# Package 'MutSeqR'

October 16, 2024

**Title** Analysis of error-corrected Next-Generation Sequencing Data for Mutation Detection **Version** 0.0.0.9000

**Description** Standard methods for analysis of mutation data following ecNGS for the purpose of mutagencity assessment. Functions include importing the mutation lists provided by a variant caller, and a set of analytical tools for statistical testing and visualization of mutation data; comparison to COSMIC and/or germline signatures; etc.

```
License MIT + file LICENSE
Depends R (>= 3.4.0)
Imports binom,
     BiocManager,
      Biostrings,
      BSgenome,
      car,
     data.table,
     doBy,
      dplyr,
     GenomeInfoDb,
      GenomicRanges,
      ggplot2,
     here,
     httr,
     IRanges,
     lme4,
     magrittr,
      openxlsx,
      patchwork,
      plyranges,
     reticulate,
      rlang,
      S4Vectors,
      stringr,
      SummarizedExperiment,
      VariantAnnotation,
```

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xml2
Suggests BSgenome.Hsapiens.UCSC.hg38,  BSgenome.Mmusculus.UCSC.mm10,  GenVisR,  knitr,  packcircles,  rmarkdown,  sigminer,  SigProfilerMatrixGeneratorR,  testthat (>= 3.0.0),  ToxicR,  trackViewer,  fs  Config/reticulate list( packages = list( list(package = ``SigProfilerAssignment"), list(package = ``SigProfilerExtractor"),
list(package = ``SigProfilerMatrixGenerator")))
Config/testthat/edition 3
Encoding UTF-8
<b>Roxygen</b> list(markdown = TRUE)
RoxygenNote 7.3.1
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add\_binom\_conf\_intervals

Add confidence intervals to mutation frequencies

# Description

Uses the binomial distribution to create confidence intervals for mutation frequencies calculated from a single point estimate using DNA sequencing

```
add_binom_conf_intervals(df, x, n, conf.level = 0.95, method = "wilson")
```

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## **Arguments**

df	The summary data frame containing the mutation frequencies
x	Column name that specifies the mutation count (e.g., mut_depth)
n	Column name that specifies the sequencing depth (e.g., total_depth)
conf.level	Confidence interval to calculate, default 95% (0.95)
method	The method used by binom::binom.confint to calculate intervals. Default is "wilson".

#### Value

A data frame with added columns indicating the confidence intervals.

annotate_CpG_sites	Annotate CpG sites

# Description

A simple method to test whether your trinucleotide context contains a CpG site. Vectorized version of Biostrings::vcountPattern is used.

## Usage

```
annotate_CpG_sites(mut_data, motif = "CG", column_query = "context", ...)
```

# Arguments

mut_data	A dataframe or GRanges object containing the genomic regions of interest in which to look for CpG sites.
motif	Default "CG", which returns CpG sites. You could in theory use an arbitrary string to look at different motifs. Use with caution. In this case the pattern being searched must be a column in the mutation data.
column_query	Default "context" but can be any column in the mutation data that you wish to look for a motif in.
	Additional arguments to vcountPattern()

# Value

A data frame with the same number of rows as there were ranges in the input, but with an additional metadata column indicating CpG sites in the target sequence of the mutation.

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bmd\_proast bmd\_proast

#### **Description**

Calculate the BMD of continuous, individual-level data. This function is an extension of the PROAST software (copyright RIVM National Institute for Public Health and the Environment).

## Usage

```
bmd_proast(
 mf_data,
  dose_col = "dose",
  response_col = "sample_MF_min",
  covariate_col = NULL,
  CES = 0.5,
  adjust_CES_to_group_SD = FALSE,
  model_averaging = TRUE,
  num\_bootstraps = 200,
  summary = TRUE,
  plot_results = FALSE,
  output_path = NULL,
  output_type = "none"
)
```

#### **Arguments**

mf\_data A data frame containing the data to be analyzed. This can be created using calculate\_mut\_freq(cols\_to\_group = "sample", retain\_metadata\_cols = [dose\_col], summary = TRUE). dose\_col The name of the column in mf\_data that contains the dose. Must be a numeric value. response\_col The name of the column(s) in mf\_data that contains the mutation frequency. Ex. c("sample\_MF\_min", "sample\_MF\_max"). The name of the column in mf\_data that contains the covariate. If no covariate covariate\_col is present, set to NULL. CES The critical effect size of the Benchmark response. The BMR in continuous data is defined as a CES-percent change in mean response (relative to the controls). Default is 0.5 (i.e. 50% change). adjust\_CES\_to\_group\_SD

A logical value indicating whether the group standard deviation should be used as the CES.

model\_averaging

A logical value indicating whether confidence intervals should be calculated using model averaging.

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num\_bootstraps The number of bootstrap resamples to be used in the model averaging. Default is 200.

summary A logical value indicating whether a summary of the results should be returned. If FALSE, raw results from the PROAST analysis are returned.

plot\_results A logical value indicating whether the results should be plotted. Plots can either be saved as a svg to the output\_path or displayed in the R plot viewer.

output\_path The filepath to save the plots' .svg files. If NULL, the plots will be saved in the current working directory.

output\_type svg or none. If svg, the plots will be saved as .svg files in the output\_path. If none, the plots will NOT be saved, but be displayed in plot viewer.

#### **Details**

This function is a modified version of the original interactive PROAST software (https://www.rivm.nl/en/proast that allows for batch processing of data. The function is designed to be used with the output of the calculate\_mut\_freq function for the purpose of calculating the Benchmark Dose of mutation frequency data. As such, some functionality of the original PROAST software has been removed.

This function will accept continuous data, with an observation for each individual subject. It is assumed that data are lognormally distributed. The response data is log-transformed, then backtransformed after the statistical analysis. The function will fit model 3 or 5 from various families of models (Exponential, Hill, Inverse Exponential, LogNormal). It will then compare the fits of models 3 and 5 for each model family and select the model with the lowest AIC. The BMD 90% confidence intervals will be calculated based on the selected model (3 or 5) for each model family using the profile likelihood method. The BMD 90% confidence interval may also be calculated using the bootstrap method if model\_averaging = TRUE. It is recommended to use 200 bootstraps for model averaging.

The function gives you the option to return either a summary of the results or the raw results output from the PROAST analysis. The summary will include the following for each response variable and covariate subgroup (if applicable):

- Selected.Model: The m3 or m5 model selected for each model family (Exponential, Hill, Inverse Exponential, LogNormal).
- Response: The response variable.
- Covariate: The covariate subgroup, if applicable.
- CES: The critical effect size of the Benchmark Response.
- CED: The Benchmark Dose, in original dose units, estimated for the given model.
- CEDL: The lower bound of the 90% confidence interval for the BMD, calculated by the profile likelihood method.
- CEDU: The upper bound of the 90% confidence interval for the BMD, calculated by the profile likelihood method.
- AIC: The Akaike Information Criterion for the selected model. Lower values indicate a better fit. It is advised to choose the BMD value from the model with the lowest AIC.
- Log.Likelihood: the log-likelihood of the given model.
- Var: The residual variance around the fitted curve on the natural log-scale.

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- a: Model Parameter the background response
- d: Model Parameter the slope of the curve.
- weights: The weight of the model in the model averaging process, if applicable.

If there is no significant response in the data, the function will return an empty data frame.

If plot\_results = TRUE the function will create the following plots for each response variable:

- Expon. The fitted curve of the selected (3 or 5) exponential model. Data is log-transformed. Individual data points are plotted using small triangles. The geometric mean (median) at each dose is plotted as a large triangle. The BMD is indicated by the dotted line. If applicable, the covariate subgroup is indicated by color.
- Hill. The fitted curve of the selected (3 or 5) Hill model. Data is log-transformed. Individual data points are plotted using small triangles. The geometric mean (median) at each dose is plotted as a large triangle. The BMD is indicated by the dotted line. If applicable, the covariate subgroup is indicated by color.
- InvExpon. The fitted curve of the selected (3 or 5) Inverse-exponential model. Data is log-transformed. Individual data points are plotted using small triangles. The geometric mean (median) at each dose is plotted as a large triangle. The BMD is indicated by the dotted line. If applicable, the covariate subgroup is indicated by color.
- LN. The fitted curve of the selected (3 or 5) LogNormal model. Data is log-transformed. Individual data points are plotted using small triangles. The geometric mean (median) at each dose is plotted as a large triangle. The BMD is indicated by the dotted line. If applicable, the covariate subgroup is indicated by color.
- ma.plot for model\_averaging = TRUE The bootstrap curves based on model averaging. The geometric mean (median) at each dose is plotted as a large triangle. Data is log-transformed.
- cleveland plot for model\_averaging = TRUE The BMD estimate for each model is plotted with error bars representing the 90% confidence interval. The size of the point represents the model weight.

To replicate these results in the PROAST interactive software, select the following menu options:

- 1. f.proast(mf data)
- 2. What type of response data do you want to consider? 1: continuous, individual data
- 3. Do you want to fit a single model or fit various nested families of models? 3: select model 3 or 5 from various families of models
- 4. Q1: Which variable do you want to consider as the independent variable? #: dose\_col
- 5. Give number(s) of the response(s) you want to analyse. #: response\_col
- 6. Give number of factor serving as potential covariate (e.g.sex) type 0 if none. #: covariate\_col
- 7. Do you want to adjust CES to within group SD? 1: no, 2: yes | adjust\_CES\_to\_group\_SD: FALSE/TRUE
- 8. Give value for CES (always positive) type 0 to avoid calculation of CIs. CES
- 9. Do you want to calculate the BMD confidence interval by model averaging? 1: no 2: yes | model\_averaging: FALSE/TRUE
- 10. give number of bootstrap runs for calculating BMD confidence interval based on MA (e.g. 200) *num\_bootstraps*
- 11. Which models do you want to be fitted? 4: previous option with lognormal DR model added

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## Value

If summary is TRUE, a data frame of final results. If summary is FALSE, a list of the raw results from the PROAST analysis.

bmd\_toxicr

Fit a BMD model for mutation frequency data

# Description

This function fits a continuous BMD model to dose-response data for mutation frequency.

