not run:
match_alleles(df)
## End(Not run)
```

match_by_alleles

Match the variants in the snpset by their alleles

Description

```
match_by_alleles()
```

Usage

```
match_by_alleles(df, verbose = NULL, show_full_output = FALSE)
```

match_by_pos 35

Arguments

df A dataframe that is in the snpset format (like returned by the get_snpset func-

tion)

verbose A logical scalar (default: FALSE). Assign to TRUE to get information on which

alleles are matched and which are not.

show_full_output

A logical scalar (default:FALSE). Assign to TRUE to show the full output from

this function

Value

The input dataframe containing only those variants with matched alleles in the snpset

Examples

```
## Not run:
CD_UKBB_lead_snps <- get_lead_snps(CD_UKBB)
snpset <- get_snpset(CD_UKBB_lead_snps, CD_FINNGEN)
match_by_alleles(snpset$found)
## End(Not run)</pre>
```

match_by_pos

Get variants that overlap between two datasets

Description

```
match_by_pos()
```

Usage

```
match_by_pos(df1, df2, verbose = NULL, show_full_output = FALSE)
```

Arguments

df1 A dataframe of variants, has to contain CHROM and POS df2 A dataframe of variants, has to contain CHROM and POS

verbose A logical scalar (default: FALSE). Assign to TRUE to get information on which

alleles are matched and which are not.

show_full_output

A logical scalar (default:FALSE). Assign to TRUE to show the full output from

this function

Value

A list containing two dataframes, one of overlapping snps and the other snps not found in the second input dataset

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Examples

```
## Not run:
CD_UKBB_index_snps <- get_lead_snps(CD_UKBB)
match_by_pos(CD_UKBB_index_snps, CD_FINNGEN)
## End(Not run)</pre>
```

qqtopr

Create a quantile quantile (QQ) plot

Description

qqtopr() displays QQ plots for association data. Required parameter is at least one dataset (dataframe) containing the association data (with columns CHROM, POS, P

Usage

```
qqtopr(
  dat,
  scale = 1,
  n_{variants} = 0,
 breaks = 15,
  title = NULL,
  color = get_topr_colors(),
  size = 1,
  legend_name = "",
  legend_position = "right",
  legend_labels = NULL,
  axis_text_size = 11,
  axis_title_size = 12,
  title_text_size = 13,
  legend_title_size = 12,
  legend_text_size = 12,
  verbose = NULL,
  diagonal_line_color = "#808080"
)
```

Arguments

dat	Dataframe or a list of dataframes (required columns is P)) of association results.
scale	An integer, plot elements scale, default: 1
n_variants	An integer, total number of variants used in the study
breaks	A number setting the breaks for the axes
title	A string to set the plot title
color	A string or vector of strings setting the color's for the input datasets

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size A number or a vector of numbers, setting the size of the plot points (default:

size=1.2)

legend_name A string, use to change the name of the legend (default: None)

legend_position

A string, top,bottom,left or right

legend_labels A string or vector of strings representing legend labels for the input datasets

axis_text_size A number, size of the x and y axes tick labels (default: 12)

axis_title_size

A number, size of the x and y title labels (default: 12)

title_text_size

A number, size of the plot title (default: 13)

legend_title_size

A number, size of the legend title

 $legend_text_size$

A number, size of the legend text

verbose A logical scalar (default: NULL). Set to FALSE to suppress printed messages

diagonal_line_color

A string setting the color of the diagonal line on the plot

Value

ggplot

Examples

```
## Not run:
qqtopr(CD_UKBB)
## End(Not run)
```

R2_CD_UKBB

Example dataset including the R2 column for the locuszoom plot function

Description

The dataset is a subset of CD_UKBB and only includes variants above and near the IL23R gene on chromosome 1

Usage

R2_CD_UKBB

Format

A data frame with 329 rows and 5 variables:

CHROM Chromosome, written as for example chr1 or 1

POS genetic position of the variant

ID Variant identifier, e.g. rsid

P P-value from Plink run, additive model, regression model GLM_FIRTH

R2 variant correlation (r^2)

Source

A subset of the CD_UKBB dataset

regionplot

Create a regionplot

Description

regionplot() displays the association results for a smaller genetic regions within one chromosome. Required parameter is at least one dataset (dataframe) containing the association data (with columns CHROM, POS, P in upper or lowercase) and either a variant ID, gene name or the genetic region represented as a chromosome together with start and stop positions (either as a single string or as three separate arguments).

All other input parameters are optional

Usage

```
regionplot(
  df,
  ntop = 10,
  annotate = NULL,
  xmin = 0,
  size = 2,
  shape = 19,
  alpha = 1,
  label_size = 4,
  annotate_with = "ID",
  color = get_topr_colors(),
  axis_text_size = 11,
  axis_title_size = 12,
  title_text_size = 13,
  show_genes = NULL,
  show_overview = TRUE,
  show_exons = NULL,
 max\_genes = 200,
```

