

# Package ‘sigminer’

August 9, 2022

**Title** Extract, Analyze and Visualize Mutational Signatures for Genomic Variations

**Version** 2.1.6

**Description** Genomic alterations including single nucleotide substitution, copy number alteration, etc. are the major force for cancer initialization and development. Due to the specificity of molecular lesions caused by genomic alterations, we can generate characteristic alteration spectra, called 'signature' (Wang, Shixiang, et al. (2021) <[DOI:10.1371/journal.pgen.1009557](https://doi.org/10.1371/journal.pgen.1009557)> & Alexandrov, Ludmil B., et al. (2020) <[DOI:10.1038/s41586-020-1943-3](https://doi.org/10.1038/s41586-020-1943-3)> & Steele Christopher D., et al. (2022) <[DOI:10.1038/s41586-022-04738-6](https://doi.org/10.1038/s41586-022-04738-6)>). This package helps users to extract, analyze and visualize signatures from genomic alteration records, thus providing new insight into cancer study.

**License** MIT + file LICENSE

**URL** <https://github.com/ShixiangWang/sigminer>

**BugReports** <https://github.com/ShixiangWang/sigminer/issues>

**Depends** R (>= 3.5)

**Imports** cli (>= 2.0.0), cowplot, data.table, dplyr, furrr (>= 0.2.0), future, ggplot2 (>= 3.3.0), ggpubr, maftools, magrittr, methods, NMF, purrr, Rcpp, rlang (>= 0.1.2), stats, tidyr

**Suggests** Biobase, Biostrings, BSgenome, BSgenome.Hsapiens.UCSC.hg19, circlize, cluster, copynumber, covr, digest, GenomicRanges, GenSA, ggalluvial, ggcorrplot, ggfittext, ggplotify, ggrepel, IRanges, knitr, lpSolve, markdown, matrixStats, nnls, parallel, patchwork, pheatmap, quadprog, R.utils, RColorBrewer, reticulate, rmarkdown, roxygen2, scales, synchronicity, testthat, tibble, UCSCXenaTools

**LinkingTo** Rcpp

**VignetteBuilder** knitr

**biocViews**

**Encoding** UTF-8

**LazyData** true

**RoxygenNote** 7.2.0

**NeedsCompilation** yes

**Author** Shixiang Wang [aut, cre] (<<https://orcid.org/0000-0001-9855-7357>>),  
 Ziyu Tao [aut] (<<https://orcid.org/0000-0003-3272-1227>>),  
 Huimin Li [aut] (<<https://orcid.org/0000-0003-1683-9057>>),  
 Tao Wu [aut] (<<https://orcid.org/0000-0002-8999-9628>>),  
 Xue-Song Liu [aut, ctb] (<<https://orcid.org/0000-0002-7736-0077>>),  
 Anand Mayakonda [ctb]

**Maintainer** Shixiang Wang <w\_shixiang@163.com>

**Repository** CRAN

**Date/Publication** 2022-08-09 15:20:02 UTC

## R topics documented:

add_h_arrow . . . . .	4
add_labels . . . . .	5
bp . . . . .	7
centromeres.hg19 . . . . .	14
centromeres.hg38 . . . . .	14
centromeres.mm10 . . . . .	15
centromeres.mm9 . . . . .	15
chromsize.hg19 . . . . .	16
chromsize.hg38 . . . . .	16
chromsize.mm10 . . . . .	17
chromsize.mm9 . . . . .	17
CN.features . . . . .	18
CopyNumber-class . . . . .	18
cosine . . . . .	19
cytobands.hg19 . . . . .	19
cytobands.hg38 . . . . .	20
cytobands.mm10 . . . . .	20
cytobands.mm9 . . . . .	21
enrich_component_strand_bias . . . . .	21
get_adj_p . . . . .	22
get_Aneuploidy_score . . . . .	23
get_bayesian_result . . . . .	25
get_cn_freq_table . . . . .	26
get_cn_ploidy . . . . .	26
get_genome_annotation . . . . .	27
get_groups . . . . .	28
get_group_comparison . . . . .	30
get_intersect_size . . . . .	31
get_pLOH_score . . . . .	32
get_shannon_diversity_index . . . . .	33
get_sig_cancer_type_index . . . . .	34

get_sig_db . . . . .	35
get_sig_exposure . . . . .	37
get_sig_feature_association . . . . .	38
get_sig_rec_similarity . . . . .	39
get_sig_similarity . . . . .	40
get_tidy_association . . . . .	42
group_enrichment . . . . .	43
handle_hyper_mutation . . . . .	45
hello . . . . .	46
MAF-class . . . . .	46
output_bootstrap . . . . .	47
output_fit . . . . .	48
output_sig . . . . .	49
output_tally . . . . .	50
read_copynumber . . . . .	50
read_copynumber_ascat . . . . .	52
read_copynumber_seqz . . . . .	53
read_maf . . . . .	53
read_sv_as_rs . . . . .	54
read_vcf . . . . .	55
read_xena_variants . . . . .	56
report_bootstrap_p_value . . . . .	57
same_size_clustering . . . . .	57
scoring . . . . .	58
show_catalogue . . . . .	60
show_cn_circos . . . . .	61
show_cn_components . . . . .	62
show_cn_distribution . . . . .	63
show_cn_features . . . . .	65
show_cn_freq_circos . . . . .	66
show_cn_group_profile . . . . .	67
show_cn_profile . . . . .	69
show_cor . . . . .	70
show_cosmic . . . . .	72
show_cosmic_sig_profile . . . . .	72
show_groups . . . . .	74
show_group_comparison . . . . .	74
show_group_distribution . . . . .	76
show_group_enrichment . . . . .	78
show_group_mapping . . . . .	79
show_sig_bootstrap . . . . .	81
show_sig_consensusmap . . . . .	85
show_sig_exposure . . . . .	86
show_sig_feature_corrplot . . . . .	87
show_sig_fit . . . . .	89
show_sig_profile . . . . .	91
show_sig_profile_heatmap . . . . .	94
show_sig_profile_loop . . . . .	96

sigminer . . . . .	97
sigprofiler . . . . .	98
sig_auto_extract . . . . .	99
sig_convert . . . . .	102
sig_estimate . . . . .	103
sig_extract . . . . .	107
sig_fit . . . . .	109
sig_fit_bootstrap . . . . .	112
sig_fit_bootstrap_batch . . . . .	115
sig_operation . . . . .	116
sig_tally . . . . .	117
sig_unify_extract . . . . .	121
simulated_catalogs . . . . .	122
simulation . . . . .	123
subset.CopyNumber . . . . .	124
transcript.hg19 . . . . .	125
transcript.hg38 . . . . .	125
transcript.mm10 . . . . .	126
transcript.mm9 . . . . .	126
transform_seg_table . . . . .	127
use_color_style . . . . .	128

**Index****129**


---

add_h_arrow	<i>Add Horizontal Arrow with Text Label to a ggplot</i>
-------------	---

---

**Description**

Add Horizontal Arrow with Text Label to a ggplot

**Usage**

```
add_h_arrow(
  p,
  x,
  y,
  label = "optimal number",
  space = 0.01,
  vjust = 0.3,
  seg_len = 0.1,
  arrow_len = unit(2, "mm"),
  arrow_type = c("closed", "open"),
  font_size = 5,
  font_family = c("serif", "sans", "mono"),
  font_face = c("plain", "bold", "italic")
)
```

**Arguments**

p	a ggplot.
x	position at x axis.
y	position at y axis.
label	text label.
space	a small space between arrow and text.
vjust	vertical adjustment, set to 0 to align with the bottom, 0.5 for the middle, and 1 (the default) for the top.
seg_len	length of the arrow segment.
arrow_len	length of the arrow.
arrow_type	type of the arrow.
font_size	font size.
font_family	font family.
font_face	font face.

**Value**

a ggplot object.

---

add_labels	<i>Add Text Labels to a ggplot</i>
------------	------------------------------------

---

**Description**

Add text labels to a ggplot object, such as the result from [show\\_sig\\_profile](#).

**Usage**

```
add_labels(  
  p,  
  x,  
  y,  
  y_end = NULL,  
  n_label = NULL,  
  labels = NULL,  
  revert_order = FALSE,  
  font_size = 5,  
  font_family = "serif",  
  font_face = c("plain", "bold", "italic"),  
  ...  
)
```

**Arguments**

p	a ggplot.
x	position at x axis.
y	position at y axis.
y_end	end position of y axis when n_label is set.
n_label	the number of label, when this is set, the position of labels at y axis is auto-generated according to y and y_end.
labels	text labels or a similarity object from <a href="#">get_sig_similarity</a> .
revert_order	if TRUE, revert label order.
font_size	font size.
font_family	font family.
font_face	font face.
...	other parameters passing to <a href="#">ggplot2::annotate</a> .

**Value**

a ggplot object.

**Examples**

```
# Load mutational signature
load(system.file("extdata", "toy_mutational_signature.RData",
  package = "sigminer", mustWork = TRUE
))
# Show signature profile
p <- show_sig_profile(sig2, mode = "SBS")

# Method 1
p1 <- add_labels(p,
  x = 0.75, y = 0.3, y_end = 0.9, n_label = 3,
  labels = paste0("text", 1:3)
)
p1

# Method 2
p2 <- add_labels(p,
  x = c(0.15, 0.6, 0.75), y = c(0.3, 0.6, 0.9),
  labels = paste0("text", 1:3)
)
p2

# Method 3
sim <- get_sig_similarity(sig2)
p3 <- add_labels(p,
  x = c(0.15, 0.6, 0.75), y = c(0.25, 0.55, 0.8),
  labels = sim, font_size = 2
)
p3
```

## Description

These functions are combined to provide a best practice for optimally identifying mutational signatures and attributing their activities (exposures) in tumor samples. They are listed in order to use.

- `bp_extract_signatures()` for extracting signatures.
- `bp_show_survey()` for showing measures change under different signature numbers to help user select optimal signature number. At default, an aggregated score (named `score`) is generated to suggest the best solution.
- `bp_show_survey2()` for showing simplified signature number survey like `show_sig_number_survey()`.
- `bp_get_sig_obj()` for get a (list of) Signature object which is common used in **sigminer** for analysis and visualization.
- `bp_attribute_activity()` for optimizing signature activities (exposures). NOTE: the activities from extraction step may be better! You can also use `sig_extract` to get optimal NMF result from multiple NMF runs. Besides, you can use `sig_fit` to quantify exposures based on signatures extracted from `bp_extract_signatures()`.
- `bp_extract_signatures_iter()` for extracting signature in a iteration way.
- `bp_cluster_iter_list()` for clustering (hclust with average linkage) iterated signatures to help collapse multiple signatures into one. The result cluster can be visualized by `plot()` or `factoextra::fviz_dend()`.
- `bp_get_clustered_sigs()` for getting clustered (grouped) mean signatures from signature clusters.
- Extra: `bp_get_stats()` for obtaining stats for signatures and samples of a solution. These stats are aggregated (averaged) as the stats for a solution (specific signature number).
- Extra: `bp_get_rank_score()` for obtaining rank score for all signature numbers.