# Usage

```
bmd_toxicr(
    mf_data,
    dose_col = "dose",
    response_col = c("sample_MF_min", "sample_MF_max"),
    model_types = c("hill", "exp-3", "exp-5", "power", "polynomial"),
    BMR = 0.5
)
```

# Arguments

mf_data	A data frame with columns "dose" and "MFmin"
dose_col	A character string specifying the column in data to be used to identify dose.
response_col	A character string specifying the column in data to be used to identify response.
model_types	A string specifying the model type. Options are "hill", "exp-3", "exp-5", "power", "polynomial"
BMR	A numeric value specifying the benchmark response. Default is 0.5.

## Value

A list with the following components:

bmd\_toxicr\_ma

bmd\_toxicr\_ma

Fit a model averaged continuous BMD model

# Description

This function fits a model averaged continuous BMD model to dose-response data for mutation frequency.

# Usage

```
bmd_toxicr_ma(
    mf_data,
    data_type = "individual",
    dose_col = "dose",
    response_cols = c("sample_MF_min", "sample_MF_max"),
    sd_col = NULL,
    n_col = NULL,
    bmr_type = "rel",
    bmr = 0.5,
    fit = "laplace",
    a = 0.025,
    ...
)
```

# Arguments

mf_data	A data frame containing the dose-response data. Data may be individual for each sample or averaged over dose groups. Required columns for individual data are "dose" and "response". Multiple response columns are allowed. Required columns for summarised data are "dose", "mean response", "sample size", and "standard deviation". Only one response column is allowed for summarised data.
data_type	A string specifying the type of response data. Data may be response per individual or summarised across dose groups. ("individual", "summary").
dose_col	A character string specifying the column in mf_data containing the dose data.
response_cols	A character vector specifying the columns in mf_data containing the response data. For summarised data types, this should be the mean response for each dose group.
sd_col	A character string specifying the column in mf_data containing the standard deviation of the response data. This is only required for summarised data types.
n_col	A character string specifying the column in mf_data containing the sample size of each dose group. This is only required for summarised data types.
bmr_type	A string specifying the type of benchmark response. For continuous models, there are four types of BMD definitions that are commonly used:

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> • Relative deviation (default; 'BMR\_TYPE = "rel"'). This defines the BMD as the dose that changes the control mean/median a certain percentage from the background dose, i.e. it is the dose, BMD that solves | f(dose)  $f(0) = (1 \pm BMR)f(0)$

- Standard deviation ('BMR\_TYPE = "sd"'). This defines the BMD as the dose associated with the mean/median changing a specified number of standard deviations from the mean at the control dose., i.e., it is the dose, BMD, that solves  $|f(dose) - f(0)| = BMR \times \sigma$
- Hybrid deviation ('BMR\_TYPE = "hybrid"'). This defines the BMD that changes the probability of an adverse event by a stated amount relative to no exposure (i.e 0). That is, it is the dose, BMD, that solves  $\frac{Pr(X>x|dose)-Pr(X>x|0)}{Pr(X<x|0)} =$ BMR. For this definition,  $Pr(X < x|0) = 1 - Pr(X > X|0) = \pi_0$ , where  $0 \le \pi_0 < 1$  is defined by the user as "point\_p," and it defaults to 0.01. Note: this discussion assumed increasing data. The fitter determines the direction of the data and inverts the probability statements for decreasing data.
- Absolute deviation ('BMR TYPE="abs"'). This defines the BMD as an absolute change from the control dose of zero by a specified amount. That is the BMD is the dose that solves the equation |f(dose) - f(0)| = BMR.

A numeric value specifying the benchmark response. The BMR is defined in bmr relation to the calculation requested in bmr\_type. Default is 0.5.

fit A string specifying the method used to fit the model. Options are ("laplace", "mle", or "mcmc"). Default is "laplace".

> The specified nominal coverage rate for computation of the lower bound on the BMDL and BMDU, i.e., one computes a  $100 \times (1 - \alpha)\%$  confidence interval. For the interval (BMDL,BMDU) this is a  $100 \times (1 - 2\alpha)\%$ . By default, it is set to 0.025 for a CI of 95%.

Additional arguments to be passed to the model fitting function. See ma continuous fit for details.

#### **Details**

Model averaging is done over the default models described in The European Food Safety Authority's (2022) Guidance on the use of the benchmark dose approach in risk assessment. These models are:

- "exp-aerts":  $f(x) = a(1 + (c-1)(1 \exp(-bx^d)))$
- "invexp-aerts":  $f(x) = a(1 + (c 1)(\exp(-bx^{-d})))$
- "hill-aerts":  $f(x) = a(1 + (c-1)(1 \frac{b^d}{b^d + x^d}))$
- "lognormal-aerts":  $f(x) = a \{1 + (c-1) (\Phi(\ln(b) + d \times \ln(x)))\}$
- "gamma-efsa":  $f(x) = a(1 + (c-1)(\Gamma(bx;d)))$
- "LMS":  $f(x) = a(1 + (c 1)(1 \exp(-bx dx^2)))$
- "probit-aerts":  $f(x) = c \left( \Phi(a + b \times x^d) \right)$
- "logistic-aerts":  $f(x) = \frac{c}{1 + \exp(-a b \times x^d)}$

Here:  $\Phi(\cdot)$  is the standard normal distribution and  $\Phi_{SN}(\cdot;\cdot)$  is the skew-normal distribution

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#### Value

A list with the following components:

• BMD: A data frame containing the BMD values and the  $100 \times (1-2\alpha)\%$  confidence intervals for each response column.

- summary: A list containing the summary statistics for each repsonse column.
- model\_plots: A list containing the model plots for each response column.
- cleveland\_plots: A list containing the Cleveland plots for each response column.
- models: A list containing the model fit values for each response column. Values include:
  - Individual\_Model\_X: Individual model fits for each model, X, used for model averaging.
     See single\_continuous\_fit for details on the model object class structure.
  - ma\_bmd: The CDF of the model averaged BMD distribution.
  - posterior\_probs: the posterior probabilities used in the model averaging.

calculate\_mut\_freq

Calculate mutation frequency

## **Description**

Calculates the mutation frequency for arbitrary groupings and creates a new dataframe with the results. Mutation frequency is # mutations / total bases, but this can be subset in different ways: e.g., by mutation context. In this case, it is necessary to change the denominator of total bases to reflect the sequencing depth at the proper reference bases under consideration.

# Usage

```
calculate_mut_freq(
  mutation_data,
  cols_to_group = "sample",
  subtype_resolution = "none",
  variant_types = c("snv", "deletion", "insertion", "complex", "mnv", "symbolic"),
  filter_germ = TRUE,
  summary = TRUE,
  retain_metadata_cols = NULL
)
```

#### **Arguments**

mutation data

The data frame to be processed containing mutation data. Required columns are listed below. Synonymous names for these columns are accepted.

- contig: The reference sequence name.
- start: 0-based start position of the feature in contig.
- sample: The sample name.
- alt\_depth: The read depth supporting the alternate allele.

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- total\_depth: The total read depth at this position (excluding N-calls).
- is\_germline: A logical variable indicating whether the mutation is a germline mutation.
- variation\_type: The category to which this variant is assigned.
- subtype\_col: The column containing the mutation subtype. This column depends on the subtype\_resolution parameter.
- reference\_col: The column containing the referene base(s) for the mutation. This column depends on the subtype\_resolution parameter.
- metadata\_cols for grouping: all columns across which you want to calculate the mutation frequency. Ex. c("tissue", "dose"). These columns should be listed in cols\_to\_group.

cols\_to\_group

A vector of grouping variables: this should be the groups of interest that you want to calculate a frequency for. For instance, getting the frequency by sample. Other options might include dose, locus, or, c("sample", "locus"). All listed variables must be a column in the mutation\_data.

subtype\_resolution

The resolution at which the frequencies are calculated. Options are

- "none" calculates mutation frequencies across all selected grouping columns. The reference\_col is not needed.
- "type" calculates mutation frequencies across all selected grouping columns for each variation\_type seperately; snv, mnv, deletion, insertion, complex, symbolic. The reference\_col is not needed.
- "base\_6" calculates mutation frequencies across all selected grouping columns for each variation\_type with snv mutations separated by normalized\_subtype; C>A, C>G, C>T, T>A, T>C, T>G. The reference\_col is normalized\_ref.
- "base\_12" calculates mutation frequencies across all selected grouping columns for each variation\_type with snv mutations separated by subtype; A>C, A>G, A>T, C>A, C>G, C>T, G>A, G>C, G>T, T>A, T>C, T>G. The reference\_col is short\_ref.
- "base\_96" calculates mutation frequencies across all selected grouping columns for each variation\_type with snv mutations separated by normalized\_context\_with\_mutation, i.e. the 96-base trinucleotide context. Ex. A\[C>T\]A. The reference\_col is normalized\_context.
- "base\_192" calculates mutation frequencies across all selected grouping columns for each variation\_type with snv mutations separated by context\_with\_mutation, i.e. the 192-base trinucleotide context. Ex A\[G>A\]A. The reference\_col is context.

variant\_types

Include these variant types in mutation counts. A vector of one or more variation\_types. Options are: "snv", "complex", "deletion", "insertion", "mnv", "symbolic", "no\_variant". Default includes all variants.

filter\_germ

A logical variable. If TRUE, exclude rows from the mutation count that were flagged as germline mutations in is\_germline. Default is TRUE.

summary

A logical variable, whether to return a summary table (i.e., where only relevant columns for frequencies and groupings are returned). Setting this to false returns all columns in the original mutation\_data, which might make plotting more difficult, but may provide additional flexibility to power users.

retain\_metadata\_cols

a character vector that contains the names of the metadata columns that you would like to retain in the summary table. This may be useful for plotting your summary data. Ex. retain the "dose" column when summarising by "sample". clonality\_cutoff NOT CURRENTLY IMPLEMENTED! Up for consideration. This value determines the fraction of reads that is considered a constitutional variant. If a mutation is present at a fraction higher than this value, the reference base will be swapped, and the alt\_depth recalculated. 0.3 (30%) would be a sane default?