```
sign_{thresh} = 5e-08,
sign_thresh_color = "red",
sign_thresh_label_size = 3.5,
xmax = NULL,
ymin = NULL,
ymax = NULL,
protein_coding_only = FALSE,
region_size = 1e+06,
gene_padding = 1e+05,
angle = 0,
legend_title_size = 12,
legend_text_size = 11,
nudge_x = 0.01,
nudge_y = 0.01,
rsids = NULL,
variant = NULL,
rsids_color = NULL,
legend_name = "",
legend_position = "right",
chr = NULL,
vline = NULL,
show_gene_names = NULL,
legend_labels = NULL,
gene = NULL,
title = NULL,
label_color = NULL,
locuszoomplot = FALSE,
region = NULL,
legend_nrow = NULL,
gene_label_size = NULL,
scale = 1,
show_legend = TRUE,
sign_thresh_linetype = "dashed",
sign_thresh_size = 0.5,
rsids_with_vline = NULL,
annotate_with_vline = NULL,
show_gene_legend = TRUE,
unit_main = 7,
unit_gene = 2,
unit_overview = 1.25,
verbose = NULL,
gene_color = NULL,
segment.size = 0.2,
segment.color = "black",
segment.linetype = "solid",
max.overlaps = 10,
unit_ratios = NULL,
extract_plots = FALSE,
```