## Usage

```
bp_extract_signatures(  
  nmf_matrix,  
  range = 2:5,  
  n_bootstrap = 20L,  
  n_nmf_run = 50,  
  RTOL = 0.001,  
  min_contribution = 0,  
  cores = min(4L, future::availableCores()),  
  cores_solution = min(cores, length(range)),  
  seed = 123456L,  
  handle_hyper_mutation = TRUE,
```

```

    report_integer_exposure = FALSE,
    only_core_stats = nrow(nmf_matrix) > 100,
    cache_dir = file.path(tempdir(), "sigminer_bp"),
    keep_cache = FALSE,
    pynmf = FALSE,
    use_conda = TRUE,
    py_path = "/Users/wsx/anaconda3/bin/python"
)

bp_extract_signatures_iter(
  nmf_matrix,
  range = 2:5,
  sim_threshold = 0.95,
  max_iter = 10L,
  n_bootstrap = 20L,
  n_nmf_run = 50,
  RTOL = 0.001,
  min_contribution = 0,
  cores = min(4L, future::availableCores()),
  cores_solution = min(cores, length(range)),
  seed = 123456L,
  handle_hyper_mutation = TRUE,
  report_integer_exposure = FALSE,
  only_core_stats = nrow(nmf_matrix) > 100,
  cache_dir = file.path(tempdir(), "sigminer_bp"),
  keep_cache = FALSE,
  pynmf = FALSE,
  use_conda = FALSE,
  py_path = "/Users/wsx/anaconda3/bin/python"
)

bp_cluster_iter_list(x, k = NULL, include_final_iteration = TRUE)

bp_get_clustered_sigs(SigClusters, cluster_label)

bp_get_sig_obj(obj, signum = NULL)

bp_get_stats(obj)

bp_get_rank_score(obj)

bp_show_survey2(
  obj,
  x = "signature_number",
  left_y = "silhouette",
  right_y = "L2_error",
  left_name = left_y,
  right_name = right_y,

```



```

    left_color = "black",
    right_color = "red",
    left_shape = 16,
    right_shape = 18,
    shape_size = 4,
    highlight = NULL
)

bp_show_survey(
  obj,
  add_score = FALSE,
  scales = c("free_y", "free"),
  fixed_ratio = TRUE
)

bp_attribute_activity(
  input,
  sample_class = NULL,
  nmf_matrix = NULL,
  method = c("bt", "stepwise"),
  bt_use_prop = FALSE,
  return_class = c("matrix", "data.table"),
  use_parallel = FALSE,
  cache_dir = file.path(tempdir(), "sigminer_attribute_activity"),
  keep_cache = FALSE
)

```

## Arguments

<code>nmf_matrix</code>	a matrix used for NMF decomposition with rows indicate samples and columns indicate components.
<code>range</code>	a numeric vector containing the ranks of factorization to try. Note that duplicates are removed and values are sorted in increasing order. The results are notably returned in this order.
<code>n_bootstrap</code>	number of bootstrapped (resampling) catalogs used. When it is 0, the original (input) mutation catalog is used for NMF decomposition, this is not recommended, just for testing, user should not set it to 0.
<code>n_nmf_run</code>	number of NMF runs for each bootstrapped or original catalog. At default, in total <code>n_bootstrap</code> x <code>n_nmf_run</code> (i.e. 1000) NMF runs are used for the task.
<code>RTOL</code>	a threshold proposed by Nature Cancer paper to control how to filter solutions of NMF. Default is 0.1% (from reference #2), only NMF solutions with KLD (KL deviance) $\leq 100.1\%$ minimal KLD are kept.
<code>min_contribution</code>	a component contribution threshold to filter out small contributed components.
<code>cores</code>	number of cpu cores to run NMF.
<code>cores_solution</code>	cores for processing solutions, default is equal to argument <code>cores</code> .
<code>seed</code>	a random seed to make reproducible result.

handle_hyper_mutation	default is TRUE, handle hyper-mutant samples.
report_integer_exposure	if TRUE, report integer signature exposure by bootstrapping technique.
only_core_stats	if TRUE, only calculate the core stats for signatures and samples.
cache_dir	a directory for keep temp result files.
keep_cache	if TRUE, keep cache results.
pymnf	if TRUE, use Python NMF driver <a href="#">Nimfa</a> . The seed currently is not used by this implementation, so the only way to reproduce your result is setting keep_cache = TRUE.
use_conda	if TRUE, create an independent conda environment to run NMF.
py_path	path to Python executable file, e.g. <code>'/Users/wsx/anaconda3/bin/python'</code> . In my test, it is more stable than use_conda=TRUE. You can install the Nimfa package by yourself or set use_conda to TRUE to install required Python environment, and then set this option.
sim_threshold	a similarity threshold for selecting samples to auto-rerun the extraction procedure (i.e. <code>bp_extract_signatures()</code> ), default is 0.95.
max_iter	the maximum iteration size, default is 10, i.e., at most run the extraction procedure 10 times.
x	result from <code>bp_extract_signatures_iter()</code> or a list of Signature objects.
k	an integer sequence specifying the cluster number to get silhouette.
include_final_iteration	if FALSE, exclude final iteration result from clustering for input from <code>bp_extract_signatures_iter()</code> , not applied if input is a list of Signature objects.
SigClusters	result from <code>bp_cluster_iter_list()</code> .
cluster_label	cluster labels for a specified cluster number, obtain it from <code>SigClusters\$sil_df</code> .
obj	a <code>ExtractionResult</code> object from <code>bp_extract_signatures()</code> .
signum	a integer vector to extract the corresponding Signature object(s). If it is NULL (default), all will be returned.
left_y	column name for left y axis.
right_y	column name for right y axis.
left_name	label name for left y axis.
right_name	label name for right y axis.
left_color	color for left axis.
right_color	color for right axis.
left_shape, right_shape, shape_size	shape setting.
highlight	a integer to highlight a x.
add_score	if FALSE, don't show score and label optimal points by rank score.
scales	one of "free_y" (default) and "free" to control the scales of plot facet.

fixed_ratio	if TRUE (default), make the x/y axis ratio fixed.
input	result from <code>bp_extract_signatures()</code> or a Signature object.
sample_class	a named string vector whose names are sample names and values are class labels (i.e. cancer subtype). If it is NULL (the default), treat all samples as one group.
method	one of 'bt' (use bootstrap exposure median, from reference #2, <b>the most recommended way in my personal view</b> ) or 'stepwise' (stepwise reduce and update signatures then do signature fitting with last signature sets, from reference #2, the result tends to assign the contribution of removed signatures to the remaining signatures, <b>maybe I misunderstand the paper method? PAY ATTENTION</b> ).
bt_use_prop	this parameter is only used for bt method to reset low contributing signature activity (relative activity $< 0.01$ ). If TRUE, use empirical P value calculation way (i.e. proportion, used by reference #2), otherwise a t. test is applied.
return_class	string, 'matrix' or 'data.table'.
use_parallel	if TRUE, use parallel computation based on <b>furrr</b> package. It can also be an integer for specifying cores.

### Details

The signature extraction approach is adopted from reference #1, #2, and the whole best practice is adopted from the pipeline used by reference #3. I implement the whole procedure with R code based on the method description of papers. The code is well organized, tested and documented so user will find it pretty simple and useful. Besides, the structure of the results is very clear to see and also visualize like other approaches provided by **sigminer**.

### Value

It depends on the called function.

### Measure Explanation in Survey Plot

The survey plot provides a pretty good way to facilitate the signature number selection. A score measure is calculated as the weighted mean of selected measures and visualized as the first sub-plot. The optimal number is highlighted with red color dot and the best values for each measures are also highlighted with orange color dots. The detail of 6 measures shown in plot are explained as below.

- score - an aggregated score based on rank scores from selected measures below. The higher, the better. When two signature numbers have the same score, the larger signature number is preferred (this is a rare situation, you have to double check other measures).
- silhouette - the average silhouette width for signatures, also named as ASW in reference #2. The signature number with silhouette decreases sharply is preferred.
- distance - the average sample reconstructed cosine distance, the lower value is better.
- error - the average sample reconstructed error calculated with L2 formula (i.e. L2 error). This lower value is better. This measure represents a similar concept like distance above, they are all used to quantify how well sample mutation profiles can be reconstructed from signatures, but distance cares the whole mutation profile similarity while error here cares value difference.

- `pos_cor` - the average positive signature exposure correlation coefficient. The lower value is better. This measure is constructed based on my understanding about signatures: mutational signatures are typically treated as independent recurrent patterns, so their activities are less correlated.
- `similarity` - the average similarity within in a signature cluster. Like `silhouette`, the point decreases sharply is preferred. In the practice, results from multiple NMF runs are clustered with "clustering with match" algorithm proposed by reference #2. This value indicates if the signature profiles extracted from different NMF runs are similar.

### Author(s)

Shixiang Wang [w\\_shixiang@163.com](mailto:w_shixiang@163.com)

### References

Alexandrov, Ludmil B., et al. "Deciphering signatures of mutational processes operative in human cancer." *Cell reports* 3.1 (2013): 246-259.

Degasperi, Andrea, et al. "A practical framework and online tool for mutational signature analyses show intertissue variation and driver dependencies." *Nature cancer* 1.2 (2020): 249-263.

Alexandrov, Ludmil B., et al. "The repertoire of mutational signatures in human cancer." *Nature* 578.7793 (2020): 94-101.

### See Also

See [sig\\_estimate](#), [sig\\_extract](#), [sig\\_auto\\_extract](#), [sigprofiler\\_extract](#) for other approaches.

### Examples

```
data("simulated_catalogs")

# Here I reduce the values for n_bootstrap and n_nmf_run
# for reducing the run time.
# In practice, you should keep default or increase the values
# for better estimation.
#
# The input data here is simulated from 10 mutational signatures

# e1 <- bp_extract_signatures(
#   t(simulated_catalogs$set1),
#   range = 8:12,
#   n_bootstrap = 5,
#   n_nmf_run = 10
# )
#
# To avoid computation in examples,
# Here just load the result
# (e1$signature and e1$exposure set to NA to reduce package size)
load(system.file("extdata", "e1.RData", package = "sigminer"))
```

```

# See the survey for different signature numbers
# The suggested solution is marked as red dot
# with highest integrated score.
p1 <- bp_show_survey(e1)
p1
# You can also exclude plotting and highlighting the score
p2 <- bp_show_survey(e1, add_score = FALSE)
p2

# You can also plot a simplified version
p3 <- bp_show_survey2(e1, highlight = 10)
p3

# Obtain the suggested solution from extraction result
obj_suggested <- bp_get_sig_obj(e1, e1$suggested)
obj_suggested
# If you think the suggested signature number is not right
# Just pick up the solution you want
obj_s8 <- bp_get_sig_obj(e1, 8)

# Track the reconstructed profile similarity
rec_sim <- get_sig_rec_similarity(obj_s8, t(simulated_catalogs$set1))
rec_sim

# After extraction, you can assign the signatures
# to reference COSMIC signatures
# More see ?get_sig_similarity
sim <- get_sig_similarity(obj_suggested)
# Visualize the match result
if (require(pheatmap)) {
  pheatmap::pheatmap(sim$similarity)
}

# You already got the activities of signatures
# in obj_suggested, however, you can still
# try to optimize the result.
# NOTE: the optimization step may not truly optimize the result!
expo <- bp_attribute_activity(e1, return_class = "data.table")
expo$abs_activity

## Not run:
# Iterative extraction:
# This procedure will rerun extraction step
# for those samples with reconstructed catalog similarity
# lower than a threshold (default is 0.95)
e2 <- bp_extract_signatures_iter(
  t(simulated_catalogs$set1),
  range = 9:11,
  n_bootstrap = 5,
  n_nmf_run = 5,
  sim_threshold = 0.99
)

```

```
e2
# When the procedure run multiple rounds
# you can cluster the signatures from different rounds by
# the following command
# bp_cluster_iter_list(e2)

## Extra utilities
rank_score <- bp_get_rank_score(e1)
rank_score
stats <- bp_get_stats(e2$iter1)
# Get the mean reconstructed similarity
1 - stats$stats_sample$cosine_distance_mean

## End(Not run)
```

---

centromeres.hg19      *Location of Centromeres at Genome Build hg19*

---

**Description**

Location of Centromeres at Genome Build hg19

**Format**

A data.frame

**Source**

Generate from UCSC gold path

**Examples**

```
data(centromeres.hg19)
```

---

centromeres.hg38      *Location of Centromeres at Genome Build hg38*

---

**Description**

Location of Centromeres at Genome Build hg38

**Format**

A data.frame

**Source**

Generate from Genome Reference Consortium

**Examples**

```
data(centromeres.hg38)
```

---

centromeres.mm10	<i>Location of Centromeres at Genome Build mm10</i>
------------------	---