#### **Details**

Additionally, by default, the operation is run by default using both the minimum and maximum independent methods for counting mutations.

#### Value

A data frame with the mutation frequency calculated. If summary is set to TRUE, the data frame will be a summary table with the mutation frequency calculated for each group. If summary is set to FALSE, the mutation frequency will be appended to each row of the original mutation\_data.

- \_MF\_min: The mutation frequency calculated using the "min" method for mutation counting. All identical mutations within a samples are assumed to be the result of clonal expansion and are thus only counted once.
- \_MF\_max: The mutation frequency calculated using the "max" method for mutation counting.
   All identical mutations within a sample are assumed to be idenpendent mutational evens and are included in the mutation frequency calculation. Note that this does not apply for germline variants.
- proportion\_min: The proportion of each mutation subtype within the group, normalized to its read depth. Calculated using the "min" method. This is only calculated if subtype\_resolution is not "none".
- proportion\_max: The proportion of each mutation subtype within the group, normalized to
  its read depth. Calculated using the "max" method. This is only calculated if subtype\_resolution
  is not "none".

check\_required\_columns

Check that all required columns are present before proceeding with the function

#### **Description**

A utility function that will check that all required columns are present.

## Usage

check\_required\_columns(data, required\_columns)

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#### **Arguments**

## Value

an error

cluster\_spectra

Hierarchical Clustering

## **Description**

perform hierarchical clustering of samples based on the mutation spectra.

## Usage

```
cluster_spectra(
   mf_data = mf_data,
   group_col = "sample",
   response_col = "proportion_min",
   subtype_col = "normalized_subtype",
   dist = "cosine",
   cluster_method = "ward.D"
)
```

# Arguments

mf_data	A data frame containing the mutation data. This data must include a column containing the mutation subtypes, a column containing the sample/cohort names, and a column containing the response variable.
group_col	The name of the column in data that contains the sample/cohort names.
response_col	The name of the column in data that contains the response variable. Typical response variables can be the subtype frequency, proportion, or count.
subtype_col	The name of the column in data that contains the mutation subtypes.
dist	the distance measure to be used. This must be one of "cosine", "euclidean", "maximum", "manhattan", "canberra", "binary" or "minkowski". See dist for details.
cluster_method	The agglomeration method to be used. See hclust for details.

## **Details**

The cosine distance measure represents the inverted cosine similarity between samples:

Cosine Dissimilarity = 
$$1 - \frac{\mathbf{A} \cdot \mathbf{B}}{\|\mathbf{A}\| \cdot \|\mathbf{B}\|}$$

This equation calculates the cosine dissimilarity between two vectors A and B.

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## Value

A dendrogram object representing the hierarchical clustering of the samples.

denominator\_dict

Values used for denominators in frequency calculations

## **Description**

These values are used to cross reference base substitution types to their appropriate denominators for calculations. That is", "for example, the 6 base substitution frequency should be subsetted based on the normalized\_ref column which would contain only T or C (i.e., the pyrimidine context for base substitutions).

# Usage

denominator\_dict

#### **Format**

A vector with corresponding values

f.proast

Run dose-response modeling using PROAST.

## **Description**

Run dose-response modeling using PROAST.

```
f.proast(
  odt = list(),
  ans.all = 0,
  er = FALSE,
  resize = FALSE,
  scale.ans = FALSE,
  const.var = F,
  show.warnings = F,
  interactive_mode = TRUE,
  datatype = NULL,
  model_choice = NULL,
  setting_choice = NULL,
  nested_model_choice = NULL,
  indep_var_choice = NULL,
  Vyans_input = NULL,
```

```
covariates = NULL,
custom_CES = 0.05,
model_selection = NULL,
lower_dd = NULL,
upper_dd = NULL,
selected_model = NULL,
adjust_CES_to_group_SD = NULL,
model_averaging = NULL,
num_bootstraps = NULL,
display_plots = TRUE
```

## Arguments

interactive\_mode

A TRUE/FALSE value specifying whether you want to run interactively (i.e., TRUE, the default) or using command-line mode (i.e., FALSE, non-interactive)

datatype Options: continuous, individual data.

#### Value

Results from PROAST.

```
generate_bubble_plots Generate Bubble Plots
```

## **Description**

Produces a ggplot object of bubble plots from given mutation data. Optionally, bubble plots can be facetted by a specified column.

#### Usage

```
generate_bubble_plots(
  mutation_data,
  facet_col = "dose",
  circle_spacing = 1,
  color_by = "normalized_subtype",
  circle_outline = "none",
  circle_resolution = 50
)
```

## **Arguments**

mutation\_data Data frame containing the mutation data.

facet\_col A string with the column name to facet by. If NULL or not provided, no faceting is performed.

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circle\_spacing Numerical value to adjust the spacing between circles.

color\_by Character vector specifying how to color the mutations. Accepted values are

"normalized\_subtype", "subtype", and "trinucleotide\_subtype". NOT FULLY

IMPLEMENTED.

circle\_outline Color for the circle outline. Default is "none", resulting in no outline color.

Other accepted values are colors in the R language.

circle\_resolution

Number of points to use for the circle resolution. Default is 50.

#### Value

A ggplot object with the bubble plot, facetted if specified.

get\_CpG\_mutations

Get mutations at CpG sites

#### **Description**

Subset the mutation data provided and return only mutations that are found at CpG sites.

## Usage

```
get_CpG_mutations(
  regions,
  mut_data,
  variant_types = c("snv", "insertion", "deletion", "mnv", "symbolic"),
  include_no_variants = T,
  motif = "CG"
)
```

#### Arguments

regions A GRanges object containing the genomic regions of interest in which to look

for CpG sites. Must have the metadata column "sequence" populated with the raw nucleotide sequence to search for CpGs. This object can be obtained using

the get\_seq.R function.

mut\_data A dataframe or GRanges object containing the mutation data to be interrogated.

tion", "insertion", "mnv", "symbolic", "no\_variant". Default includes all vari-

ants.

include\_no\_variants

TRUE or FALSE to indicate whether the table should include CpG sites with no variants. Useful if you want to know how many of the potential sites were

mutated.

motif Default "CG", which returns CpG sites. You could in theory use an arbitrary

string to look at different motifs. Use with caution.

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#### Value

A GRanges object where each range is a mutation at a CpG site (a subset of mutations from the larger object provided to the function).

get\_CpG\_regions

Get the coordinates of the CpG sites within your genomic regions

## **Description**

Filters the ranges of your genomic regions to find a positions with a specific motif. The default is CpG sites, but can be customizable.

## Usage

```
get_CpG_regions(regions, motif = "CG")
```

## **Arguments**

regions A GRanges object containing the genomic regions of interest in which to look

for CpG sites. Must have the metadata column "sequence" populated with the raw nucleotide sequence to search for CpGs. This object can be obtained using

the get\_seq.R function.

motif Default "CG", which returns CpG sites. You could in theory use an arbitrary

string to look at different motifs. Use with caution.

#### Value

A GRanges object where each range is a CpG site (a subset of ranges from the larger object provided to the function).

get\_ref\_of\_mut

A utility function that will return the reference context of a mutation

# Description

A utility function that will return the reference context of a mutation

#### Usage

```
get_ref_of_mut(mut_string)
```

## **Arguments**

```
mut_string the mutation. Ex. T>C, A[G>T]C
```

get\_region\_seqs 19

get\_region\_seqs

Get sequence of Duplex Sequencing target regions

## **Description**

This is mostly a helper function. It imports package data to find target regions and some associated information, and further extends the table by getting raw nucleotide sequences for each region of the genome. Note that the way this is written, currently, the default genomes are hg38 and mm10 for human and mouse, respectively.

### Usage

```
get_region_seqs(species)
```

## **Arguments**

species

One of "mouse" or "human", to determine which regions to return.

#### Value

A GRanges object where each range is a target region.

get\_seq

Get sequence of genomic target regions

## **Description**

Create a GRanges object from the target metadata and import raw nucleotide sequences from the UCSC database. https://genome.ucsc.edu

```
get_seq(
  regions = c("TSpanel_human", "TSpanel_mouse", "TSpanel_rat", "custom_interval"),
  custom_regions_file = NULL,
  rg_sep = "\t",
  genome = NULL,
  is_0_based = TRUE,
  padding = 0
)
```

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#### **Arguments**

regions "TSpanel\_human", "TSpanel\_mouse", "TSpanel\_rat, or "custom\_interval". The

argument refers to the TS Mutagenesis panel of the specified species, or to a

custom panel. If custom, provide file path in custom\_regions\_file.

custom\_regions\_file

"filepath". If regions is set to custom\_interval, provide the file path for the

tab-delimited file containing regions metadata. Required columns are "contig",

"start", and "end".

rg\_sep The delimiter for importing the custom\_regions\_file. The default is tab-delimited.

genome If a custom regions file is provided, indicate the genome assembly for the refer-

ence genome. Please refer to the UCSC genomes. Ex.Human GRCh38 = hg38 | Human GRCh37 = hg19 | Mouse GRCm38 = mm10 | Mouse GRCm39 = mm39

| Rat RGSC 6.0 = rn6 | Rat mRatBN7.2 = rn7

is\_0\_based TRUE or FALSE. Indicates whether the target region coordinates are 0 based

(TRUE) or 1 based (FALSE). If TRUE, ranges will be converted to 1-based.

padding An integer value by which the function will extend the range of the target se-

quence on both sides. Modified region ranges will be reported in seq\_start and

seq\_end. Default is 0.