```
label_fontface = "plain",
label_family = "",
gene_label_fontface = "plain",
gene_label_family = "",
build = 38,
label_alpha = 1,
vline_color = "grey",
vline_linetype = "dashed",
vline_alpha = 1,
vline_size = 0.5,
log_trans_p = TRUE
```

Arguments

sign_thresh_color

df	Dataframe or a list of dataframes (required columns are CHROM, POS, P), in upper- or lowercase) of association results.
ntop	An integer, number of datasets (GWAS results) to show on the top plot
annotate	A number (p-value). Display annotation for variants with p-values below this threshold
xmin, xmax	Integer, setting the chromosomal range to display on the x-axis
size	A number or a vector of numbers, setting the size of the plot points (default: size=1.2)
shape	A number of a vector of numbers setting the shape of the plotted points
alpha	A number or a vector of numbers setting the transparency of the plotted points
label_size	An number to set the size of the plot labels (default: label_size=3)
annotate_with	A string. Annotate the variants with either Gene_Symbol or ID (default: "Gene_Symbol")
color	A string or a vector of strings, for setting the color of the datapoints on the plot
<pre>axis_text_size</pre>	A number, size of the x and y axes tick labels (default: 12)
axis_title_size	
	A number, size of the x and y title labels (default: 12)
title_text_size	
	A number, size of the plot title (default: 13)
show_genes	A logical scalar, show genes instead of exons (default show_genes=FALSE)
show_overview	A logical scalar, shows/hides the overview plot (default= TRUE)
show_exons	Deprecated : A logical scalar, show exons instead of genes (default show_exons=FALSE)
max_genes	An integer, only label the genes if they are fewer than max_genes (default values is 200).
sign_thresh	A number or vector of numbers, setting the horizontal significance threshold (default: sign_thresh=5e-8). Set to NULL to hide the significance threshold.

A string or vector of strings to set the color/s of the significance threshold/s

sign_thresh_label_size

A number setting the text size of the label for the significance thresholds (default

text size is 3.5)

ymin, ymax Integer, min and max of the y-axis, (default values: ymin=0, ymax=max(-log10(df\$P)))

protein_coding_only

A logical scalar, if TRUE, only protein coding genes are used for annotation

region_size An integer (default = 20000000) (or a string represented as 200kb or 2MB)

indicating the window size for variant labeling. Increase this number for sparser

annotation and decrease for denser annotation.

gene_padding An integer representing size of the region around the gene, if the gene argument

was used (default = 100000)

angle A number, the angle of the text label

legend_title_size

A number, size of the legend title

legend_text_size

A number, size of the legend text

nudge_x A number to vertically adjust the starting position of each gene label (this is a

ggrepel parameter)

nudge_y A number to horizontally adjust the starting position of each gene label (this is

a ggrepel parameter)

rsids A string (rsid) or vector of strings to highlight on the plot, e.g. rsids=c("rs1234,

rs45898")

variant A string representing the variant to zoom in on. Can be either an rsid, or a

dataframe (with the columns CHROM,POS,P)

rsids_color A string, the color of the variants in variants_id (default color is red)

legend_name A string, use to change the name of the legend (default: None)

legend_position

A string, top,bottom,left or right

chr A string or integer, the chromosome to plot (i.e. chr15), only required if the

input dataframe contains results from more than one chromosome

vline A number or vector of numbers to add a vertical line to the plot at a specific

chromosomal position, e.g vline=204000066. Multiple values can be provided

in a vector, e.g vline=c(204000066,100500188)

show_gene_names

A logical scalar, if set to TRUE, gene names are shown even though they exceed

the max genes count

legend_labels A string or vector of strings representing legend labels for the input datasets

gene A string representing the gene to zoom in on (e.g. gene=FTO)

title A string to set the plot title

label_color A string or a vector of strings. To change the color of the gene or variant labels

locuszoomplot A logical scalar set to FALSE. Only set to TRUE by calling the locuszoom

function

region A string representing a genetic region, e.g. chr1:67038906-67359979 An integer, sets the number of rows allowed for the legend labels legend_nrow gene_label_size A number setting the size of the gene labels shown at the bottom of the plot scale A number, to change the size of the title and axes labels and ticks at the same time (default: 1) A logical scalar, set to FALSE to hide the legend (default : TRUE) show_legend sign_thresh_linetype A string, the line-type of the horizontal significance threshold (default : dashed) sign_thresh_size A number, sets the size of the horizontal significance threshold line (default : 1) rsids_with_vline A string (rsid) or vector of strings to highlight on the plot with their rsids and vertical lines further highlighting their positions annotate_with_vline A number (p-value). Display annotation and vertical lines for variants with pvalues below this threshold show_gene_legend A logical scalar, set to FALSE to hide the gene legend (default value is TRUE) the height unit of the main plot (default = 7) unit_main unit_gene the height unit of the gene plot (default= 2) unit_overview the height unit of the overview plot (default = 1.25) Logical, set to FALSE to get suppress printed information verbose A string representing a color, can be used to change the color of the genes/exons gene_color on the geneplot segment.size line segment color (ggrepel argument) segment.color line segment thickness (ggrepel argument) segment.linetype line segment solid, dashed, etc.(ggrepel argument) max.overlaps Exclude text labels that overlap too many things. Defaults to 10 (ggrepel arguunit_ratios A string of three numbers separated by ":", for the overview, main and gene plots height ratios e.g 1.25:7:2 extract_plots Logical, FALSE by default. Set to TRUE to extract the three plots separately in a list label_fontface A string or a vector of strings. Label font "plain", "bold", "italic", "bold.italic" (ggrepel argument) label_family A string or a vector of strings. Label font name (default ggrepel argument is "") gene_label_fontface Gene label font "plain", "bold", "italic", "bold.italic" (ggrepel argument) gene_label_family

Gene label font name (default ggrepel argument is "")

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A number representing the genome build or a data frame. Set to 37 to change to build build (GRCh37). The default is build 38 (GRCh38). label_alpha An number or vector of numbers to set the transparency of the plot labels (default: label_alpha=1) vline_color A string. The color of added vertical line/s (default: grey) vline_linetype A string. The linetype of added vertical line/s (default : dashed) vline_alpha A number. The alpha of added vertical line/s (default: 1) vline_size A number. The size of added vertical line/s (default: 0.5) A logical scalar (default: TRUE). By default the p-values in the input datasets log_trans_p are log transformed using -log10. Set this argument to FALSE if the p-values in the datasets have already been log transformed.

Value

plots within ggplotGrobs, arranged with egg::gtable_frame

Examples

```
## Not run:
regionplot(CD_UKBB, gene="IL23R")
## End(Not run)
```

topr

topr

Description

A package for viewing and annotating genetic association data

topr functions

The main plotting functions are:

- manhattan to create Manhattan plot of association results
- regionplot to create regional plots of association results for smaller genetic regions

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See Also

Useful links:

- https://github.com/totajuliusd/topr
- Report bugs at https://github.com/totajuliusd/topr/issues

Examples

```
library(topr)
# Create a manhattan plot using
manhattan(CD_UKBB)

# Create a regional plot
regionplot(CD_UKBB, gene="IL23R")
```

UC_UKBB

UKBB Ulcerative colitis (ICD 10 code K51)

Description

Dataset retrieved from the UK biobank including of 5,452 UC cases (K51) and 481,862 controls. The dataset has been filtered on variants with P<1e-03.

Usage

UC_UKBB

Format

A data frame with 45,012 rows and 8 variables

CHROM Chromosome, written as for example chr1 or 1

POS genetic position of the variant

REF the reference allele

ALT the alternative allele

ID Variant identifier, e.g. rsid

P P-value from Plink run, additive model, regression model GLM_FIRTH

OR Odds Ratio

AF Allele frequency

Source

Ulcerative Colitis UKBB ICD10 code K51, only including variants with P<1e-03

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