---

**Description**

Location of Centromeres at Genome Build mm10

**Format**

A data.frame

**Source**

Generate from <https://hgdownload.soe.ucsc.edu/goldenPath/mm10/database/gap.txt.gz>

**Examples**

```
data(centromeres.mm10)
```

---

centromeres.mm9	<i>Location of Centromeres at Genome Build mm9</i>
-----------------	--

---

**Description**

Location of Centromeres at Genome Build mm9

**Format**

A data.frame

**Source**

Generate from <https://hgdownload.soe.ucsc.edu/goldenPath/mm9/database/> with code:

```
for i in $(seq 1 19) X Y;  
do  
wget https://hgdownload.soe.ucsc.edu/goldenPath/mm9/database/chr${i}_gap.txt.gz  
done
```

**Examples**

```
data(centromeres.mm9)
```

---

`chromsize.hg19`*Chromosome Size of Genome Build hg19*

---

**Description**

Chromosome Size of Genome Build hg19

**Format**

A data.frame

**Source**

Generate from UCSC gold path

**Examples**

```
data(chromsize.hg19)
```

---

`chromsize.hg38`*Chromosome Size of Genome Build hg38*

---

**Description**

Chromosome Size of Genome Build hg38

**Format**

A data.frame

**Source**

Generate from UCSC gold path

**Examples**

```
data(chromsize.hg38)
```



---

chromsize.mm10	<i>Chromosome Size of Genome Build mm10</i>
----------------	---

---

**Description**

Chromosome Size of Genome Build mm10

**Format**

A data.frame

**Source**

Generate from UCSC gold path <http://hgdownload.cse.ucsc.edu/goldenPath/mm10/bigZips/mm10.chrom.sizes>

**Examples**

```
data(chromsize.mm10)
```

---

chromsize.mm9	<i>Chromosome Size of Genome Build mm9</i>
---------------	--

---

**Description**

Chromosome Size of Genome Build mm9

**Format**

A data.frame

**Source**

Generate from UCSC gold path <http://hgdownload.cse.ucsc.edu/goldenPath/mm9/bigZips/mm9.chrom.sizes>

**Examples**

```
data(chromsize.mm9)
```

---

CN.features	<i>Classification Table of Copy Number Features Devised by Wang et al. for Method 'W'</i>
-------------	---

---

**Description**

Classification Table of Copy Number Features Devised by Wang et al. for Method 'W'

**Format**

A data.table with "sigminer.features" class name

**Source**

Generate from code under data\_raw/

**Examples**

```
data(CN.features)
```

---

CopyNumber-class	<i>Class CopyNumber</i>
------------------	-------------------------

---

**Description**

S4 class for storing summarized absolute copy number profile.

**Slots**

data data.table of absolute copy number calling.

summary.per.sample data.table of copy number variation summary per sample.

genome\_build genome build version, should be one of 'hg19' or 'hg38'.

genome\_measure Set 'called' will use autosomo called segments size to compute total size for CNA burden calculation, this option is useful for WES and target sequencing. Set 'wg' will autosome size from genome build, this option is useful for WGS, SNP etc..

annotation data.table of annotation for copy number segments.

dropoff.segs data.table of copy number segments dropped from raw input.

---

cosine *Calculate Cosine Measures*

---

**Description**

Calculate Cosine Measures

**Usage**

```
cosine(x, y)
```

**Arguments**

x a numeric vector or matrix with column representing vector to calculate similarity.  
y must be same format as x.

**Value**

a numeric value or matrix.

**Examples**

```
x <- c(1, 1, 1, 0, 0, 0, 0, 0, 0, 0, 0, 0)
y <- c(0, 0, 1, 1, 1, 1, 1, 0, 1, 0, 0, 0)
z1 <- cosine(x, y)
z1
z2 <- cosine(matrix(x), matrix(y))
z2
```

---

cytobands.hg19 *Location of Chromosome Cytobands at Genome Build hg19*

---

**Description**

Location of Chromosome Cytobands at Genome Build hg19

**Format**

A data.frame

**Source**

from UCSC

**Examples**

```
data(cytobands.hg19)
```

---

`cytobands.hg38`*Location of Chromosome Cytobands at Genome Build hg38*

---

**Description**

Location of Chromosome Cytobands at Genome Build hg38

**Format**

A data.frame

**Source**

from UCSC

**Examples**

```
data(cytobands.hg38)
```

---

`cytobands.mm10`*Location of Chromosome Cytobands at Genome Build mm10*

---

**Description**

Location of Chromosome Cytobands at Genome Build mm10

**Format**

A data.frame

**Source**

from UCSC <http://hgdownload.cse.ucsc.edu/goldenpath/mm10/database/cytoBand.txt.gz>

**Examples**

```
data(cytobands.mm10)
```

---

`cytobands.mm9`*Location of Chromosome Cytobands at Genome Build mm9*

---

**Description**

Location of Chromosome Cytobands at Genome Build mm9

**Format**

A data.frame

**Source**

from UCSC <http://hgdownload.cse.ucsc.edu/goldenpath/mm9/database/cytoBand.txt.gz>

**Examples**

```
data(cytobands.mm9)
```

---

`enrich_component_strand_bias`*Performs Strand Bias Enrichment Analysis for a Given Sample-by-Component Matrix*

---

**Description**

See [sig\\_tally](#) for examples.

**Usage**

```
enrich_component_strand_bias(mat)
```

**Arguments**

`mat` a sample-by-component matrix from [sig\\_tally](#) with strand bias labels "T:" and "B:".

**Value**

a data.table sorted by p\_value.

---

 get\_adj\_p

 Get Adjust P Values from Group Comparison
 

---

### Description

Setting `aes(label=..p.adj..)` in `ggpubr::compare_means()` does not show adjust p values. The returned result of this function can be combined with `ggpubr::stat_pvalue_manual()` to fix this problem.

### Usage

```
get_adj_p(
  data,
  .col,
  .grp = "Sample",
  comparisons = NULL,
  method = "wilcox.test",
  p.adjust.method = "fdr",
  p.digits = 3L,
  ...
)
```

### Arguments

<code>data</code>	a data.frame containing column for groups and column for comparison.
<code>.col</code>	column name for comparison.
<code>.grp</code>	column name for groups.
<code>comparisons</code>	Default is NULL, use all combination in group column. It can be a list of length-2 vectors. The entries in the vector are either the names of 2 values on the x-axis or the 2 integers that correspond to the index of the groups of interest, to be compared.
<code>method</code>	a character string indicating which method to be used for comparing means. It can be 't.test', 'wilcox.test' etc..
<code>p.adjust.method</code>	correction method, default is 'fdr'. Run <code>p.adjust.methods</code> to see all available options.
<code>p.digits</code>	how many significant digits are to be used.
<code>...</code>	other arguments passed to <code>ggpubr::compare_means()</code>

### Details

More info see `ggpubr::compare_means()`, `ggpubr::stat_compare_means()` and `stats::p.adjust()`.

### Value

a data.frame containing comparison result

**Source**

<https://github.com/kassambara/ggpubr/issues/143>

**Examples**

```
library(ggpubr)
# T-test
stat.test <- compare_means(
  len ~ dose,
  data = ToothGrowth,
  method = "t.test",
  p.adjust.method = "fdr"
)
stat.test
# Create a simple box plot
p <- ggboxplot(ToothGrowth, x = "dose", y = "len")
p

# Add p values
my_comparisons <- list(c("0.5", "1"), c("1", "2"), c("0.5", "2"))
p + stat_compare_means(method = "t.test", comparisons = my_comparisons)

# Try adding adjust p values
# proposed by author of ggpubr
# however it does not work
p + stat_compare_means(aes(label = ..p.adj..), method = "t.test", comparisons = my_comparisons)

# Solution:
# calculate adjust p values and their location
# then use stat_pvalue_manual() function
p_adj <- get_adj_p(ToothGrowth, .col = "len", .grp = "dose")
p_adj
p + stat_pvalue_manual(p_adj, label = "p.adj")

# Show selected comparisons
# Of note, p value is adjusted
# for three comparisons, but only
# two are showed in figure
p_adj <- get_adj_p(ToothGrowth,
  .col = "len", .grp = "dose",
  comparisons = list(c("0.5", "1"), c("1", "2")))
p + stat_pvalue_manual(p_adj, label = "p.adj")
```

## Description

This implements a Cohen-Sharir method (see reference) like "Aneuploidy Score" computation. You can read the source code to see how it works. Basically, it follows the logic of Cohen-Sharir method but with some difference in detail implementation. Their results should be counterpart, but with no data validation for now. **Please raise an issue if you find problem/bugs in this function.**

## Usage

```
get_Aneuploidy_score(
  data,
  ploidy_df = NULL,
  genome_build = "hg19",
  rm_black_arms = FALSE
)
```

## Arguments

data	a CopyNumber object or a data.frame containing at least 'chromosome', 'start', 'end', 'segVal', 'sample' these columns.
ploidy_df	default is NULL, compute ploidy by segment-size weighted copy number across autosome, see <a href="#">get_cn_ploidy</a> . You can also provide a data.frame with 'sample' and 'ploidy' columns.
genome_build	genome build version, should be 'hg19', 'hg38', 'mm9' or 'mm10'.
rm_black_arms	if TRUE, remove short arms of chr13/14/15/21/22 from calculation as documented in reference #3.

## Value

A data.frame

## References

- Cohen-Sharir, Y., McFarland, J. M., Abdusamad, M., Marquis, C., Bernhard, S. V., Kazachkova, M., ... & Ben-David, U. (2021). Aneuploidy renders cancer cells vulnerable to mitotic checkpoint inhibition. *Nature*, 1-6.
- Logic reference: [https://github.com/quevedor2/aneuploidy\\_score/](https://github.com/quevedor2/aneuploidy_score/).
- Taylor, Alison M., et al. "Genomic and functional approaches to understanding cancer aneuploidy." *Cancer cell* 33.4 (2018): 676-689.

## Examples

```
# Load copy number object
load(system.file("extdata", "toy_copynumber.RData",
  package = "sigminer", mustWork = TRUE
))

df <- get_Aneuploidy_score(cn)
df
```



```
df2 <- get_Aneuploidy_score(cn@data)
df2

df3 <- get_Aneuploidy_score(cn@data,
  ploidy_df = get_cn_ploidy(cn@data)
)
df3
```

---

get\_bayesian\_result    *Get Specified Bayesian NMF Result from Run*

---

### Description

Sometimes, we may want to use or inspect specified run result from [sig\\_auto\\_extract](#). This function is designed for this purpose.

### Usage

```
get_bayesian_result(run_info)
```

### Arguments

run\_info            a data.frame with 1 row and two necessary columns Run and file.

### Value

a list.

### Author(s)

Shixiang Wang

### Examples

```
load(system.file("extdata", "toy_copynumber_tally_W.RData",
  package = "sigminer", mustWork = TRUE
))

res <- sig_auto_extract(cn_tally_W$nmf_matrix, result_prefix = "Test_copynumber", nrun = 1)

# All run info are stored in res$Raw$summary_run
# Obtain result of run 1
res_run1 <- get_bayesian_result(res$Raw$summary_run[1, ])
```

---

get\_cn\_freq\_table      *Get CNV Frequency Table*

---

### Description

Get CNV Frequency Table

### Usage

```
get_cn_freq_table(
  data,
  genome_build = "hg19",
  cutoff = 2L,
  resolution_factor = 1L
)
```

### Arguments

**data**            a CopyNumber object or a data.frame containing at least 'chromosome', 'start', 'end', 'segVal', 'sample' these columns.

**genome\_build**   genome build version, used when data is a data.frame, should be 'hg19' or 'hg38'.

**cutoff**            copy number value cutoff for splitting data into AMP and DEL. The values equal to cutoff are discarded. Default is 2, you can also set a length-2 vector, e.g. c(2, 2).