#### Value

a GRanges object with sequences and metadata of targeted regions. Region ranges coordinates will become 1-based.

import\_mut\_data

Import a .mut file

## Description

Imports a .mut file into the local R environment.

```
import_mut_data(
   mut_file,
   mut_sep = "\t",
   rsids = FALSE,
   sample_data_file = NULL,
   sd_sep = "\t",
   vaf_cutoff,
   range_buffer = 0,
   regions = c("TSpanel_human", "TSpanel_mouse", "TSpanel_rat", "custom_interval", "none"),
   custom_regions_file = NULL,
   rg_sep = "\t",
   is_0_based = TRUE,
```

import\_mut\_data 21

```
depth_calc = "take_del",
  custom_column_names = NULL,
  output_granges = FALSE
)
```

#### **Arguments**

mut\_file

"filepath". The .mut file containing mutation data to be imported. This can be either a data frame object or a filepath to a file or directory. If you specify a folder, the function will attempt to read all files in the folder and combine them into a single data frame. Required columns are listed below. Synonymous names for these columns are accepted.

- contig: The reference sequence name.
- start: 0-based start position of the feature in contig.
- end: half-open end position of the feature in contig.
- sample: The sample name.
- ref: The reference allele at this position
- alt: The left-aligned, normalized, alternate allele at this position.
- alt\_depth: The read depth supporting the alternate allele.
- depth col: The total read depth at this position. This column can be total\_depth (excluding N-calls) or depth(including N-calls; if total\_depth is not available).
- variation\_type: The category to which this variant is assigned.
- context: The local reference trinucleotide context at this position (e.g. ATC not necessarily the transcript codon)

mut\_sep

The delimiter for importing the .mut file. Default is tab-delimited.

rsids

A logical variable; whether or not the .mut file contains rsID information (existing SNPs).

sample\_data\_file

An optional file containing additional sample metadata (dose, tissue, timepoint, etc.). This can be either a data frame object or a file path to a file.

sd\_sep

The delimiter for importing sample metadata table. Default is tab-delimited.

vaf\_cutoff

The function will add is\_germline column that identifies ostensibly germline variants using a cutoff for variant allele fraction (VAF). There is no default value provided, but generally a value of 0.1 (i.e., 10%) is a good starting point. Setting this will flag variants that are present at a frequency greater than this value at a given site.

 $range\_buffer$ 

An integer >= 0. Variants that occur outside of the defined regions' ranges will be filtered out. Use the range-buffer to extend the range within which a variant can occur. The default is 1 nucleotide outside of region ranges. Ex. Structural variants and indels may start outside of the regions. Adjust the range\_buffer to include these variants. Rows that were filtered out of the mutation data will be returned in a separate data frame.

regions

Values are c("TSpanel\_human", "TSpanel\_mouse", "TSpanel\_rat" "custom\_interval", "none"). Indicates the target panel used for Duplex Sequencing. The argument refers to

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the TS Mutagenesis panel of the specified species, or to a custom panel. If "custom", provide the file path of your regions file in custom\_regions\_file.

custom\_regions\_file

"filepath". If regions is set to "custom\_interval", provide the file containing regions metadata. Required columns are contig, start, and end. This can be either a data frame object or a file path to a file.

rg\_sep

The delimiter for importing the custom\_regions\_file. Default is tab-delimited.

is\_0\_based

A logical variable. Indicates whether the custom\_regions\_file target region coordinates are 0 based (TRUE) or 1 based (FALSE). If TRUE, ranges will be converted to 1-based.

depth\_calc

Values are c("take\_del", "take\_mean"). In the instance when there are two or more calls at the same location within a sample, and the depths differ, this parameter chooses the method for resolving the difference. This occurs when a deletion is called in the data. It will be called alongside a no\_variant. "take\_mean" calculates the depth column by taking the mean of all depths in the group. "take\_del" calculates the depth column by choosing only the depth of the deletion in the group, or if no deletion is present, the complex variant. If there is no deletion or complex variant, then it takes the mean of the depths within the group. Default is "take\_del". depth\_col = total\_depth or depth.

custom\_column\_names

A list of names to specify the meaning of column headers. Since column names can vary with data, this might be necessary to digest the mutation data table properly. Typical defaults are set, but can be substituted in the form of list(total\_depth = "my\_custom\_depth\_name", sample = "my\_custom\_sample\_column\_name"). For a comprehensive list, see examples. You can change one or more of these.

output\_granges

A logical variable; whether you want the mutation data to output as a GRanges object. Default output (FALSE) is as a dataframe.

#### Value

A table where each row is a mutation, and columns indicate the location, type, and other data. If output\_granges is set to TRUE, the mutation data will be returned as a GRanges object, otherwise mutation data is returned as a dataframe. If the mutation data contains rows that are filtered out of the specified regions, results will be returned as a list. The first element will be the mutation data called mut\_dat and the second element will be the rows that were filtered from the data called rows\_outside\_regions. If no rows are filtered out, the results will be returned as a single dataframe or GRanges object.

# Output Column Definitions:

- nchar\_ref: The length (in bp) of the reference allele.
- nchar\_alt: The length (in bp) of the alternate allele.
- varlen: The length (in bp) of the variant.
- total\_depth: The total read depth at this position, excluding N-calls.
- vaf: The variant allele fraction. Calculated as alt\_depth/depth\_col where depth\_col can be total\_depth or depth.
- is\_germline: TRUE or FALSE. Flags ostensible germline mutations (vaf > vaf\_cutoff).

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• ref\_depth: The total read depth at the position calling for the reference allele. Calculated as depth\_col - alt\_depth where depth\_col can be total\_depthor depth.

- subtype: The substitution type for the snv variant (12-base spectrum; e.g. A>C)
- short\_ref: The reference base at this position.
- normalized\_subtype: The C/T-based substitution type for the snv variant (6-base spectrum; e.g. A>C -> T>G).
- normalized\_ref: The reference base in C/T-base notation for this position (e.g. A -> T).
- context\_with\_mutation: The substitution type fo the snv variant including the two flanking nucleotides (192-trinucleotide spectrum; e.g. T[A>C]G)
- normalized\_context\_with\_mutation: The C/T-based substitution type for the snv variant including the two flanking nucleotide (96-base spectrum e.g. T[A>C]G -> C[T>G]A)
- normalized\_context: The trinucleotide context in C/T base notation for this position (e.g. TAG -> CTA).
- gc\_content: % GC of the trinucleotide context at this position.

import\_vcf\_data

Import a vcf file

# **Description**

The function reads the genomic vcf file(s) and extracts the data into a dataframe. The function also reads in sample metadata if provided and joins it with the mutation data. An interval list of genomic regions can be provided to filter out variants that occur outside of the defined regions' ranges. The function will use the reference genome to extract the trinucleotide context of every position in the mutation data. The function can output the mutation data as a dataframe or a granges object.

```
import_vcf_data(
    vcf_file,
    vaf_cutoff,
    range_buffer = 1,
    sample_data_file = NULL,
    sd_sep = "\t",
    regions = c("TSpanel_human", "TSpanel_mouse", "TSpanel_rat", "custom_interval", "none"),
    custom_regions_file = NULL,
    rg_sep = "\t",
    genome = NULL,
    species = NULL,
    masked_BS_genome = FALSE,
    depth_calc = "take_del",
    output_granges = FALSE
)
```

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#### **Arguments**

vcf\_file

The path to the genomic .vcf or .vcf.gz file(s) to be imported. If you specify a folder, the function will attempt to read all files in the folder and combine them into on dataset. Multisample vcf files are not supported; vcf files must contain one sample each. Required fields are listed below

- FIXED FIELDS:
- CHROM: The reference sequence name. Equivalent to contig
- POS: 0-based start position of the feature in contig.
- REF: The reference allele at this position
- ALT: The left-aligned, normalized, alternate allele at this position.
- FORMAT FIELDS:
- AD: The allelic depths for the reference and alternate alleles in the order listed.
- DP: The total read depth at this position (including N-calls). Equivalent to depth.
- VD: Variant Depth. Equivalent to alt\_depth.
- INFO FIELDS
- TYPE: The category to which this variant is assigned. Equivalent to variation\_type.
- END: The half-open end position of the feature in contig.
- sample: An identifying field for your samples; either in the INFO field or as the header to the FORMAT field.
- SUGGESTED INFO FIELDS:
- SVTYPE: Structural variant types; INV DUP DEL INS FUS.
- SVLEN: Length of the structural variant in base pairs.

vaf\_cutoff

Add is\_germline column that identifies ostensibly germline variants using a cutoff for variant allele fraction (VAF). There is no default value provided, but generally a value of 0.01 - 0.1 (i.e., 1% - 10%) is a good starting point. Setting this flag variants that are present at a frequency greater than this value at a given site.

