

Package ‘DESNP’

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Type Package

Title Differentially Expressed Single Nucleotide Polymorphism

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Description Provides a framework for the identification and analysis of Differentially Expressed Single Nucleotide Polymorphisms (deSNPs) using high-throughput sequencing data. It enables users to import SNP count data from variant files, perform allele-specific read count extraction, and statistically detect SNPs showing significant differences in allele expression between biological conditions or sample groups. This package contains tools for calculating SNP-index and Delta SNP-index from VCF-derived allele depth data with statistical testing and filtering, including sliding-window analysis of genomic regions.

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Encoding UTF-8

Depends R (>= 4.1.0)

Imports vcfR, dplyr, tidyr, magrittr, tidyselect, VGAM, stats, utils,
GenomicRanges, IRanges, S4Vectors, GenomeInfoDb, tools, ggplot2

NeedsCompilation no

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DeltaSNPindex	<i>Compute Delta SNP-index</i>
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Description

DeltaSNPindex calculates the difference in SNP-index values between two experimental conditions using output from the SNPindex() function. The Delta SNP-index (Δ SNP-index) is computed as the subtraction of SNP-index values between groups for each SNP position, representing allele frequency changes between conditions. This approach is based on the QTL-seq strategy described by Takagi et al. (2013). The function also supports automatic group detection and optional threshold-based filtering to identify SNPs showing substantial allele frequency differentiation.

Usage

```
DeltaSNPindex(
  snpindex_input,
  meta_file = NULL,
  filter_deltaSNP = TRUE,
  deltaSNP_threshold = 0.5,
  filter_direction = c("both", "greater", "lesser"),
  column_filter = c("all", "any")
)
```

Arguments

`snpindex_input` Either:

- Path to a tab-delimited SNP-index file (e.g., output of SNPindex()), or
- A data frame containing SNP-index columns.

The input must contain columns beginning with SNPindex_.

`meta_file` Optional path to a metadata file describing condition groupings. The file must contain columns:

- `column` – SNP-index column name
- `condition` – condition label

Optional:

- `group` – allows multiple independent group comparisons

If NULL, automatic control detection mode is used.

`filter_deltaSNP` Logical; whether to apply threshold-based filtering. Default: TRUE.

deltaSNP_threshold Numeric threshold for filtering Δ SNP-index values. Default: 0.5.
filter_direction Filtering direction:
 "both" Select SNPs with $|\Delta$ SNP-index \geq threshold
 "greater" Select SNPs with Δ SNP-index \geq threshold
 "lesser" Select SNPs with Δ SNP-index \leq -threshold
column_filter Determines filtering behavior when multiple delta SNP-index columns exist:
 "any" SNP passes if any delta SNP-index column satisfies threshold
 "all" SNP passes only if all delta SNP-index columns satisfy threshold

Details

Δ SNP-index is calculated as:

$$\Delta SNPindex = SNPindex_{condition2} - SNPindex_{condition1}$$

Modes of operation:

1. Metadata mode

If `meta_file` is provided:

- If a group has exactly 2 conditions \rightarrow single Δ SNP-index column
- If more than 2 conditions \rightarrow all pairwise comparisons are generated

2. Auto mode

If `meta_file` = NULL, the function:

- Automatically detects control columns (control/ctrl/c)
- Computes treatment vs control Δ SNP-index
- Falls back to pairwise comparisons if no control is detected

The function returns all computed Δ SNP-index values and optionally filtered results.

Value

Returns a list containing:

all_delta_SNPs Data frame containing all computed Δ SNP-index values

filtered_delta_SNPs Filtered SNPs based on threshold

summary Summary statistics of the analysis

Author(s)

Sayanti Guha Majumdar, Saima Bano, Nandita Banerjee, Rahul Kumar Tiwari, Dipro Sinha, Subham Ghosh, Sanjeev Kumar

References

Takagi, H., Abe, A., Yoshida, K., Kosugi, S., Natsume, S., Mitsui, H., Uemura, A., Utsushi, H., Tamiru, M., Takahashi, R., Goto, K., Terauchi, R. (2013). QTL-seq: rapid mapping of quantitative trait loci in rice by whole genome resequencing of DNA from two bulked populations. *Plant Journal*, 74(1), 174–183. doi:10.1111/tpj.12105