**resolution\_factor**   an integer to control the resolution. When it is 1 (default), compute frequency in each cytoband. When it is 2, use compute frequency in each half cytoband.

### Value

a data.table.

---

get\_cn\_ploidy            *Get Ploidy from Absolute Copy Number Profile*

---

### Description

Get Ploidy from Absolute Copy Number Profile

### Usage

```
get_cn_ploidy(data)
```

**Arguments**

data            a [CopyNumber](#) object or a data.frame containing at least 'chromosome', 'start', 'end', 'segVal' these columns.

**Value**

a value or a data.table

**Examples**

```
# Load copy number object
load(system.file("extdata", "toy_copynumber.RData",
  package = "sigminer", mustWork = TRUE
))

df <- get_cn_ploidy(cn)
df
```

---

get\_genome\_annotation *Get Genome Annotation*

---

**Description**

Get Genome Annotation

**Usage**

```
get_genome_annotation(
  data_type = c("chr_size", "centro_loc", "cytobands", "transcript"),
  chrs = paste0("chr", c(1:22, "X", "Y")),
  genome_build = c("hg19", "hg38", "mm10", "mm9")
)
```

**Arguments**

data\_type        'chr\_size' for chromosome size, 'centro\_loc' for location of centromeres, 'cytobands' for location of chromosome cytobands and 'transcript' for location of transcripts.

chrs            chromosomes start with 'chr'

genome\_build    one of 'hg19', 'hg38'

**Value**

a data.frame containing annotation data

**Examples**

```
df1 <- get_genome_annotation()
df1

df2 <- get_genome_annotation(genome_build = "hg38")
df2

df3 <- get_genome_annotation(data_type = "centro_loc")
df3

df4 <- get_genome_annotation(data_type = "centro_loc", genome_build = "hg38")
df4

df5 <- get_genome_annotation(data_type = "cytobands")
df5

df6 <- get_genome_annotation(data_type = "cytobands", genome_build = "hg38")
df6
```

---

get\_groups

*Get Sample Groups from Signature Decomposition Information*


---

**Description**

One of key results from signature analysis is to cluster samples into different groups. This function takes Signature object as input and return the membership in each cluster.

**Usage**

```
get_groups(
  Signature,
  method = c("consensus", "k-means", "exposure", "samples"),
  n_cluster = NULL,
  match_consensus = TRUE
)
```

**Arguments**

Signature	a Signature object obtained either from <a href="#">sig_extract</a> or <a href="#">sig_auto_extract</a> . Now it can be used to relative exposure result in <code>data.table</code> format from <a href="#">sig_fit</a> .
method	grouping method, more see details, could be one of the following: <ul style="list-style-type: none"> <li>'consensus' - returns the cluster membership based on the hierarchical clustering of the consensus matrix, it can only be used for the result obtained by <a href="#">sig_extract()</a> with multiple runs using <b>NMF</b> package.</li> <li>'k-means' - returns the clusters by k-means.</li> <li>'exposure' - assigns a sample into a group whose signature exposure is dominant.</li> </ul>

- 'samples' - returns the cluster membership based on the contribution of signature to each sample, it can only be used for the result obtained by `sig_extract()` using NMF package.
- n\_cluster            only used when the method is 'k-means'.
- match\_consensus    only used when the method is 'consensus'. If TRUE, the result will match order as shown in consensus map.

### Details

Users may find there are bigger differences between using method 'samples' and 'exposure' but they use a similar idea to find dominant signature, here goes the reason:

Method 'samples' using data directly from NMF decomposition, this means the two matrix W (basis matrix or signature matrix) and H (coefficient matrix or exposure matrix) are the results of NMF. For method 'exposure', it uses the signature exposure loading matrix. In this situation, each signature represents a number of mutations (alterations) about implementation please see source code of `sig_extract()` function.

### Value

a data.table object

### See Also

`NMF::predict()`, `show_groups`.

### Examples

```
# Load copy number prepare object
load(system.file("extdata", "toy_copynumber_tally_W.RData",
  package = "sigminer", mustWork = TRUE
))
# Extract copy number signatures
library(NMF)
sig <- sig_extract(cn_tally_W$nmf_matrix, 2,
  nrun = 10
)

# Methods 'consensus' and 'samples' are from NMF::predict()
g1 <- get_groups(sig, method = "consensus", match_consensus = TRUE)
g1
g2 <- get_groups(sig, method = "samples")
g2

# Use k-means clustering
g3 <- get_groups(sig, method = "k-means")
g3
```

---

get\_group\_comparison *Get Comparison Result between Signature Groups*

---

## Description

Compare genotypes/phenotypes based on signature groups (samples are assigned to several groups). For categorical type, calculate fisher p value (using `stats::fisher.test`) and count table. In larger than 2 by 2 tables, compute p-values by Monte Carlo simulation. For continuous type, calculate anova p value (using `stats::aov`), summary table and Tukey Honest significant difference (using `stats::TukeyHSD`). The result of this function can be plotted by `show_group_comparison()`.

## Usage

```
get_group_comparison(  
  data,  
  col_group,  
  cols_to_compare,  
  type = "ca",  
  NAs = NA,  
  verbose = FALSE  
)
```

## Arguments

data	a <code>data.frame</code> containing signature groups and genotypes/phenotypes (including categorical and continuous type data) want to analyze. User need to construct this <code>data.frame</code> by him/herself.
col_group	column name of signature groups.
cols_to_compare	column names of genotypes/phenotypes want to summarize based on groups.
type	a character vector with length same as <code>cols_to_compare</code> , 'ca' for categorical type and 'co' for continuous type.
NAs	default is NA, filter NAs for categorical columns. Otherwise a value (either length 1 or length same as <code>cols_to_compare</code> ) fill NAs.
verbose	if TRUE, print extra information.

## Value

a list contains data, summary, p value etc..

## Author(s)

Shixiang Wang [w\\_shixiang@163.com](mailto:w_shixiang@163.com)

**Examples**

```

load(system.file("extdata", "toy_copynumber_signature_by_W.RData",
  package = "sigminer", mustWork = TRUE
))

# Assign samples to clusters
groups <- get_groups(sig, method = "k-means")

set.seed(1234)

groups$prob <- rnorm(10)
groups$new_group <- sample(c("1", "2", "3", "4", NA), size = nrow(groups), replace = TRUE)

# Compare groups (filter NAs for categorical columns)
groups.cmp <- get_group_comparison(groups[, -1],
  col_group = "group",
  cols_to_compare = c("prob", "new_group"),
  type = c("co", "ca"), verbose = TRUE
)

# Compare groups (Set NAs of categorical columns to 'Rest')
groups.cmp2 <- get_group_comparison(groups[, -1],
  col_group = "group",
  cols_to_compare = c("prob", "new_group"),
  type = c("co", "ca"), NAs = "Rest", verbose = TRUE
)

```

---

get\_intersect\_size      *Get Overlap Size between Interval x and y*

---

**Description**

Get Overlap Size between Interval x and y

**Usage**

```
get_intersect_size(x.start, x.end, y.start, y.end)
```

**Arguments**

x.start	start position of interval x.
x.end	start position of interval x.
y.start	start position of interval x.
y.end	start position of interval x.

**Value**

a numeric vector.

**Examples**

```
o1 <- get_intersect_size(1, 5, 3, 20)
o1
o2 <- get_intersect_size(3, 20, 1, 10)
o2
o3 <- get_intersect_size(c(1, 2, 1), c(10, 4, 6), c(4, 2, 5), c(10, 3, 22))
o3
```

---

get_pLOH_score	<i>Get proportions of pLOH score from Allele Specific Copy Number Profile</i>
----------------	---

---

**Description**

pLOH score represents the genome that displayed LOH.

**Usage**

```
get_pLOH_score(data, rm_chrs = c("chrX", "chrY"), genome_build = "hg19")
```

**Arguments**

data	a CopyNumber object or a data.frame containing at least 'chromosome', 'start', 'end', 'segVal', "minor_cn", 'sample' these columns.
rm_chrs	chromosomes to be removed in calculation. Default is sex chromosomes (recommended).
genome_build	genome build version, should be 'hg19', 'hg38', 'mm9' or 'mm10'.

**Value**

A data.frame

**References**

Steele, Christopher D., et al. "Signatures of copy number alterations in human cancer." bioRxiv (2021).



**Examples**

```
# Load toy dataset of absolute copynumber profile
load(system.file("extdata", "toy_segTab.RData",
  package = "sigminer", mustWork = TRUE
))

set.seed(1234)
segTabs$minor_cn <- sample(c(0, 1), size = nrow(segTabs), replace = TRUE)
cn <- read_copynumber(segTabs,
  seg_cols = c("chromosome", "start", "end", "segVal"),
  genome_measure = "wg", complement = TRUE, add_loh = TRUE
)

df <- get_pLOH_score(cn)
df

df2 <- get_pLOH_score(cn@data)
df2
```

---

```
get_shannon_diversity_index
```

*Get Shannon Diversity Index for Signatures*

---

**Description**

$$H = - \sum_{i=1}^n p_i \ln(p_i)$$

where  $n$  is the number of signatures identified in the signature with exposure > cutoff, and  $p_i$  is the normalized exposure of the  $i$ th signature with exposure > cutoff. Exposures of signatures were normalized to sum to 1.

**Usage**

```
get_shannon_diversity_index(rel_expo, cutoff = 0.001)
```

**Arguments**

**rel\_expo** a data.frame with numeric columns indicating **relative** signature exposures for each sample. Typically this data can be obtained from [get\\_sig\\_exposure\(\)](#).

**cutoff** a relative exposure cutoff for filtering signatures, default is 0.1%.

**Value**

a data.frame

## References

Steele, Christopher D., et al. "Undifferentiated sarcomas develop through distinct evolutionary pathways." *Cancer Cell* 35.3 (2019): 441-456.

## Examples

```
# Load mutational signature
load(system.file("extdata", "toy_mutational_signature.RData",
  package = "sigminer", mustWork = TRUE
))
# Get signature exposure
rel_expo <- get_sig_exposure(sig2, type = "relative")
rel_expo
diversity_index <- get_shannon_diversity_index(rel_expo)
diversity_index
```

---

```
get_sig_cancer_type_index
```

*Obtain Signature Index for Cancer Types*

---

## Description

Obtain Signature Index for Cancer Types

## Usage

```
get_sig_cancer_type_index(
  sig_type = c("legacy", "SBS", "DBS", "ID"),
  seq_type = c("WGS", "WES"),
  source = c("PCAWG", "TCGA", "nonPCAWG"),
  keyword = NULL
)
```

## Arguments

sig_type	signature type.
seq_type	sequencing type.
source	data source.
keyword	keyword to search in the signature index database.

## Value

a list.

### Examples

```

11 <- get_sig_cancer_type_index()
12 <- get_sig_cancer_type_index(sig_type = "SBS")
13 <- get_sig_cancer_type_index(sig_type = "DBS", source = "PCAWG", seq_type = "WGS")
14 <- get_sig_cancer_type_index(sig_type = "ID")
15 <- get_sig_cancer_type_index(keyword = "breast")
11
12
13
14
15

```

---

get\_sig\_db

*Get Curated Reference Signature Database*


---

### Description

Reference mutational signatures and their aetiologies, mainly obtained from COSMIC database (SigProfiler results) and cleaned before saving into **sigminer** package. You can obtain:

- COSMIC legacy SBS signatures.
- COSMIC v3 SBS signatures.
- COSMIC v3 DBS signatures.
- COSMIC v3 ID (indel) signatures.
- SBS and RS (rearrangement) signatures from Nik lab 2020 Nature Cancer paper.
- RS signatures from BRCA560 and USARC cohorts.
- Copy number signatures from USARC cohort and TCGA.