range\_buffer

An integer >= 0 .Required if using a targetted approach. Variants that occur outside of the defined regions' ranges will be filtered out. Use the range-buffer to extend the range outside of a region within which a variant can occur. The default is 1 nucleotide outside of region ranges. Ex. Structural variants and indels may start outside of the regions. Adjust the range\_buffer to include these variants in the final mutation data. Variants that are filtered out of the data are returned in a separate dataframe.

sample\_data\_file

An optional file containing additional sample metadata (dose, timepoint, etc.). This can be a data frame or a file path.

sd\_sep

The delimiter for importing sample metadata tables. Default is tab-delimited

regions

"TSpanel\_human", "TSpanel\_mouse", "TSpanel\_rat", "custom\_interval" or "none". The 'TSpanel\_' argument refers to the TS Mutagenesis panel of the specified species, or to a custom regions interval file. If set to 'custom\_interval', please provide the file path in custom\_regions\_file and the genome assembly version

install\_ref\_genome 25

of the reference genome using the 'genome' parameter. If you are not using a targeted approach, set regions to none, and supply the species and genome assembly of the reference genome using the 'species' and 'genome' parameters respectively.

custom\_regions\_file

"filepath". If regions is set to custom\_interval, provide the file path for the file containing regions metadata. Required columns are "contig", "start", and "end"

rg\_sep The delimiter for importing the custom\_regions\_file. Default is tab-delimited

genome The genome assembly of the reference genome. For a ####### complete list,

refer to https://genome.ucsc.edu. Ex.Human GRCh38 = hg38 | Human GRCh37 = hg19 | Mouse GRCm38 = mm10 | Mouse GRCm39 = mm39 | Rat RGSC 6.0

 $= rn6 \mid Rat \ mRatBN7.2 = rn7$ 

species The species of the reference genome. Required if regions is set to none. The

value can be the common name of the species or the scientific name. Ex. "hu-

man" or "Homo sapiens".

sample, and the depths differ, this parameter chooses the method of calculation for the total\_depth. take\_mean calculates the total\_depth by taking the mean reference depth and then adding all the alt depths. take\_del calculates the total\_depth by choosing only the reference depth of the deletion in the group, or if no deletion is present, the complex variant, then adding all alt depths. If there is no deletion or complex variant, it takes the mean of the reference depths. Default

is "take\_del".

output\_granges TRUE or FALSE; whether you want the mutation data to output as a GRanges

object. Default output is as a dataframe.

## Value

A data frame or a GRanges object where each row is a mutation, and columns indicate the location, type, and other data.

install\_ref\_genome

Install the reference genome for the specified organism.

## **Description**

This function will use BSgenome to install the reference genome for a specified organism and assembly version.

```
install_ref_genome(organism, genome, masked = FALSE)
```

26 load\_regions\_file

# Arguments

organism the name of the organism for which to install the reference genome. This can be

the scientific name or a common name. For example Homo Sapiens, H. sapiens,

or human

genome The reference genome assembly version. Ex. hg18, mm10, rn6.

masked Logical value. Whether to search for the 'masked' BSgenome. Default is

FALSE.

#### Value

a BSgenome object

load\_regions\_file

Imports the regions file

# **Description**

A helper function to import the regions metadata file. It is used in import\_mut\_data and get\_seq.

## Usage

```
load_regions_file(regions, custom_regions_file = NULL, rg_sep = "\t")
```

## **Arguments**

regions "TSpanel\_human", "TSpanel\_mouse", 'TSpanel\_rat', or "custom\_interval". The

argument refers to the TS Mutagenesis panel of the specified species, or to a cus-

tom panel. If custom, provide file path in custom\_regions\_file.

custom\_regions\_file

"filepath". If regions is set to 'custom\_interval', provide the file path for the

tab-delimited file containing regions metadata. Required columns are "contig",

"start", and "end".

rg\_sep The delimiter for importing the custom\_regions\_file. The default is tab-delimited

lollipop\_mutations 27

#### **Description**

TO DO: Create plt without trackViewer package. Uses the trackViewer package to plot mutations in a lollipop plot in specific regions as defined by the user input.

## Usage

```
lollipop_mutations(species = "human", mutations, ...)
```

#### **Arguments**

species One of "human" or "mouse"

mutations A GRanges object with mutation data

... Additional arguments to trackViewer::lolliplot (e.g., ranges = GRanges ("chr1",

IRanges (104, 109)) ) Suggests track Viewer lolliplot

make\_CpG\_summary\_table

Summarize CpG sites

# Description

Creates a summary table of CpG sites based on groupings of interest. This is basically a convenience function that wraps calculate\_mut\_freqs() over CpG data (or any data). See the documentation for that function for parameters. It is up to the user to supply proper data to the function.

## Usage

```
make_CpG_summary_table(cpg_muts, ...)
```

## Arguments

cpg\_muts A data frame containing CpG mutations TO DO: cpg\_muts = df "cpg\_mutations"

is created in the .Rmd file, but is not created by any other function.

. . . Additional arguments to calculate\_mut\_freqs()

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migrate\_mut

Validate a mut file

# Description

Not tested yet. TODO! Could avoid data.table dependency here I think.

## Usage

```
migrate_mut(mut_table, op = MutSeqR::op)
```

## **Arguments**

mut\_table mutation data

op Default is MutSeqR::op, a list of options (see also, op, TODO)

model\_mf

Perform linear modelling on mutation frequency for given fixed and random effects

#### **Description**

model\_mf will fit a linear model to analyse the effect(s) of given factor(s) on mutation frequency and perform specified pairwise comparisons. This function will fit either a generalized linear model (glm) or, if supplied random effects, a generalized linear mixed-effects model (glmer) . Pairwise comparisons are conducted using the doBy library (esticon) and estimates are then backtransformed. The delta method is employed to approximate the back-transformed standard-errors. A Sidak correction is applied to adjust p-values for multiple comparisons.

```
model_mf(
    mf_data,
    fixed_effects,
    test_interaction = TRUE,
    random_effects = NULL,
    reference_level,
    muts = "sample_sum_min",
    total_count = "sample_group_depth",
    contrasts = NULL,
    cont_sep = "\t",
    ...
)
```

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#### **Arguments**

mf\_data The data frame containing the mutation frequency data. Mutation counts and

total sequencing depth should be summarized per sample alongside columns for

your fixed effects. This data can be obtained using calculate\_mut\_freq(summary=TRUE).

 $\label{lem:fixed_effects} \textbf{The name}(s) \ of \ the \ column(s) \ that \ will \ act \ as \ the \ fixed\_effects \ (factor/independent) \ act \$ 

variable) for modelling mutation frequency.

test\_interaction

a logical value. Whether or not your model should include the interaction between the fixed\_effects.

random\_effects The name of the column(s) to be analysed as a random effect in the model.

Providing this effect will cause the function to fit a generalized linear mixed-

effects model.

reference\_level

Refers to one of the levels within each of your fixed\_effects. The coefficient for the reference level will represent the baseline effect. The coefficients of the other levels will be interpreted in relation to the reference\_level as deviations

from the baseline effect.

muts The column containing the mutation count per sample.

total\_count The column containing the sequencing depth per sample.

contrasts a data frame or a filepath to a file that will provide the information necessary

to make pairwise comparisons between groups. The table must consist of two columns. The first column will be a group within your fixed\_effects and the second column must be the group that it will be compared to. The values must correspond to entries in your mf\_data column for each fixed effect. Put the group that you expect to have the higher mutation frequency in the 1st column and the group that you expect to have a lower mutation frequency in the second column. For multiple fixed effects, separate the levels of each fixed\_effect of a group with a colon. Ensure that all fixed\_effects are represented in each entry for

the table. See details for examples.

cont\_sep The delimiter for importing the contrast table file. Default is tab-delimited.

... Extra arguments for glm or glmer. The glmer function is used when a random\_effect

is supplied, otherwise, the model uses the glm function.

#### **Details**

fixed\_effects are variables that have a direct and constant effect on the dependent variable (ie mutation frequency). They are typically the experimental factors or covariates of interest for their impact on the dependent variable. One or more fixed\_effect may be provided. If you are providing more than one fixed effect, avoid using correlated variables; each fixed effect must independently predict the dependent variable. Ex. fixed\_effects = c("dose", "genomic\_target", "tissue", "age", etc).

Interaction terms enable you to examine whether the relationship between the dependent and independent variable changes based on the value of another independent variable. In other words, if an interaction is significant, then the relationship between the fixed effects is not constant across all levels of each variable. Ex. Consider investigating the effect of dose group and tissue on mutation

30 model\_mf

frequency. An interaction between dose and tissue would capture whether the dose response differs between tissues.

random\_effects account for the unmeasured sources of statistical variance that affect certain groups in the data. They help account for unobserved heterogeneity or correlation within groups. Ex. If your model uses repeated measures within a sample, random\_effects = "sample".

Setting a reference\_level for your fixed effects enhances the interpretability of the model. Ex. Consider a fixed\_effect "dose" with levels 0, 25, 50, and 100 mg/kg. Intuitively, the reference\_level would refer to the negative control dose, "0" since we are interested in testing how the treatment might change mutation frequency relative to the control.

## Examples of contrasts:

If you have a fixed\_effect "dose" with dose groups 0, 25, 50, 100, then the first column would contain the treated groups (25, 50, 100), while the second column would be 0, thus comparing each treated group to the control group.

250

500

1000

Alternatively, if you would like to compare mutation frequency between treated dose groups, then the contrast table would look as follows, with the lower dose always in the second column, as we expect it to have a lower mutation frequency. Keeping this format aids in interpretability of the estimates for the pairwise comparisons. Should the columns be reversed, with the higher group in the second column, then the model will compute the fold-decrease instead of the fold-increase.