Examples

```
snp_file <- system.file("extdata", "All_SNPs_Index.tsv",
                        package = "DESNP")

meta_file <- system.file("extdata", "meta_columns.tsv",
                        package = "DESNP")

result_deltasnpindex <- DeltaSNPindex(
  snpindex_input = snp_file,
  meta_file = meta_file,
  deltaSNP_threshold = 0,
  filter_direction = "both",
  column_filter = "any"
)

head(result_deltasnpindex$all_delta_SNPs)
head(result_deltasnpindex$filtered_delta_SNPs)
result_deltasnpindex$summary
```

DeltaSNPWindow

Sliding Window Analysis of Δ SNP-index

Description

DeltaSNPWindow compute Δ SNP-index values across chromosomes using an overlapping sliding-window approach to identify genomic regions exhibiting allele frequency differentiation between experimental conditions. The method follows the QTL-seq framework described by Takagi et al. (2013), which builds upon the SNP-index strategy introduced by Abe et al. (2012). The function computes the mean Δ SNP-index within each window to highlight consistent allele frequency differences and optionally applies a Wilcoxon signed-rank test to evaluate whether window-level deviations significantly differ from zero. Multiple testing correction is performed using the Benjamini–Hochberg method. Regions showing strong and statistically supported Δ SNP-index signals represent candidate loci enriched for biologically relevant allele frequency shifts.

Usage

```
DeltaSNPWindow(
  deltasnp_input,
  delta_cols = NULL,
  window_size = 1e6,
```

```
    step_size = 1e5,  
    run_wilcoxon = TRUE  
  )
```

Arguments

deltasnp_input	A file path (TSV format) or a data frame containing SNP data. The input must contain at least the following columns: CHROM (chromosome name) and POS (genomic position).
delta_cols	Character vector or numeric indices specifying Δ SNP-index columns. If NULL, columns containing both "delta" and "snp" (case-insensitive) and numeric values are automatically detected.
window_size	Size of sliding windows in base pairs. Default is 1e6 (1 Mb).
step_size	Step size for sliding windows in base pairs. Default is 1e5 (100 kb).
run_wilcoxon	Logical value indicating whether to perform Wilcoxon signed-rank test for each window. If TRUE, p-values and FDR-adjusted p-values (Benjamini-Hochberg method) are computed.

Details

The function:

- Generates sliding windows for each chromosome.
- Calculates mean Δ SNP-index within each window.
- Optionally performs Wilcoxon signed-rank test to test deviation from zero.
- Adjusts p-values using Benjamini-Hochberg FDR correction.

Value

A data frame containing:

- seqnames – Chromosome name
- start – Window start position
- end – Window end position
- width – Window width
- strand – Genomic strand
- Mean Δ SNP-index columns (prefixed with mean_)
- Wilcoxon p-values (if enabled)
- FDR-adjusted p-values (if enabled)

Author(s)

Sayanti Guha Majumdar, Saima Bano, Nandita Banerjee, Rahul Kumar Tiwari, Dipro Sinha, Subham Ghosh, Sanjeev Kumar

References

Abe, A., Kosugi, S., Yoshida, K., Natsume, S., Takagi, H., Kanzaki, H., Matsumura, H., Yoshida, K., Mitsuoka, C., Tamiru, M., Innan, H., Cano, L. M., Kamoun, S., Terauchi, R. (2012). Genome sequencing reveals agronomically important loci in rice using MutMap. *Nature Biotechnology*, 30(2), 174–178. doi:10.1038/nbt.2095

Takagi, H., Abe, A., Yoshida, K., Kosugi, S., Natsume, S., Mitsui, H., Uemura, A., Utsushi, H., Tamiru, M., Takahashi, R., Goto, K., Terauchi, R. (2013). QTL-seq: rapid mapping of quantitative trait loci in rice by whole genome resequencing of DNA from two bulked populations. *Plant Journal*, 74(1), 174–183. doi:10.1111/tpj.12105

Benjamini, Y., and Hochberg, Y. (1995). Controlling the false discovery rate: a practical and powerful approach to multiple testing. *Journal of the Royal Statistical Society: Series B*, 57(1), 289–300.