### Usage

```
get_sig_db(sig_db = "legacy")
```

### Arguments

`sig_db` default 'legacy', it can be 'legacy' (for **COSMIC v2 'SBS'**), 'SBS', 'DBS', 'ID' and 'TSB' (for **COSMIV v3.1 signatures**) for small scale mutations. For more specific details, it can also be 'SBS\_hg19', 'SBS\_hg38', 'SBS\_mm9', 'SBS\_mm10', 'DBS\_hg19', 'DBS\_hg38', 'DBS\_mm9', 'DBS\_mm10' to use COSMIC v3 reference signatures from Alexandrov, Ludmil B., et al. (2020) (reference #1). In addition, it can be one of "SBS\_Nik\_lab\_Organ", "RS\_Nik\_lab\_Organ", "SBS\_Nik\_lab", "RS\_Nik\_lab" to refer reference signatures from Degasper, Andrea, et al. (2020) (reference #2); "RS\_BRCA560", "RS\_USARC" to reference signatures from BRCA560 and USARC cohorts; "CNS\_USARC" (40 categories), "CNS\_TCGA" (48 categories) to reference copy number signatures from USARC cohort and TCGA. **UPDATE**, the latest version of reference version can be automatically downloaded and loaded from <https://cancer>.

[sanger.ac.uk/signatures/downloads/](https://www.sanger.ac.uk/signatures/downloads/) when a option with latest\_prefix is specified (e.g. "latest\_SBS\_GRCh37"). **Note:** the signature profile for different genome builds are basically same. And specific database (e.g. 'SBS\_mm10') contains less signatures than all COSMIC signatures (because some signatures are not detected from Alexandrov, Ludmil B., et al. (2020)). For all available options, check the parameter setting.

## Value

a list.

## References

- Steele, Christopher D., et al. "Signatures of copy number alterations in human cancer." bioRxiv (2021).
- Alexandrov, Ludmil B., et al. "The repertoire of mutational signatures in human cancer." Nature 578.7793 (2020): 94-101.
- Steele, Christopher D., et al. "Undifferentiated sarcomas develop through distinct evolutionary pathways." Cancer Cell 35.3 (2019): 441-456.

## See Also

[get\\_sig\\_similarity](#), [sig\\_fit](#) and [show\\_cosmic\\_sig\\_profile](#).

## Examples

```
s1 <- get_sig_db()
s2 <- get_sig_db("SBS")
s3 <- get_sig_db("DBS")
s4 <- get_sig_db("DBS_mm10")
s5 <- get_sig_db("SBS_Nik_lab")
s6 <- get_sig_db("ID")
s7 <- get_sig_db("RS_BRCA560")
s8 <- get_sig_db("RS_USARC")
s9 <- get_sig_db("RS_Nik_lab")
s10 <- get_sig_db("CNS_USARC")
s11 <- get_sig_db("CNS_TCGA")
s1
s2
s3
s4
s5
s6
s7
s8
s9
s10
s11
```

---

get\_sig\_exposure      *Get Signature Exposure from 'Signature' Object*

---

## Description

The expected number of mutations (or copy number segment records) with each signature was determined after a scaling transformation  $V \sim WH = W'H'$  where  $W' = WU'$  and  $H' = UH$ . The scaling matrix  $U$  is a  $K \times K$  diagonal matrix ( $K$  is signature number,  $U'$  is the inverse of  $U$ ) with the element corresponding to the L1-norm of column vectors of  $W$  (ie. the sum of the elements of the vector). As a result, the  $k$ -th row vector of the final matrix  $H'$  represents the absolute exposure (activity) of the  $k$ -th process across samples (e.g., for SBS, the estimated (or expected) number of mutations generated by the  $k$ -th process). Of note, for copy number signatures, only components of feature CN was used for calculating  $H'$ .

## Usage

```
get_sig_exposure(
  Signature,
  type = c("absolute", "relative"),
  rel_threshold = 0.01
)
```

## Arguments

Signature	a Signature object obtained either from <a href="#">sig_extract</a> or <a href="#">sig_auto_extract</a> , or just a raw exposure matrix with column representing samples (patients) and row representing signatures.
type	'absolute' for signature exposure and 'relative' for signature relative exposure.
rel_threshold	only used when type is 'relative', relative exposure less than ( $\leq$ ) this value will be set to 0 and thus all signature exposures may not sum to 1. This is similar to this argument in <a href="#">sig_fit</a> .

## Value

a `data.table`

## Author(s)

Shixiang Wang [w\\_shixiang@163.com](mailto:w_shixiang@163.com)

## References

Kim, Jaegil, et al. "Somatic ERCC2 mutations are associated with a distinct genomic signature in urothelial tumors." *Nature genetics* 48.6 (2016): 600.

**Examples**

```
# Load mutational signature
load(system.file("extdata", "toy_mutational_signature.RData",
  package = "sigminer", mustWork = TRUE
))
# Get signature exposure
expo1 <- get_sig_exposure(sig2)
expo1
expo2 <- get_sig_exposure(sig2, type = "relative")
expo2
```

---

```
get_sig_feature_association
```

*Calculate Association between Signature Exposures and Other Features*

---

**Description**

Association of signature exposures with other features will be performed using one of two procedures: for a continuous association variable (including ordinal variable), correlation is performed; for a binary association variable, samples will be divided into two groups and Mann-Whitney U-test is performed to test for differences in signature exposure medians between the two groups. See [get\\_tidy\\_association](#) for cleaning association result.

**Usage**

```
get_sig_feature_association(
  data,
  cols_to_sigs,
  cols_to_features,
  type = "ca",
  method_co = c("spearman", "pearson", "kendall"),
  method_ca = stats::wilcox.test,
  min_n = 0.01,
  verbose = FALSE,
  ...
)
```

**Arguments**

data	a data.frame contains signature exposures and other features
cols_to_sigs	colnames for signature exposure
cols_to_features	colnames for other features
type	a character vector containing 'ca' for categorical variable and 'co' for continuous variable, it must have the same length as cols_to_features.

method_co	method for continuous variable, default is "spearman", could also be "pearson" and "kendall".
method_ca	method for categorical variable, default is "wilcox.test"
min_n	a minimal fraction (e.g. 0.01) or a integer number (e.g. 10) for filtering some variables with few positive events. Default is 0.01.
verbose	if TRUE, print extra message.
...	other arguments passing to test functions, like cor.test.

**Value**

a list. For 'co' features, 'measure' means correlation coefficient. For 'ca' features, 'measure' means difference in means of signature exposure.

**See Also**

[get\\_tidy\\_association](#)

---

get\_sig\_rec\_similarity

*Get Reconstructed Profile Cosine Similarity, RSS, etc.*

---

**Description**

See [bp\\_extract\\_signatures](#) for examples.

**Usage**

```
get_sig_rec_similarity(Signature, nmf_matrix)
```

**Arguments**

Signature	a Signature object.
nmf_matrix	a matrix used for NMF decomposition with rows indicate samples and columns indicate components.

**Value**

a data.table.

---

get_sig_similarity	<i>Calculate Similarity between Identified Signatures and Reference Signatures</i>
--------------------	--

---

## Description

The reference signatures can be either a Signature object specified by Ref argument or known COSMIC signatures specified by sig\_db argument. Two COSMIC databases are used for comparisons - "legacy" which includes 30 signatures, and "SBS" - which includes updated/refined 65 signatures. This function is modified from compareSignatures() in **maftools** package. **NOTE:** all reference signatures are generated from gold standard tool: SigProfiler.

## Usage

```
get_sig_similarity(
  Signature,
  Ref = NULL,
  sig_db = c("legacy", "SBS", "DBS", "ID", "TSB", "SBS_Nik_lab", "RS_Nik_lab",
    "RS_BRCA560", "RS_USARC", "CNS_USARC", "CNS_TCGA", "SBS_hg19", "SBS_hg38", "SBS_mm9",
    "SBS_mm10", "DBS_hg19", "DBS_hg38", "DBS_mm9", "DBS_mm10", "SBS_Nik_lab_Organ",
    "RS_Nik_lab_Organ", "latest_SBS_GRCh37", "latest_DBS_GRCh37", "latest_ID_GRCh37",
    "latest_SBS_GRCh38", "latest_DBS_GRCh38", "latest_SBS_mm9", "latest_DBS_mm9",
    "latest_SBS_mm10", "latest_DBS_mm10", "latest_SBS_rn6", "latest_DBS_rn6"),
  db_type = c("", "human-exome", "human-genome"),
  method = "cosine",
  normalize = c("row", "feature"),
  feature_setting = sigminer::CN.features,
  set_order = TRUE,
  pattern_to_rm = NULL,
  verbose = TRUE
)
```

## Arguments

Signature	a Signature object or a component-by-signature matrix/data.frame (sum of each column is 1) or a normalized component-by-sample matrix/data.frame (sum of each column is 1). More please see examples.
Ref	default is NULL, can be a same object as Signature.
sig_db	default 'legacy', it can be 'legacy' (for <b>COSMIC v2 'SBS'</b> ), 'SBS', 'DBS', 'ID' and 'TSB' (for <b>COSMIV v3.1 signatures</b> ) for small scale mutations. For more specific details, it can also be 'SBS_hg19', 'SBS_hg38', 'SBS_mm9', 'SBS_mm10', 'DBS_hg19', 'DBS_hg38', 'DBS_mm9', 'DBS_mm10' to use COSMIC v3 reference signatures from Alexandrov, Ludmil B., et al. (2020) (reference #1). In addition, it can be one of "SBS_Nik_lab_Organ", "RS_Nik_lab_Organ", "SBS_Nik_lab", "RS_Nik_lab" to refer reference signatures from Degasper, Andrea, et al. (2020) (reference #2); "RS_BRCA560", "RS_USARC" to reference signatures from BRCA560 and USARC cohorts; "CNS_USARC" (40



categories), "CNS\_TCGA" (48 categories) to reference copy number signatures from USARC cohort and TCGA. **UPDATE**, the latest version of reference version can be automatically downloaded and loaded from <https://cancer.sanger.ac.uk/signatures/downloads/> when a option with latest\_prefix is specified (e.g. "latest\_SBS\_GRCh37"). **Note**: the signature profile for different genome builds are basically same. And specific database (e.g. 'SBS\_mm10') contains less signatures than all COSMIC signatures (because some signatures are not detected from Alexandrov, Ludmil B., et al. (2020)). For all available options, check the parameter setting.

db_type	only used when sig_db is enabled. "" for keeping default, "human-exome" for transforming to exome frequency of component, and "human-genome" for transforming to whole genome frequency of component. Currently only works for 'SBS'.
method	default is 'cosine' for cosine similarity.
normalize	one of "row" and "feature". "row" is typically used for common mutational signatures. "feature" is designed by me to use when input are copy number signatures.
feature_setting	a data.frame used for classification. <b>Only used when method is "Wang" ("W")</b> . Default is <code>CN.features</code> . Users can also set custom input with "feature", "min" and "max" columns available. Valid features can be printed by <code>unique(CN.features\$feature)</code> .
set_order	if TRUE, order the return similarity matrix.
pattern_to_rm	patterns for removing some features/components in similarity calculation. A vector of component name is also accepted. The remove operation will be done after normalization. Default is NULL.
verbose	if TRUE, print extra info.

### Value

a list containing similarities, aetiologies if available, best match and RSS.

### Author(s)

Shixiang Wang [w\\_shixiang@163.com](mailto:w_shixiang@163.com)

### References

- Alexandrov, Ludmil B., et al. "The repertoire of mutational signatures in human cancer." *Nature* 578.7793 (2020): 94-101.
- Degasperi, Andrea, et al. "A practical framework and online tool for mutational signature analyses show intertissue variation and driver dependencies." *Nature cancer* 1.2 (2020): 249-263.
- Steele, Christopher D., et al. "Undifferentiated sarcomas develop through distinct evolutionary pathways." *Cancer Cell* 35.3 (2019): 441-456.
- Nik-Zainal, Serena, et al. "Landscape of somatic mutations in 560 breast cancer whole-genome sequences." *Nature* 534.7605 (2016): 47-54.

Steele, Christopher D., et al. "Signatures of copy number alterations in human cancer." bioRxiv (2021).