100 25

100 50

50 25

Ex. Consider the scenario where the fixed\_effects are "dose" (0, 25, 50, 100) and "genomic\_target" ("chr1", "chr2"). To compare the three treated dose groups to the control for each genomic target, the contrast table would look like:

25:chr1 0:chr1

50:chr1 0:chr1

100:chr1 0:chr1

25:chr2 0:chr2

50:chr2 0:chr2

100:chr2 0:chr2

Troubleshooting: If you are having issues with convergence for your generalized linear mixed- effects model, it may be advisable to increase the tolerance level for convergence checking during model fitting. This is done through the control argument for the lme4::glmer function. The default tolerance is tol = 0.002. Add this argument as an extra argument in the model\_mf function. Ex. control = lme4::glmerControl(check.conv.grad = lme4::.makeCC("warning", tol = 3e-3, relTol = NULL))

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#### Value

Model results are output as a list. Included are:

• model\_data: the supplied mf\_data with added column for model residuals.

- summary: the summary of the model.
- anova: the analysis of variance for models with two or more effects. Anova (model)
- residuals\_histogram: the model residuals plotted as a histogram. This is used to check whether the variance is normally distributed. A symmetric bell-shaped histogram, evenly distributed around zero indicates that the normality assumption is likely to be true.
- residuals\_qq\_plot: the model residuals plotted in a quantile-quantile plot. For a normal distribution, we expect points to roughly follow the y=x line.
- point\_estimates\_matrix: the contrast matrix used to generate point-estimates for the fixed effects.
- point\_estimates: the point estimates for the fixed effects.
- pairwise\_comparisons\_matrix: the contrast matrix used to conduct the pairwise comparisons specified in the contrasts.
- pairwise\_comparisons: the results of pairwise comparisons specified in the contrasts.

ор

Column names for mut tables

## **Description**

A list of column specifications

## Usage

op

#### **Format**

A list with potential variable column names

32 plot\_mean\_mf

plot\_CI

plot\_CI

## Description

Plot confidence intervals

## Usage

```
plot_CI(data, order = "none", custom_order = NULL)
```

## **Arguments**

data A data frame with the results of the BMD analysis. Data must contain columns

"Response", "BMD", "BMDL", and "BMDU". BMD values can be NA.

order Indicates how the responses should be ordered. Options are "none" (default),

"asc" for ascending BMD values, "desc" for descending BMD values, or a cus-

tom order.

custom\_order A character vector with the custom order of the Responses.

#### Value

a ggplot object

plot\_mean\_mf

Plot the Mean Mutatation Frequency

## **Description**

This function calculates the mean mutation frequency across samples for given groups and plots the results.

```
plot_mean_mf(
    mf_data,
    group_col = "dose",
    mf_type = "both",
    plot_type = "line",
    plot_error_bars = TRUE,
    plot_indiv_vals = TRUE,
    group_order = "none",
    group_order_input = NULL,
    add_labels = "mean_count",
    scale_y_axis = "linear",
```

plot\_mean\_mf 33

```
x_lab = NULL,
y_lab = NULL,
plot_title = NULL,
custom_palette = NULL)
```

## Arguments

mf\_data A data frame containing the mutation frequency data. This is obtained from the

calculate\_mut\_freq function with SUMMARY = TRUE.

group\_col The column in mf\_data by which to calculate the mean. Ex. "dose" or c("dose",

"tissue").

mf\_type The type of mutation frequency to plot. Options are "min", "max", "both", or

"stacked". If "both", the min and max mutation frequencies are plotted side by side. "stacked" can be chosen for bar plot\_type only. If "stacked", the difference between the min and max MF is stacked on top of the min MF such that the total

height of both bars represent the max MF. Default is "min".

plot\_type The type of plot to create. Options are "bar" or "line". Default is "bar". plot\_error\_bars

Whether to plot the error bars. Default is TRUE. Error bars are standard error of the mean

plot\_indiv\_vals

Whether to plot the individual values as data points. Default is FALSE.

group\_order The order of the groups along the x-axis. 'Options include:

• none: No ordering is performed. Default.

- smart: Groups are ordered based on the sample names.
- arranged: Groups are ordered based on one or more factor column(s) in mf\_data. Factor column names are passed to the function using the group\_order\_input.
- custom: Groups are ordered based on a custom vector of group names. The custom vector is passed to the function using the group\_order\_input.

group\_order\_input

The order of the groups if group\_order is "custom". The column name by which to arrange groups if group\_order is "arranged".

add\_labels The data labels to display on the plot. Either "indiv\_count", "indiv\_MF", "mean\_count",

"mean\_MF", or "none". Count labels display the number of mutations, MF labels display the mutation frequency. Mean plots the mean value. Indiv plots the labels for individual data points (only if plot\_indiv\_vals = TRUE). Default

is "none".

scale\_y\_axis The scale of the y axis. Either "linear" or "log". Default is "linear".

plot\_title The title of the plot. Default is "Mean Mutation Frequency".

custom\_palette A custom color palette to use for the plot. Values that can be customized in-

clude "Mean", "Individual", "Mean min", "Mean max", "Individual min", and

"Individual max". Default is NULL. @return a ggplot object

xlab The x-axis label. Default is the value of group\_col.

ylab The y-axis label. Default is "Mutation Frequency (mutations/bp)".

plot\_mf

plot\_mf

Plot the Mutation Frequency

# Description

This function creates a plot of the mutation frequency.

## Usage

```
plot_mf(
    mf_data,
    group_col,
    plot_type = "bar",
    mf_type = "min",
    fill_col = NULL,
    custom_palette = NULL,
    group_order = "none",
    group_order_input = NULL,
    labels = "count",
    scale_y_axis = "linear",
    x_lab = NULL,
    y_lab = NULL,
    title = NULL
)
```

# Arguments

mf_data	A data frame containing the mutation frequency data. This is obtained from the calculate_mut_freq function with SUMMARY = TRUE.
group_col	The name of the column containing the sample/group names for the x-axis.
plot_type	The type of plot to create. Options are "bar" or "point".
mf_type	The type of mutation frequency to plot. Options are "min", "max", "both", or "stacked". If "both", the min and max mutation frequencies are plotted side by side. "stacked" can be chosen for bar plot_type only. If "stacked", the difference between the min and max MF is stacked on top of the min MF such that the total height of both bars represent the max MF.
fill_col	The name of the column containing the fill variable.
custom_palette	A character vector of colour codes to use for the plot. If NULL, a default palette is used
group_order	The order of the samples/groups along the x-axis. ' Options include:

- none: No ordering is performed. Default.
- smart: Samples are ordered based on the sample names.
- arranged: Samples are ordered based on one or more factor column(s) in mf\_data. Factor column names are passed to the function using the group\_order\_input.

plot\_model\_mf 35

• custom: Samples are ordered based on a custom vector of sample names. The custom vector is passed to the function using the group\_order\_input.

```
group_order_input
```

The order of the samples/groups if group\_order is "custom". The column name by which to arrange samples/groups if group\_order is "arranged"

labels

The data labels to display on the plot. Either "count", "MF", or "none". Count labels display the number of mutations, MF labels display the mutation frequency.

scale\_y\_axis

The scale of the y axis. Either "linear" or "log".

x\_lab The label for the x axis.

y\_lab The label for the y axis.

title The title of the plot.

#### Value

A ggplot object

plot\_model\_mf

Plot your mf model

# Description

Provide a visualization of the point estimates derived using model\_mf()

```
plot_model_mf(
  model,
  plot_type = "point",
  x_effect = NULL,
  plot_error_bars = TRUE,
  plot_signif = TRUE,
  ref_effect = NULL,
  x_order = NULL,
  fill_order = NULL,
  x_label = NULL,
  y_label = NULL,
  plot_title = NULL,
  fill_label = NULL,
  custom_palette = NULL
)
```

36 plot\_spectra

#### **Arguments**

model A model object created using model\_mf()

plot\_type The type of plot to create. Options are "bar" or "point".

x\_effect If there are multiple fixed effects in the model, specify the fixed effect to plot on

the x-axis. The other will be used in the fill aesthetic. Currently, only 2 fixed

effects are supported.

plot\_error\_bars

Logical. If TRUE, the estimated standard error will be added to the plot.

plot\_signif

Logical. If TRUE, will add significance labels based on the pairwise\_comparisons data frame in the model object. This is only valid if you supplied a contrasts table to model\_mf(). Symbols will be applied to plotted values that are significantly different from the reference. Your contrasts table is structured as a data frame with two columns, each containing levels of the fixed effects to be contrasted. When adding significance labels, symbols will be added to the values defined in the first column, while the second column will represent the reference. A different symbol will be used for each unique reference level. If a single plotted value has been contrasted against multiple references, then it will gain multiple symbols for each significance difference.

x\_order A character vector indicating the order of the levels for the x\_effect.

fill\_order A character vector indicating the order of the levels for the fill aesthetic, if ap-

plicable.

x\_label The label for the x-axis.

y\_label The label for the y-axis.

plot\_title The title of the plot.

fill\_label The label for the fill aesthetic, if applicable.

custom\_palette A vector of colors to use for the fill and color aesthetics. If not provided, a

default palette will be used. When plotting a model that has a single fixed effect, you can specify colors for "fill" and "color" using a named vector. Likewise, when plotting a model with two fixed effects, you can specify colors for the

levels within your fill variable.

#### Value

A ggplot object.