Examples

```
example_file <- system.file(
  "extdata",
  "All_delta_SNPs_Index.tsv",
  package = "DESNP"
)

window_result <- DeltaSNPWindow(
  deltasnp_input = example_file,
  window_size = 5e5,
  step_size = 1e5,
  run_wilcoxon = TRUE
)

head(window_result)
```

manhattan_plot

Genome-wide Manhattan Plot for SNP Index and Δ SNP Index

Description

Generates a customizable genome-wide Manhattan plot for SNP index (SNP_{index}) or Δ SNP index (delta_snp_index) values. The function automatically detects appropriate SNP index columns or allows the user to specify columns manually. It computes cumulative genomic positions across chromosomes and produces a ggplot object. Faceting is supported when multiple SNP index columns are plotted.

Usage

```
manhattan_plot(
  input_data,
  plot_cols = NULL,
  plot_title = "Genome-wide Manhattan Plot of SNP Index",
```

```

x_label = "Chromosome",
y_label = NULL,
y_limits = c(0, 1),
y_breaks = NULL,
x_expand = 0.01,
point_size = 0.7,
point_alpha = 0.7,
hline_value = 0.5,
hline_linetype = "dashed",
hline_color = "black",
title_size = 14,
title_face = "bold",
axis_title_size = 12,
axis_text_size = 8,
axis_text_angle = 40,
facet_ncol = 1,
facet_scales = "free_y"
)

```

Arguments

input_data	<p>Either a path to a tab-delimited SNP result file or a data frame containing SNP data.</p> <p>The data must contain:</p> <ul style="list-style-type: none"> • CHROM — Chromosome name • POS — Genomic position • One or more SNP index or ΔSNP index columns <p>Column names are case-insensitive for CHROM and POS.</p>
plot_cols	<p>Optional character vector specifying which columns to plot.</p> <p>If NULL (default):</p> <ul style="list-style-type: none"> • The function first searches for columns matching <code>^delta_snp_index(.*?)?\$</code> • If none found, searches for <code>^SNPindex_</code> • Delta SNP index columns are prioritized
plot_title	Main title of the Manhattan plot.
x_label	Label for x-axis.
y_label	Label for y-axis. Automatically set to Δ SNP index for delta columns, or "SNP index" for SNPindex columns if NULL.
y_limits	Numeric vector specifying y-axis limits. Default: <code>c(0, 1)</code>
y_breaks	Custom y-axis tick marks. If NULL, ggplot default is used.
x_expand	Expansion factor for x-axis spacing.
point_size	Size of plotted SNP points.
point_alpha	Transparency level of points (0–1).
hline_value	Y-value for horizontal reference line. Default: 0.5
hline_linetype	Line type for horizontal reference line.

<code>hline_color</code>	Color of horizontal reference line.
<code>title_size</code>	Font size of plot title.
<code>title_face</code>	Font face of title (e.g., "bold").
<code>axis_title_size</code>	Font size of axis titles.
<code>axis_text_size</code>	Font size of axis tick labels.
<code>axis_text_angle</code>	Rotation angle of chromosome labels.
<code>facet_ncol</code>	Number of columns when faceting multiple SNP index types.
<code>facet_scales</code>	Scaling for faceted plots ("fixed", "free", "free_y").

Details

Automatic Column Detection

If `plot_cols = NULL`, the function:

1. Searches for columns matching `^delta_snp_index(.*)*$`
2. If none found, searches for `^SNPindex_`
3. Stops if no matching columns are detected.

Cumulative Genome Position

To create a genome-wide Manhattan plot across multiple chromosomes:

- Chromosome lengths are calculated using maximum POS.
- Cumulative offsets are computed.
- SNP positions are shifted accordingly.

This ensures chromosomes appear sequentially along the x-axis.

Faceting

If more than one SNP index column is plotted:

- Separate panels are created using `facet_wrap()`
- Y-axis scaling can be controlled via `facet_scales`

Plot Output

The function **returns a ggplot object**. Use `ggsave()` if you want to save PNG, PDF, or TIFF files manually.

Value

A ggplot object representing the Manhattan plot.

Author(s)

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Examples

```
example_file <- system.file(
  "extdata",
  "All_delta_SNPs_Index.tsv",
  package = "DESNP"
)

manhattan_plot(input_data = example_file)

# Plot specific column
manhattan_plot(
  input_data = example_file,
  plot_cols = c("delta_snp_index_SNPindex_trt_200dai_vs_SNPindex_control_200dai")
)

# Custom y-limits and horizontal line
manhattan_plot(
  input_data = example_file,
  y_limits = c(-1, 1),
  hline_value = 0
)
```

SNPindex

SNP-index Calculation

Description

Computes SNP-index values from allele depth (AD) fields of the VCF file, performs statistical comparison between experimental conditions, and identifies significantly differentiated SNPs following the SNP-index framework described by Abe et al. (2012). The function supports beta-binomial regression and Fisher's exact test, with automatic test selection based on replicate availability.