### Examples

```
# Load mutational signature
load(system.file("extdata", "toy_mutational_signature.RData",
  package = "sigminer", mustWork = TRUE
))

s1 <- get_sig_similarity(sig2, Ref = sig2)
s1

s2 <- get_sig_similarity(sig2)
s2
s3 <- get_sig_similarity(sig2, sig_db = "SBS")
s3

# Set order for result similarity matrix
s4 <- get_sig_similarity(sig2, sig_db = "SBS", set_order = TRUE)
s4

## Remove some components
## in similarity calculation
s5 <- get_sig_similarity(sig2,
  Ref = sig2,
  pattern_to_rm = c("T[T>G]C", "T[T>G]G", "T[T>G]T")
)
s5

## Same to DBS and ID signatures
x1 <- get_sig_db("DBS_hg19")
x2 <- get_sig_db("DBS_hg38")
s6 <- get_sig_similarity(x1$db, x2$db)
s6
```

---

get\_tidy\_association *Get Tidy Signature Association Results*

---

### Description

Get Tidy Signature Association Results

### Usage

```
get_tidy_association(cor_res, p_adjust = FALSE, method = "fdr")
```

**Arguments**

cor_res	data returned by <a href="#">get_sig_feature_association()</a>
p_adjust	logical, if TRUE, adjust p values by data type.
method	p value correction method, see <a href="#">stats::p.adjust</a> for more detail.

**Value**

a data.frame

**See Also**

[get\\_sig\\_feature\\_association](#)

---

group_enrichment	<i>General Group Enrichment Analysis</i>
------------------	--

---

**Description**

This function takes a data.frame as input, compares proportion of positive cases or mean measure in one subgroup and the remaining samples.

**Usage**

```
group_enrichment(
  df,
  grp_vars = NULL,
  enrich_vars = NULL,
  cross = TRUE,
  co_method = c("t.test", "wilcox.test"),
  ref_group = NA
)
```

**Arguments**

df	a data.frame.
grp_vars	character vector specifying group variables to split samples into subgroups (at least 2 subgroups, otherwise this variable will be skipped).
enrich_vars	character vector specifying measure variables to be compared. If variable is not numeric, only binary cases are accepted in the form of TRUE/FALSE or P/N (P for positive cases and N for negative cases). Of note, NA values set to negative cases.
cross	logical, default is TRUE, combine all situations provided by grp_vars and enrich_vars. For examples, c('A', 'B') and c('C', 'D') will construct 4 combinations(i.e. "AC", "AD", "BC" and "BD"). A variable can not be in both grp_vars and enrich_vars, such cases will be automatically drop. If FALSE, use pairwise combinations, see section "examples" for use cases.

co\_method        test method for continuous variable, default is 't.test'.  
 ref\_group        reference group set in grp\_vars.

### Value

a `data.table` with following columns:

- `grp_var`: group variable name.
- `enrich_var`: enrich variable (variable to be compared) name.
- `grp1`: the first group name, should be a member in `grp_var` column.
- `grp2`: the remaining samples, marked as 'Rest'.
- `grp1_size`: sample size for `grp1`.
- `grp1_pos_measure`: for binary variable, it stores the proportion of positive cases in `grp1`; for continuous variable, it stores mean value.
- `grp2_size`: sample size for `grp2`.
- `grp2_pos_measure`: same as `grp1_pos_measure` but for `grp2`.
- `measure_observed`: for binary variable, it stores odds ratio; for continuous variable, it stores scaled mean ratio.
- `measure_tested`: only for binary variable, it stores estimated odds ratio and its 95% CI from `fisher.test()`.
- `p_value`: for binary variable, it stores p value from `fisher.test()`; for continuous variable, it stores value from `wilcox.test()` or `t.test()`.
- `type`: one of "binary" and "continuous".
- `method`: one of "fish.test", "wilcox.test" and "t.test".

### See Also

[show\\_group\\_enrichment](#)

### Examples

```
set.seed(1234)
df <- dplyr::tibble(
  g1 = factor(abs(round(rnorm(99, 0, 1)))),
  g2 = rep(LETTERS[1:4], c(50, 40, 8, 1)),
  e1 = sample(c("P", "N"), 99, replace = TRUE),
  e2 = rnorm(99)
)

print(str(df))
print(head(df))

# Compare g1:e1, g1:e2, g2:e1 and g2:e2
x1 <- group_enrichment(df, grp_vars = c("g1", "g2"), enrich_vars = c("e1", "e2"))
x1

# Only compare g1:e1, g2:e2
```

```
x2 <- group_enrichment(df,
  grp_vars = c("g1", "g2"),
  enrich_vars = c("e1", "e2"),
  co_method = "wilcox.test",
  cross = FALSE
)
x2

# Visualization
p1 <- show_group_enrichment(x1, fill_by_p_value = TRUE)
p1
p2 <- show_group_enrichment(x1, fill_by_p_value = FALSE)
p2
p3 <- show_group_enrichment(x1, return_list = TRUE)
p3
```

---

handle\_hyper\_mutation *Handle Hypermutant Samples*

---

### Description

This can be used for SNV/INDEL count matrix. For copy number analysis, please skip it.

### Usage

```
handle_hyper_mutation(nmf_matrix)
```

### Arguments

`nmf_matrix` a matrix used for NMF decomposition with rows indicate samples and columns indicate components.

### Value

a matrix.

### References

Kim, Jaegil, et al. "Somatic ERCC2 mutations are associated with a distinct genomic signature in urothelial tumors." *Nature genetics* 48.6 (2016): 600.

---

hello	<i>Say Hello to Users</i>
-------	---------------------------

---

**Description**

Say Hello to Users

**Usage**

hello()

**Examples**

hello()

---

MAF-class	<i>Class MAF</i>
-----------	------------------

---

**Description**

S4 class for storing summarized MAF. It is from `maftools` package.

**Details**

More about MAF object please see [maftools](#).

**Slots**

`data` `data.table` of MAF file containing all non-synonymous variants.

`variants.per.sample` table containing variants per sample

`variant.type.summary` table containing variant types per sample

`variant.classification.summary` table containing variant classification per sample

`gene.summary` table containing variant classification per gene

`summary` table with basic MAF summary stats

`maf.silent` subset of main MAF containing only silent variants

`clinical.data` clinical data associated with each sample/Tumor\_Sample\_Barcode in MAF.

---

output_bootstrap	<i>Output Signature Bootstrap Fitting Results</i>
------------------	---

---

### Description

Output Signature Bootstrap Fitting Results

### Usage

```
output_bootstrap(x, result_dir, mut_type = "SBS", sig_db = mut_type)
```

### Arguments

x	result from <a href="#">sig_fit_bootstrap_batch</a> .
result_dir	a result directory.
mut_type	one of 'SBS', 'DBS', 'ID' or 'CN'.
sig_db	default 'legacy', it can be 'legacy' (for <b>COSMIC v2 'SBS'</b> ), 'SBS', 'DBS', 'ID' and 'TSB' (for <b>COSMIV v3.1 signatures</b> ) for small scale mutations. For more specific details, it can also be 'SBS_hg19', 'SBS_hg38', 'SBS_mm9', 'SBS_mm10', 'DBS_hg19', 'DBS_hg38', 'DBS_mm9', 'DBS_mm10' to use COSMIC v3 reference signatures from Alexandrov, Ludmil B., et al. (2020) (reference #1). In addition, it can be one of "SBS_Nik_lab_Organ", "RS_Nik_lab_Organ", "SBS_Nik_lab", "RS_Nik_lab" to refer reference signatures from Degasperi, Andrea, et al. (2020) (reference #2); "RS_BRCA560", "RS_USARC" to reference signatures from BRCA560 and USARC cohorts; "CNS_USARC" (40 categories), "CNS_TCGA" (48 categories) to reference copy number signatures from USARC cohort and TCGA. <b>UPDATE</b> , the latest version of reference version can be automatically downloaded and loaded from <a href="https://cancer.sanger.ac.uk/signatures/downloads/">https://cancer.sanger.ac.uk/signatures/downloads/</a> when a option with latest_ prefix is specified (e.g. "latest_SBS_GRCh37"). <b>Note</b> : the signature profile for different genome builds are basically same. And specific database (e.g. 'SBS_mm10') contains less signatures than all COSMIC signatures (because some signatures are not detected from Alexandrov, Ludmil B., et al. (2020)). For all available options, check the parameter setting.

### Value

Nothing.

---

output_fit	<i>Output Signature Fitting Results</i>
------------	---

---

**Description**

Output Signature Fitting Results

**Usage**

```
output_fit(x, result_dir, mut_type = "SBS", sig_db = mut_type)
```

**Arguments**

x	result from <a href="#">sig_fit</a> .
result_dir	a result directory.
mut_type	one of 'SBS', 'DBS', 'ID' or 'CN'.
sig_db	default 'legacy', it can be 'legacy' (for <b>COSMIC v2 'SBS'</b> ), 'SBS', 'DBS', 'ID' and 'TSB' (for <b>COSMIV v3.1 signatures</b> ) for small scale mutations. For more specific details, it can also be 'SBS_hg19', 'SBS_hg38', 'SBS_mm9', 'SBS_mm10', 'DBS_hg19', 'DBS_hg38', 'DBS_mm9', 'DBS_mm10' to use COSMIC v3 reference signatures from Alexandrov, Ludmil B., et al. (2020) (reference #1). In addition, it can be one of "SBS_Nik_lab_Organ", "RS_Nik_lab_Organ", "SBS_Nik_lab", "RS_Nik_lab" to refer reference signatures from Degasperi, Andrea, et al. (2020) (reference #2); "RS_BRCA560", "RS_USARC" to reference signatures from BRCA560 and USARC cohorts; "CNS_USARC" (40 categories), "CNS_TCGA" (48 categories) to reference copy number signatures from USARC cohort and TCGA. <b>UPDATE</b> , the latest version of reference version can be automatically downloaded and loaded from <a href="https://cancer.sanger.ac.uk/signatures/downloads/">https://cancer.sanger.ac.uk/signatures/downloads/</a> when a option with latest_ prefix is specified (e.g. "latest_SBS_GRCh37"). <b>Note</b> : the signature profile for different genome builds are basically same. And specific database (e.g. 'SBS_mm10') contains less signatures than all COSMIC signatures (because some signatures are not detected from Alexandrov, Ludmil B., et al. (2020)). For all available options, check the parameter setting.

**Value**

Nothing.



---

output_sig	<i>Output Signature Results</i>
------------	---------------------------------

---

**Description**

Output Signature Results

**Usage**

```
output_sig(sig, result_dir, mut_type = "SBS", sig_db = mut_type)
```

**Arguments**

sig	a Signature object.
result_dir	a result directory.
mut_type	one of 'SBS', 'DBS', 'ID' or 'CN'.
sig_db	default 'legacy', it can be 'legacy' (for <b>COSMIC v2 'SBS'</b> ), 'SBS', 'DBS', 'ID' and 'TSB' (for <b>COSMIV v3.1 signatures</b> ) for small scale mutations. For more specific details, it can also be 'SBS_hg19', 'SBS_hg38', 'SBS_mm9', 'SBS_mm10', 'DBS_hg19', 'DBS_hg38', 'DBS_mm9', 'DBS_mm10' to use COSMIC v3 reference signatures from Alexandrov, Ludmil B., et al. (2020) (reference #1). In addition, it can be one of "SBS_Nik_lab_Organ", "RS_Nik_lab_Organ", "SBS_Nik_lab", "RS_Nik_lab" to refer reference signatures from Degasperi, Andrea, et al. (2020) (reference #2); "RS_BRCA560", "RS_USARC" to reference signatures from BRCA560 and USARC cohorts; "CNS_USARC" (40 categories), "CNS_TCGA" (48 categories) to reference copy number signatures from USARC cohort and TCGA. <b>UPDATE</b> , the latest version of reference version can be automatically downloaded and loaded from <a href="https://cancer.sanger.ac.uk/signatures/downloads/">https://cancer.sanger.ac.uk/signatures/downloads/</a> when a option with latest_ prefix is specified (e.g. "latest_SBS_GRCh37"). <b>Note</b> : the signature profile for different genome builds are basically same. And specific database (e.g. 'SBS_mm10') contains less signatures than all COSMIC signatures (because some signatures are not detected from Alexandrov, Ludmil B., et al. (2020)). For all available options, check the parameter setting.