## Description

Given a data frame construct a plot displaying the mutation subtypes observed in a cohort.

plot\_spectra 37

## Usage

```
plot_spectra(
   mutation_data,
   group_col = "sample",
   subtype_resolution = "base_6",
   response = c("frequency", "proportion", "sum"),
   mf_type = c("min", "max"),
   variant_types = c("snv", "deletion", "insertion", "complex", "mnv", "symbolic"),
   group_order = "none",
   group_order_input = NULL,
   dist = "cosine",
   cluster_method = "ward.D",
   palette = NULL,
   x_lab = NULL,
   y_lab = NULL
)
```

## **Arguments**

mutation\_data

A data frame containing the mutation data. This data must include a column containing the mutation subtypes, a column containing the sample/cohort names, and a column containing the response variable. Typical response variables can be the subtype frequency, proportion, or count.

group\_col The name of the column in data that contains the sample/cohort names. subtype\_resolution

The resolution of the mutation spectra. Default is base\_6.

response

The name of the column in data that contains the response variable. Typical response variables can be the subtype frequency, proportion, or count.

mf\_type

The type of mutation frequency to use. Default is min.

variant\_types

A character vector of the mutation types to include.

group\_order

The method for ordering the samples within the plot. Options include:

- none: No ordering is performed. Default.
- smart: Groups are ordered based on the group names.
- arranged: Groups are ordered based on one or more factor column(s) in mf\_data. Column names are passed to the function using the group\_order\_input.
- custom: Groups are ordered based on a custom vector of group names. The custom vector is passed to the function using the group\_order\_input.
- clustered: Groups are ordered based on hierarchical clustering. The dissimilarity matrix can be specified using the dist argument. The agglomeration method can be specified using the cluster\_method argument.

group\_order\_input

A character vector specifying details for the group order method. If group\_order is arranged, group\_order\_input should contain the column name(s) to be used for ordering the samples. If group\_order is custom, group\_order\_input should contain the custom vector of group names.

38 plot\_trinucleotide

dist	The dissimilarity matrix for hierarchical clustering. Options are cosine, euclidean, maximum, manhattan, canberra, binary or minkowski. The default is cosine. See dist for details.
cluster_method	The agglomeration method for hierarchical clustering. Options are ward.D, ward.D2, single, complete, average (= $UPGMA$ ), mcquitty (= $WPGMA$ ), median (= $WPGMC$ ) or centroid (= $UPGMC$ ). The default is Ward.D. See hclust for details.
palette	A named vector of colors to be used for the mutation subtypes. The names of the vector should correspond to the mutation subtypes in the data. The default is a set of colors from the RColorBrewer package.
x_lab	The label for the x-axis. Default is the value of group_col.
y_lab	The label for the y-axis. Default is the value of response_col.

plot\_trinucleotide

Plot the trinucleotide spectrum

#### **Description**

Creates barplots of the trinucleotide spectrum for all levels of a given group based on the mutation data. All plots are exported.

#### Usage

```
plot_trinucleotide(
   mutation_data,
   response = c("frequency", "proportion", "sum"),
   mf_type = "min",
   group_col = "dose",
   max_y = c("individual", "group"),
   sum_totals = TRUE,
   output_path = NULL,
   output_type = "svg"
)
```

#### **Arguments**

mutation\_data A data frame containing mutation data. This can be obtained using the 'im-

port\_mut\_data' or 'read\_vcf' functions.

response A character string specifying the type of response to plot. Must be one of 'fre-

quency', 'proportion', or 'sum'.

mf\_type A character string specifying the mutation count method to plot. Must be one of

'min' or 'max'. Default is 'min'.

group\_col A character string specifying the column(s) in 'mutation\_data' to group the data

by. Default is 'sample'. The sum, proportion, or frequency will be calculated and a plot will be generated for all unique levels of this group. You can specify

more than one column to group by.

max_y	A character string specifying the max response value for the y-axis. Must be one of 'individual' or 'group'.'individual' will adjust the maximum y-axis value for each level of the group independently of the others. 'group' will set the maximum y-axis value based on the entire dataset such that all plots will have the same scale. Default is 'group'.
sum_totals	A logical value specifying whether to sum the total mutations.
output_path	A character string specifying the path to save the output plot. Default is NULL. This will create an output directory in the current working directory.
output_type	A character string specifying the type of output file. Options are 'jpeg', 'pdf', 'png', 'svg', or 'tiff'. Default is 'svg'.

#### **Details**

The function calculates the mutation frequency and plots the trinucleotide spectrum for all levels of a given group based on the mutation data. The function calculates the mutation frequency using the 'calculate\_mut\_freq' function with "cols\_to\_group" set to 'group\_col' and "subtype\_resolution" set to 'base\_96'. For a given group, mutation counts and total informative duplex bases are summed across all samples. Mutation frequency is calculated by dividing the total mutation counts by the total number of duplex bases. For a given mutation subtype, proportion is calculated as the proportion of total mutation counts normalized to the total number of duplex bases for a given group and subtype. ^ should just explain this in calculate mutation freq and refer to that function.

```
plot_trinucleotide_heatmap

Create a heatmap plot
```

# Description

This function creates a heatmap plot using the provided data file.

```
plot_trinucleotide_heatmap(
    mf_data,
    group_var = "dose",
    mf_type = "min",
    mut_proportion_scale = "turbo",
    max = 0.2,
    rescale_data = FALSE,
    condensed = FALSE
)
```

40 print\_ascii\_art

## **Arguments**

mf\_data The data file

group\_var The variable to group by.

mf\_type The type of mutation frequency to plot. Options are "min" or "max". (Default:

"min")

mut\_proportion\_scale

The scale option for the mutation proportion. Options are passed to viridis::scale\_fill\_viridis\_c.

One of # inferno, magma, plasma, viridis, cividis, turbo, mako, or rocket. We highly recomend the default for its ability to disciminate hard to see patterns.

(Default: "turbo")

max Maximum value used for plotting the relative contributions. Contributions that

are higher will have the maximum colour. (Default: 0.2)

rescale\_data Logical value indicating whether to rescale the mutation proportions to increase

the dynamic range of colors shown on the plot. (Default: TRUE)

condensed More condensed plotting format. Default = F.

#### Value

A ggplot object representing the heatmap plot.

# **Examples**

```
create_heatmap(mutation_data, dose, "inferno")
```

print\_ascii\_art

This function prints ASCII art when the package is loaded

# **Description**

This function prints ASCII art when the package is loaded

```
print_ascii_art()
```

radar\_plot 41

rauar prot Create a radar prot	radar_plot	Create a radar plot
--------------------------------	------------	---------------------

# **Description**

Create a radar plot

# Usage

```
radar_plot(mf_data, response_col, label_col, facet_col)
```

# **Arguments**

mf\_data A data frame with the data to plot response\_col The column with the response values

label\_col The column with the labels for the radar plot.

facet\_col The column with the group to facet the radar plots.

#### Value

A radar plot

rename_columns	Map column names of mutation data to default column names. A utility
	function that renames columns of mutation data to default columns

names.

#### **Description**

Map column names of mutation data to default column names. A utility function that renames columns of mutation data to default columns names.

# Usage

```
rename_columns(data, column_map = op$column)
```

# Arguments

data mutation data

column\_map a list that maps synonymous column names to their default.

# Value

the mutation data with column names changed to match default.

42 reverseComplement

render\_report

Read configuration file and render R Markdown document

#### **Description**

This function reads a configuration file in YAML format, extracts the parameters, and renders an R Markdown document using the specified parameters.

#### Usage

```
render_report(config_filepath, output_file)
```

# Arguments

#### Value

None

#### **Examples**

```
read_config_and_render("config.yaml", "output.html")
```

reverseComplement

Get the reverse complement of a DNA or RNA sequence.

# Description

Get the reverse complement of a DNA or RNA sequence.

#### Usage

```
reverseComplement(
    x,
    content = c("dna", "rna"),
    case = c("lower", "upper", "as is")
)
```

#### **Arguments**

```
x A character vector of DNA or RNA sequences.

content c("dna", "rna") The type of sequence to be reversed.

case c("lower", "upper", "as is") The case of the output sequence.
```

sidak 43

#### **Details**

This file is part of the source code for SPGS: an R package for identifying statistical patterns in genomic sequences. Copyright (C) 2015 Universidad de Chile and INRIA-Chile A copy of Version 2 of the GNU Public License is available in the share/licenses/gpl-2 file in the R installation directory or from http://www.R-project.org/Licenses/GPL-2. reverseComplement.R

sidak

Correct p-values for multiple comparisons

## **Description**

Correct p-values for multiple comparisons

#### Usage

sidak(vecP)

#### **Arguments**

vecP

vector of p-values

## Details

This function corrects a vector of probabilities for multiple testing using the Bonferroni (1935) and Sidak (1967) corrections. References: Bonferroni (1935), Sidak (1967), Wright (1992). Bonferroni, C. E. 1935. Il calcolo delle assicurazioni su gruppi di teste. Pp. 13-60 in: Studi in onore del Professore Salvatore Ortu Carboni. Roma. Sidak, Z. 1967. Rectangular confidence regions for the means of multivariate normal distributions. Journal of the American Statistical Association 62:626-633. Wright, S. P. 1992. Adjusted P-values for simultaneous inference. Biometrics 48: 1005-1013. Pierre Legendre, May 2007

#### Value

adjusted p-values

signature\_analysis\_sigminer

Run COSMIC signatures comparison

## **Description**

After cleaning the mutation data input, runs several Alexandrov Lab tools for COSMIC signature analysis (assigns signatures to best explain the input data).