Usage

```
SNPindex(
  vcf_file,
  metadata_file,
  min_depth = 10,
  test_method = c("auto", "beta_binomial", "fisher_exact"),
  PVALUE = 0.05,
  Pvalue_filter = c("all", "any")
)
```

Arguments

`vcf_file` Path to the input VCF file. The VCF file must contain allele depth (AD) values in the genotype (FORMAT) field.

`metadata_file` Path to a metadata file describing sample information. The file must be tab- or whitespace-separated with a header row.

Required columns:

- `sample` – Sample names exactly matching the genotype column names of the VCF file (case-sensitive).
- `condition` – Experimental group label (e.g., control, treatment).

Optional columns:

Any additional columns (e.g., `tissue`, `batch`, `replicate`) are automatically treated as grouping variables for stratified analysis.

Important notes:

- Sample names must match VCF file sample names exactly.
- At least two unique condition values are required.
- Mismatched samples will be removed during merging.

Example without grouping variables:

sample	condition
S1	control
S2	control
S3	treatment
S4	treatment

Example with grouping variables:

sample	condition	tissue	batch
S1	control	leaf	1
S2	control	root	1
S3	treatment	leaf	1
S4	treatment	root	2

`min_depth` Minimum total read depth (REF + ALT) required for a SNP to be retained. Default: 10.

`test_method` Statistical method used to compare allele frequencies between conditions.

"auto" Automatically selects beta-binomial if ≥ 2 replicates per condition are detected; otherwise Fisher's exact test is used.

"beta_binomial" Forces beta-binomial regression using `VGAM::vglm()`.

"fisher_exact" Forces Fisher's exact test using `stats::fisher.test()`.

`PVALUE` Significance threshold for identifying significant SNPs. Default: 0.05. Filtering is based on raw p-values.

`Pvalue_filter` Filtering behavior when grouping variables are present.

"all" SNP must be significant in all groups.

"any" SNP is significant if significant in at least one group.

Details

The SNP-index is calculated as:

$$SNPindex = ALT / (REF + ALT)$$

SNPs with total depth below `min_depth` are removed before testing.

Statistical testing strategy:

- If ≥ 2 biological replicates are detected per condition, beta-binomial regression is applied.
- If replicates are insufficient, Fisher's exact test is used.
- In "auto" mode, the test is selected automatically.

P-values are adjusted for multiple testing using the Benjamini–Hochberg (BH) method and reported as FDR.

Value

Returns a list containing:

all_SNPs Data frame of all analyzed SNPs including allele counts, SNP-index values, p-values, and FDR.

significant_SNPs Subset of SNPs passing the PVALUE threshold.

summary Summary statistics including total SNPs, number of significant SNPs, test used, depth threshold, and PVALUE.

Author(s)

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References

Abe, A., Kosugi, S., Yoshida, K., Natsume, S., Takagi, H., Kanzaki, H., Matsumura, H., Yoshida, K., Mitsuoka, C., Tamiru, M., Innan, H., Cano, L. M., Kamoun, S., & Terauchi, R. (2012). Genome sequencing reveals agronomically important loci in rice using MutMap. *Nature Biotechnology*, 30(2), 174–178. doi:10.1038/nbt.2095

Benjamini, Y., and Hochberg, Y. (1995). Controlling the false discovery rate. *Journal of the Royal Statistical Society: Series B*, 57(1), 289–300.

See Also

[read.vcfR](#), [vglm](#), [fisher.test](#)

Examples

```
vcf_path <- system.file("extdata", "combined_top10_new_smut.vcf",
                        package = "DESNP")

meta_path <- system.file("extdata", "metadata.tsv",
                        package = "DESNP")

result_snpindex <- SNPindex(
  vcf_file = vcf_path,
  metadata_file = meta_path,
  min_depth = 10,
  test_method = "auto",
  PVALUE = 0,
  Pvalue_filter = "any"
)

head(result_snpindex$all_SNPs)
head(result_snpindex$significant_SNPs)
result_snpindex$summary
```

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