**Value**

Nothing.

---

output_tally	<i>Output Tally Result in Barplots</i>
--------------	--

---

**Description**

Output Tally Result in Barplots

**Usage**

```
output_tally(x, result_dir, mut_type = "SBS")
```

**Arguments**

x	a matrix with row representing components (motifs) and column representing samples.
result_dir	a result directory.
mut_type	one of 'SBS', 'DBS', 'ID' or 'CN'.

**Value**

Nothing.

---

read_copynumber	<i>Read Absolute Copy Number Profile</i>
-----------------	--

---

**Description**

Read **absolute** copy number profile for preparing CNV signature analysis. See detail part of [sig\\_tally\(\)](#) to see how to handle sex to get correct summary.

**Usage**

```
read_copynumber(
  input,
  pattern = NULL,
  ignore_case = FALSE,
  seg_cols = c("Chromosome", "Start.bp", "End.bp", "modal_cn"),
  samp_col = "sample",
  add_loh = FALSE,
  loh_min_len = 10000,
  loh_min_frac = 0.05,
  join_adj_seg = TRUE,
  skip_annotation = FALSE,
  use_all = add_loh,
  min_segnum = 0L,
```

```

    max_copynumber = 20L,
    genome_build = c("hg19", "hg38", "mm10", "mm9"),
    genome_measure = c("called", "wg"),
    complement = FALSE,
    ...
)

```

## Arguments

input	a data.frame or a file or a directory contains copy number profile.
pattern	an optional regular expression used to select part of files if input is a directory, more detail please see <a href="#">list.files()</a> function.
ignore_case	logical. Should pattern-matching be case-insensitive?
seg_cols	four strings used to specify chromosome, start position, end position and copy number value in input, respectively. Default use names from ABSOLUTE calling result.
samp_col	a character used to specify the sample column name. If input is a directory and cannot find samp_col, sample names will use file names (set this parameter to NULL is recommended in this case).
add_loh	if TRUE, add LOH labels to segments. <b>NOTE</b> a column 'minor_cn' must exist to indicate minor allele copy number value. Sex chromosome will not be labeled.
loh_min_len	The length cut-off for labeling a segment as 'LOH'. Default is 10Kb.
loh_min_frac	When join_adj_seg set to TRUE, only the length fraction of LOH region is larger than this value will be labeled as 'LOH'. Default is 30%.
join_adj_seg	if TRUE (default), join adjacent segments with same copy number value. This is helpful for precisely count the number of breakpoint. When set use_all=TRUE, the mean function will be applied to extra numeric columns and unique string columns will be pasted by comma for joined records.
skip_annotation	if TRUE, skip annotation step, it may affect some analysis and visualization functionality, but speed up reading data.
use_all	default is FALSE. If True, use all columns from raw input.
min_segnum	minimal number of copy number segments within a sample.
max_copynumber	bigger copy number within a sample will be reset to this value.
genome_build	genome build version, should be 'hg19', 'hg38', 'mm9' or 'mm10'.
genome_measure	default is 'called', can be 'wg' or 'called'. Set 'called' will use called segments size to compute total size for CNA burden calculation, this option is useful for WES and target sequencing. Set 'wg' will use autosome size from genome build, this option is useful for WGS, SNP etc..
complement	if TRUE, complement chromosome (except 'Y') does not show in input data with normal copy 2.
...	other parameters pass to <a href="#">data.table::fread()</a>

**Value**

a [CopyNumber](#) object.

**Author(s)**

Shixiang Wang [w\\_shixiang@163.com](mailto:w_shixiang@163.com)

**See Also**

[read\\_maf](#) for reading mutation data to [MAF](#) object.

**Examples**

```
# Load toy dataset of absolute copynumber profile
load(system.file("extdata", "toy_segTab.RData",
  package = "sigminer", mustWork = TRUE
))
cn <- read_copynumber(segTabs,
  seg_cols = c("chromosome", "start", "end", "segVal"),
  genome_build = "hg19", complement = FALSE
)
cn
cn_subset <- subset(cn, sample == "TCGA-DF-A2KN-01A-11D-A17U-01")

# Add LOH
set.seed(1234)
segTabs$minor_cn <- sample(c(0, 1), size = nrow(segTabs), replace = TRUE)
cn <- read_copynumber(segTabs,
  seg_cols = c("chromosome", "start", "end", "segVal"),
  genome_measure = "wg", complement = TRUE, add_loh = TRUE
)
# Use tally method "S" (Steele et al.)
tally_s <- sig_tally(cn, method = "S")

tab_file <- system.file("extdata", "metastatic_tumor.segtab.txt",
  package = "sigminer", mustWork = TRUE
)
cn2 <- read_copynumber(tab_file)
cn2
```

---

`read_copynumber_ascat` *Read Copy Number Data from ASCAT Result Files*

---

**Description**

Note, the result is not a [CopyNumber](#) object, you need to generate it by yourself.

**Usage**

```
read_copynumber_ascat(x)
```

**Arguments**

x                    one or more .rds format files which contains ASCAT object from result of `ascat.runAscat()` in **ASCAT** package.

**Value**

a tidy list.

---

read\_copynumber\_seqz    *Read Absolute Copy Number Profile from Sequenza Result Directory*

---

**Description**

Read Absolute Copy Number Profile from Sequenza Result Directory

**Usage**

```
read_copynumber_seqz(target_dir, return_df = FALSE, ...)
```

**Arguments**

target\_dir        a directory path.  
 return\_df        if TRUE, return a data.frame directly, otherwise return a [CopyNumber](#) object.  
 ...               other parameters passing to [read\\_copynumber\(\)](#).

**Value**

a data.frame or a CopyNumber object.

---

read\_maf                *Read MAF Files*

---

**Description**

This function is a wrapper of `maftools::read.maf`. Useless options in `maftools::read.maf` are dropped here. You can also use `maftools::read.maf` to read the data. All reference alleles and mutation alleles should be recorded in positive strand format.

**Usage**

```
read_maf(maf, verbose = TRUE)
```

**Arguments**

maf	tab delimited MAF file. File can also be gz compressed. Required. Alternatively, you can also provide already read MAF file as a dataframe.
verbose	TRUE logical. Default to be talkative and prints summary.

**See Also**

[read\\_copynumber](#) for reading copy number data to [CopyNumber](#) object.

**Examples**

```
lam1.maf <- system.file("extdata", "tcga_lam1.maf.gz", package = "maftools", mustWork = TRUE)
if (!require("R.utils")) {
  message("Please install 'R.utils' package firstly")
} else {
  lam1 <- read_maf(maf = lam1.maf)
  lam1
}
```

---

read\_sv\_as\_rs

*Read Structural Variation Data as RS object*


---

**Description**

Read Structural Variation Data as RS object

**Usage**

```
read_sv_as_rs(input)
```

**Arguments**

input	a data.frame or a file with the following columns: "sample", "chr1", "start1", "end1", "chr2", "start2", "end2", "strand1", "strand2", "svclass". NOTE: If column "svclass" already exists in input, "strand1" and "strand2" are optional. If "svclass" is not provided, read_sv_as_rs() will compute it by "strand1", "strand2"(strand1/strand2), "chr" and "chr2": <ul style="list-style-type: none"> <li>• translocation, if mates are on different chromosomes.</li> <li>• inversion (+/-) and (-/+), if mates on the same chromosome.</li> <li>• deletion (+/+), if mates on the same chromosome.</li> <li>• tandem-duplication (-/-), if mates on the same chromosome.</li> </ul>
-------	---

**Value**

a list

## Examples

```
sv <- readRDS(system.file("extdata", "toy_sv.rds", package = "sigminer", mustWork = TRUE))
rs <- read_sv_as_rs(sv)
# svclass is optional
rs2 <- read_sv_as_rs(sv[, setdiff(colnames(sv), "svclass")])
identical(rs, rs2)

tally_rs <- sig_tally(rs)
```

---

read\_vcf

*Read VCF Files as MAF Object*

---

## Description

MAF file is more recommended. In this function, we will mimic the MAF object from the key c(1, 2, 4, 5, 7) columns of VCF file.

## Usage

```
read_vcf(
  vcfs,
  samples = NULL,
  genome_build = c("hg19", "hg38", "mm10", "mm9"),
  keep_only_pass = FALSE,
  verbose = TRUE
)
```

## Arguments

vcfs	VCF file paths.
samples	sample names for VCF files.
genome_build	genome build version like "hg19".
keep_only_pass	if TRUE, keep only 'PASS' mutation for analysis.
verbose	if TRUE, print extra info.

## Value

a [MAF](#).

## See Also

[read\\_maf](#), [read\\_copynumber](#)

## Examples

```
vcfs <- list.files(system.file("extdata", package = "sigminer"), "*.vcf", full.names = TRUE)

maf <- read_vcf(vcfs)
maf <- read_vcf(vcfs, keep_only_pass = TRUE)
```

---

read\_xena\_variants      *Read UCSC Xena Variant Format Data as MAF Object*

---

## Description

Read UCSC Xena Variant Format Data as MAF Object

## Usage

```
read_xena_variants(path)
```

## Arguments

path                    a path to variant file.

## Value

a MAF object.

## Examples

```
if (requireNamespace("UCSCXenaTools")) {
  library(UCSCXenaTools)
  options(use_hiplot = TRUE)
  example_file <- XenaGenerate(subset = XenaDatasets == "mc3/ACC_mc3.txt") %>%
    XenaQuery() %>%
    XenaDownload()
  x <- read_xena_variants(example_file$destfiles)
  x@data
  y <- sig_tally(x)
  y
}
```



---

 report\_bootstrap\_p\_value

*Report P Values from bootstrap Results*


---

**Description**

See examples in [sig\\_fit\\_bootstrap](#).

**Usage**

```
report_bootstrap_p_value(x, thresholds = c(0.01, 0.05, 0.1))
```

**Arguments**

x                    a (list of) result from [sig\\_fit\\_bootstrap](#).  
 thresholds        a vector of relative exposure threshold for calculating p values.

**Value**

a (list of) matrix

---

 same\_size\_clustering    *Same Size Clustering*


---

**Description**

This is a wrapper for several implementation that classify samples into same size clusters, the details please see [this blog](#). The source code is modified based on code from the blog.

**Usage**

```
same_size_clustering(  
  mat,  
  diss = FALSE,  
  clsize = NULL,  
  algo = c("nnit", "hcbottom", "kmvar"),  
  method = c("maxd", "random", "mind", "elki", "ward.D", "average", "complete",  
             "single")  
)
```

**Arguments**

mat                a data/distance matrix.  
 diss              if TRUE, treat mat as a distance matrix.  
 clsize            integer, number of sample within a cluster.  
 algo              algorithm.  
 method            method.

**Value**

a vector.

**Examples**

```
set.seed(1234L)
x <- rbind(
  matrix(rnorm(100, sd = 0.3), ncol = 2),
  matrix(rnorm(100, mean = 1, sd = 0.3), ncol = 2)
)
colnames(x) <- c("x", "y")

y1 <- same_size_clustering(x, clsize = 10)
y11 <- same_size_clustering(as.matrix(dist(x)), clsize = 10, diss = TRUE)

y2 <- same_size_clustering(x, clsize = 10, algo = "hcbottom", method = "ward.D")

y3 <- same_size_clustering(x, clsize = 10, algo = "kmvar")
y33 <- same_size_clustering(as.matrix(dist(x)), clsize = 10, algo = "kmvar", diss = TRUE)
```

---

scoring

*Score Copy Number Profile*

---

**Description**

Returns quantification of copy number profile and events including tandem duplication and Chromothripsis etc. Only copy number data from autosome is used here. **Some of the quantification methods are rough, you use at your risk.** You should do some extra work to check the result scores.