44 signature\_fitting

#### Usage

```
signature_analysis_sigminer(
  mutations,
  project_name = "Default",
  project_genome = "BSgenome.Mmusculus.UCSC.hg38",
  group = "sample",
  run_bootstrapping = F,
  vaf_cutoff,
  ...
)
```

#### **Arguments**

mutations A data frame, imported from a .mut or .vcf file

project\_name The name of the project; used to get mutation data into the required .txt format

for SigProfiler

project\_genome A string describing the reference genome to use; e.g., GRCh38

group The column in the mutation data used to aggregate groups (e.g., sample ID,

tissue, dose)

run\_bootstrapping

TRUE or FALSE. Default FALSE. Determines if the sig\_fit\_bootstrap\_batch() function should be run. This *should* be done, but the process is slow, so it's best

to confirm that the rest of the analysis is working as expected first.

vaf\_cutoff The threshold above which variants are identified as ostensibly germline vari-

ants using a cutoff for variant allele fraction (VAF). There is no default value provided, but generally a value of 0.1 (i.e., 10%) is a good starting point. Setting this will remove variants that are present at a frequency greater than this

value at a given site.

... additional arguments may be supplied To Do: we need to document the elipsis

... in the parameters of the function in a way that doesn't cause a warning during

Check()

#### Value

Creates a subfolder in the output directory with SigProfiler tools results. Suggests: sigminer

signature\_fitting

Run COSMIC signatures comparison

## Description

After cleaning the mutation data input, runs several Alexandrov Lab tools for COSMIC signature analysis (assigns signatures to best explain the input data).

signature\_fitting 45

# Usage

```
signature_fitting(
  mutation_data,
  project_name = "Default",
  project_genome = "GRCh38",
  env_name = "MutSeqR",
  group = "sample",
  output_path = NULL,
  python_version = "3.11",
  python_path = "~/../../AppData/Local/Programs/Python/Python310/python.exe",
  python_home =
  "C:/Users/adodge/OneDrive - HC-SC PHAC-ASPC/Documents/.virtualenvs/r-reticulate")
```

# **Arguments**

mutation_data	A data frame, imported from a .mut file
project_name	The name of the project; used to get mutation data into the required .txt format for SigProfiler
<pre>project_genome</pre>	A string describing the reference genome to use; e.g. GRCh37, GRCH38, mm10, mm9, rn6 output_path The directory where output results should be written. *not a parameter of the function
env_name	The name of the virtual environment. This will be created on first use.
group	The column in the mutation data used to aggregate groups (e.g., sample ID, tissue, dose)
output_path	The filepath to the directory in which the output folder will be created to store results.
python_version	The version of python to be used.
python_path	The path to the version of python to be used with reticulate. It is important that this version of python meets the dependencies, including the SigProfiler python tools.
python_home	The path to the conda virtual environment that contains the required python dependencies

# Value

 $Creates\ a\ subfolder\ in\ the\ output\ directory\ with\ SigProfiler\ tools\ results.\ SigProfilerMatrixGeneratorR\ SigProfilerMatrixGeneratorR\ install$ 

46 spectra\_comparison

spectra\_comparison

Compare the overall mutation spectra between groups

#### **Description**

spectra\_comparison compares the mutation spectra of groups using a modified contingency table approach.

#### Usage

```
spectra_comparison(
 mutation_data,
 cols_to_group,
  subtype_resolution = "base_6",
 variant_types = c("snv", "deletion", "insertion", "complex", "mnv", "symbolic"),
 mf_type = "min",
 contrasts,
  cont_sep = "\t"
)
```

## **Arguments**

mutation\_data

A data frame containing the mutation data. This is the output from import\_mut\_data or import\_vcf\_data.

cols\_to\_group

A character vector of the column names in the mutation data that you want to group by. Ex. c("dose", "tissue"). This function will sum the mutations across groups before running the comparison.

subtype\_resolution

The resolution of the mutation spectra to be compared. Options include "base\_6", "base\_12", "base\_96", and "base\_192" and "type". See calculate\_mut\_freq for more details.

variant\_types

A character vector of the mutation types to include in the comparison. Default is all types of mutations.

mf\_type

The type of mutation frequency to use. Default is "min".

contrasts

a filepath to a tab-delimited . txt file OR a dataframe that will specify the comparisons to be made between groups. The table must consist of two columns. The first column will be a level within your group column and the second column must be the group level that it will be compared to. All values must correspond to entries in your cols\_to\_group column. For more than one group variable, separate the levels of each group with a colon. Ensure that all groups listed in cols\_to\_group are represented in each entry for the table. See details for examples.

cont\_sep

The delimiter used to import the contrasts table. Default is tab.

subtype\_dict 47

# **Details**

Examples of contrasts: If you have group = "dose" with dose groups 0, 25, 50, 100. The first column would contain the treated groups (25, 50, 100), while the second column would be 0, thus comparing each treated group to the control group.

250

500

1000

Ex. Consider two grouping variables group = c("dose", "tissue"); with levels dose (0, 25, 50, 100) and tissue("bone\_marrow", "liver"). To compare the mutation spectra between tissues for each dose group, the contrast table would look like:

0:bone\_marrow 0:liver

25:bone\_marrow 25:liver

50:bone\_marrow 50:liver

100:bone\_marrow 100:liver

#### Value

the log-likelihood statistic G2 for the specified comparisons with the p-value adjusted for multiple-comparisons.

subtype\_dict

Values accepted for mutation subtypes

# Description

These values are used to enable user input to translate to columns in a mut file

#### Usage

subtype\_dict

#### **Format**

A vector with corresponding values

write\_excel\_from\_list

subtype\_list

A list of mutation subtypes at different resolutions

# **Description**

A list of mutation subtypes at different resolutions

# Usage

```
subtype\_list
```

#### **Format**

A list with corresponding values

```
write_excel_from_list Write Excel tables
```

# Description

Takes a list of tables (data frames) and writes each one to a separate Excel sheet in a workbook. Names of tabs will be based on names in the list.

# Usage

```
write_excel_from_list(list_of_tables, output_path, workbook_name)
```

# Arguments

list\_of\_tables A named list of data frames to be written.

output\_path The directory where the Excel file should be written.

workbook\_name The file name for the Excel file.

#### Value

A saved Excel workbook.

## **Description**

Takes a single data frame and writes it to an Excel workbook.

#### Usage

```
write_excel_single_table(
  mut_data,
  output_path = "./",
  workbook_name = "Default"
)
```

#### **Arguments**

mut\_data The data frame to be written.

output\_path The directory where the Excel file should be written.

workbook\_name The file name for the Excel file.

#### Value

A saved Excel workbook.

```
write_mutational_matrix
```

Write a Mutational Matrix to input into the sigprofiler web application

# Description

Creates a .txt file from mutation data that can be used for mutational signatures analysis using the SigProfiler web application. Can handle group analyses (ex dose, tissue, etc). Currently only supports snv.

```
write_mutational_matrix(
  mutation_data,
  group = "dose",
  matrices = "base_96",
  mf_type = "min",
  filter = "somatic",
  output_path = NULL
)
```

#### **Arguments**

or read\_vcf().

group The column in the mutation data used to aggregate groups (e.g., sample, tissue,

dose)

matrices At what resolution should the sny mutation counts be calculated? Options are

"base\_96", 1536?, 384?, 6144?, DINUC? 6, 24, INDEL TO DO: add more ma-

trices options.

mf\_type Mutation counting method. Options are "min" or "max". Minimum Independent

Frequency (min): All identical mutations within a sample are assumed to be the result of clonal expansion and are thus only counted once. Maximum Independent Frequency (max): All identical mutations within a sample are assumed to be idenpendant mutational evens and are included in the mutation frequency calculation. Note that this does not apply for germline variants. Default is "min".

TO DO: clonality cut\_off?

filter Parameter allows you to choose to filter for only somatic or germline mutations.

output\_path The path to save the output file. Default is NULL, which will save the file in the

current working directory. Values = c("somatic", "germline", "none). "none"

will leave the data unfiltered.

#### Value

a .txt file that can be uploaded to the sigprofiler web application as "Mutational Matrix"

```
write_mutation_calling_file
```

Write the mutation calling file to input into the sigprofiler web application

# Description

Creates a .txt file from mutation data that can be used for mutational signatures analysis using the SigProfiler web application.Cannot group higher than sample.

```
write_mutation_calling_file(
  mutation_data,
  project_name = "test",
  project_genome = "GRCm38",
  output_path = NULL
)
```

write\_reference\_fasta 51

# **Arguments**

mutation_data	The GRanges object containing the mutation data. The output of import_mut_data() or read_vcf().
project_name	The name of the project; used to get mutation data into the required .txt format for SigProfiler
<pre>project_genome</pre>	A string describing the reference genome to use (e.g., GRCh38, GRCm38)
output_path	The path to save the output file. Default is NULL. See import_mut_data() or read_vcf() for more details.

# Value

a .txt file that can be uploaded to the sigprofiler web application as "Mutational calling file" Filters out ostensibly germline mutations identified in mutation data.

write\_reference\_fasta Write FASTA file of reference sequences

# Description

In some cases, you might want to generate an arbitrary multi-sequence FASTA file from GRanges including the reference sequences. This function allows you to do this.

# Usage

```
write_reference_fasta(ref_ranges, fasta_out = "./reference_output.fasta")
```

# Arguments

ref\_ranges A GRanges object including the sequences of the reference regions included for

the data.

fasta\_out The path for the FASTA file output.

## Value

Writes a FASTA reference file to be used in downstream applications.

write\_VCF\_from\_mut

 $write\_VCF\_from\_mut$ 

Write FASTA file of reference sequences

# Description

In some cases, you might want to generate an arbitrary multi-sequence FASTA file from GRanges including the reference sequences. This function allows you to do this.

# Usage

```
write_VCF_from_mut(mutation_data, vcf_out = "./mutation_output.vcf")
```

# Arguments

 $\mbox{mutation\_data} \quad A \ GRanges \ object \ imported \ from \ a \ .mut \ file$ 

vcf\_out The path for the VCF file output.

# Value

Writes a VCF file of mutations to be used in downstream applications.

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