**Usage**

```
scoring(object, TD_size_cutoff = c(1000, 1e+05, 2e+06), TD_cn_cutoff = Inf)
```

**Arguments**

object	a object of <a href="#">CopyNumber</a> .
TD_size_cutoff	a length-3 numeric vector used to specify the start, midpoint, end segment size for determining tandem duplication size range, midpoint is used to split TD into short TD and long TD. Default is 1Kb to 100Kb for short TD, 100Kb to 2Mb for long TD.
TD_cn_cutoff	a number defining the maximum copy number of TD, default is Inf, i.e. no cutoff.

**Value**

a data . table with following scores:

- cnaBurden: CNA burden representing the altered genomic fraction as previously reported.
- cnaLoad: CNA load representing the quantity of copy number alteration.
- MACN: mean altered copy number (MACN) reflecting the property of altered copy number segments, calculated as

$$MACN = \frac{\sum_i CN_i}{N_{cnv}}$$

where  $CN_i$  is the copy number of altered segment  $i$ ,  $N_{cnv}$  is the number of CNV.

- weightedMACN: same as MACN but weighted with segment length.

$$MACN_{weighted} = \frac{\sum_i (CN_i \times L_i)}{\sum_i L_i}$$

where  $L_i$  is the length of altered copy number segment  $i$ .

- Ploidy: ploidy, the formula is same as weightedMACN but using all copy number segments instead of altered copy number segments.
- TDP\_pnas: tandem duplication phenotype score from <https://www.pnas.org/doi/10.1073/pnas.1520010113>, the threshold  $k$  in reference is omitted.

$$TDP = - \frac{\sum_{chr} |TD_{obs} - TD_{exp}|}{TD_{total}}$$

where  $TD_{total}$  is the number of TD,  $TD_{obs}$  and  $TD_{exp}$  are observed number of TD and expected number of TD for each chromosome.

- TDP: tandem duplication score used defined by our group work, TD represents segment with copy number greater than 2.

$$TD = \frac{TD_{total}}{\sum_{chr} |TD_{obs} - TD_{exp}| + 1}$$

- sTDP: TDP score for short TD.
- lTDP: TDP score for long TD.
- TDP\_size : TDP region size (Mb).
- sTDP\_size: sTDP region size (Mb).
- lTDP\_size: lTDP region size(Mb).
- Chromoth\_state: chromothripsis state score, according to reference [doi:10.1016/j.cell.2013.02.023](https://doi.org/10.1016/j.cell.2013.02.023), chromothripsis frequently leads to massive loss of segments on the affected chromosome with segmental losses being interspersed with regions displaying normal (disomic) copy-number (e.g., copy-number states oscillating between copy-number = 1 and copy-number = 2), form tens to hundreds of locally clustered DNA rearrangements. Most of methods use both SV and CNV to infer chromothripsis, here we roughly quantify it with

$$\sum_{chr} N_{OsCN}^2$$

where  $N_{OsCN}$  is the number of oscillating copy number pattern "2-1-2" for each chromosome.

**Examples**

```
# Load copy number object
load(system.file("extdata", "toy_copynumber.RData",
  package = "sigminer", mustWork = TRUE
))

d <- scoring(cn)
d

d2 <- scoring(cn, TD_cn_cutoff = 4L)
d2
```

---

show\_catalogue

*Show Alteration Catalogue Profile*


---

**Description**

Show Alteration Catalogue Profile

**Usage**

```
show_catalogue(
  catalogue,
  mode = c("SBS", "copynumber", "DBS", "ID", "RS"),
  method = "Wang",
  normalize = c("raw", "row", "feature"),
  style = c("default", "cosmic"),
  samples = NULL,
  samples_name = NULL,
  x_lab = "Components",
  y_lab = "Counts",
  ...
)
```

**Arguments**

catalogue	result from <a href="#">sig_tally</a> or a matrix with row representing components (motifs) and column representing samples
mode	signature type for plotting, now supports 'copynumber', 'SBS', 'DBS', 'ID' and 'RS' (genome rearrangement signature).
method	method for copy number feature classification in <a href="#">sig_tally</a> , can be one of "Wang" ("W"), "S".
normalize	normalize method.
style	plot style, one of 'default' and 'cosmic'.
samples	default is NULL, show sum of all samples in one row. If not NULL, show specified samples.

samples_name	set the sample names shown in plot.
x_lab	x axis lab.
y_lab	y axis lab.
...	other arguments passing to <a href="#">show_sig_profile</a> .

**Value**

a ggplot object

**Examples**

```
data("simulated_catalogs")
p <- show_catalogue(simulated_catalogs$set1, style = "cosmic")
p
```

---

show_cn_circos	<i>Show Copy Number Profile in Circos</i>
----------------	---

---

**Description**

Another visualization method for copy number profile like [show\\_cn\\_profile](#).

**Usage**

```
show_cn_circos(
  data,
  samples = NULL,
  show_title = TRUE,
  chrs = paste0("chr", 1:22),
  genome_build = c("hg19", "hg38", "mm10", "mm9"),
  col = NULL,
  side = "inside",
  ...
)
```

**Arguments**

data	a <a href="#">CopyNumber</a> object or a data.frame containing at least 'chromosome', 'start', 'end', 'segVal' these columns.
samples	default is NULL, can be a character vector representing multiple samples or number of samples to show. If data argument is a data.frame, a column called sample must exist.
show_title	if TRUE (default), show title with sample ID.
chrs	chromosomes start with 'chr'.
genome_build	genome build version, used when data is a data.frame, should be 'hg19' or 'hg38'.

col colors for the heatmaps. If it is NULL, set to `circlize::colorRamp2(c(1, 2, 4), c("blue", "black", "red"))`.

side side of the heatmaps.

... other parameters passing to `circlize::circos.genomicHeatmap`.

**Value**

a circos plot

**Examples**

```
load(system.file("extdata", "toy_copynumber.RData",
  package = "sigminer", mustWork = TRUE
))

show_cn_circos(cn, samples = 1)
show_cn_circos(cn, samples = "TCGA-99-7458-01A-11D-2035-01")

## Remove title
show_cn_circos(cn, samples = 1, show_title = FALSE)

## Subset chromosomes
show_cn_circos(cn, samples = 1, chrs = c("chr1", "chr2", "chr3"))

## Arrange plots
layout(matrix(1:4, 2, 2))
show_cn_circos(cn, samples = 4)

layout(1) # reset layout
```

---

show\_cn\_components      *Show Copy Number Components*

---

**Description**

Show classified components ("Wang" ("W") method) for copy number data.

**Usage**

```
show_cn_components(
  parameters,
  method = "Wang",
  show_weights = TRUE,
  log_y = FALSE,
  return_plotlist = FALSE,
  base_size = 12,
  nrow = 2,
```

```

    align = "hv",
    ...
)

```

### Arguments

parameters	a data.frame contain parameter components, obtain this from <a href="#">sig_tally</a> function.
method	method for feature classification, can be one of "Wang" ("W"), "S" (for method described in Steele et al. 2019).
show_weights	default is TRUE, show weights for each component. Only used when method is "Macintyre".
log_y	logical, if TRUE, show log <sub>10</sub> based y axis, only works for input from "Wang" ("W") method.
return_plotlist	if TRUE, return a list of ggplot objects but a combined plot.
base_size	overall font size.
nrow	(optional) Number of rows in the plot grid.
align	(optional) Specifies whether graphs in the grid should be horizontally ("h") or vertically ("v") aligned. Options are "none" (default), "hv" (align in both directions), "h", and "v".
...	other options pass to <a href="#">plot_grid</a> function of <b>cowplot</b> package.

### Value

a ggplot object

### Author(s)

Shixiang Wang [w\\_shixiang@163.com](mailto:w_shixiang@163.com)

---

show\_cn\_distribution    *Show Copy Number Distribution either by Length or Chromosome*

---

### Description

Visually summarize copy number distribution either by copy number segment length or chromosome. Input is a [CopyNumber](#) object, genome\_build option will read from genome\_build slot of object.

**Usage**

```
show_cn_distribution(  
  data,  
  rm_normal = TRUE,  
  mode = c("ld", "cd"),  
  fill = FALSE,  
  scale_chr = TRUE,  
  base_size = 14  
)
```

**Arguments**

data	a <a href="#">CopyNumber</a> object.
rm_normal	logical. Whether remove normal copy (i.e. "segVal" equals 2), default is TRUE.
mode	either "ld" for distribution by CN length or "cd" for distribution by chromosome.
fill	when mode is "cd" and fill is TRUE, plot percentage instead of count.
scale_chr	logical. If TRUE, normalize count to per Megabase unit.
base_size	overall font size.

**Value**

a ggplot object

**Author(s)**

Shixiang Wang [w\\_shixiang@163.com](mailto:w_shixiang@163.com)

**Examples**

```
# Load copy number object  
load(system.file("extdata", "toy_copynumber.RData",  
  package = "sigminer", mustWork = TRUE  
)  
)  
# Plot distribution  
p1 <- show_cn_distribution(cn)  
p1  
p2 <- show_cn_distribution(cn, mode = "cd")  
p2  
p3 <- show_cn_distribution(cn, mode = "cd", fill = TRUE)  
p3
```



---

 show\_cn\_features      *Show Copy Number Feature Distributions*


---

**Description**

Show Copy Number Feature Distributions

**Usage**

```
show_cn_features(
  features,
  method = "Wang",
  rm_outlier = FALSE,
  ylab = NULL,
  log_y = FALSE,
  return_plotlist = FALSE,
  base_size = 12,
  nrow = 2,
  align = "hv",
  ...
)
```

**Arguments**

features	a feature list generate from <a href="#">sig_tally</a> function.
method	method for feature classification, can be one of "Wang" ("W"), "S" (for method described in Steele et al. 2019).
rm_outlier	default is FALSE, if TRUE, remove outliers. Only works when method is "Wang" ("W").
ylab	lab of y axis.
log_y	logical, if TRUE, show log <sub>10</sub> based y axis, only works for input from "Wang" ("W") method.
return_plotlist	if TRUE, return a list of ggplot objects but a combined plot.
base_size	overall font size.
nrow	(optional) Number of rows in the plot grid.
align	(optional) Specifies whether graphs in the grid should be horizontally ("h") or vertically ("v") aligned. Options are "none" (default), "hv" (align in both directions), "h", and "v".
...	other options pass to <a href="#">plot_grid</a> function of cowplot package.

**Value**

a ggplot object

---

show\_cn\_freq\_circos     *Show Copy Number Variation Frequency Profile with Circos*

---

## Description

Show Copy Number Variation Frequency Profile with Circos

## Usage

```
show_cn_freq_circos(
  data,
  groups = NULL,
  cutoff = 2L,
  resolution_factor = 1L,
  title = c("AMP", "DEL"),
  chrs = paste0("chr", 1:22),
  genome_build = c("hg19", "hg38", "mm10", "mm9"),
  cols = NULL,
  plot_ideogram = TRUE,
  track_height = 0.5,
  ideogram_height = 1,
  ...
)
```

## Arguments

data	a CopyNumber object or a data.frame containing at least 'chromosome', 'start', 'end', 'segVal', 'sample' these columns.
groups	a named list or a column name for specifying groups.
cutoff	copy number value cutoff for splitting data into AMP and DEL. The values equal to cutoff are discarded. Default is 2, you can also set a length-2 vector, e.g. c(2, 2).
resolution_factor	an integer to control the resolution. When it is 1 (default), compute frequency in each cytoband. When it is 2, use compute frequency in each half cytoband.
title	length-2 titles for AMP and DEL.
chrs	chromosomes start with 'chr'.
genome_build	genome build version, used when data is a data.frame, should be 'hg19' or 'hg38'.
cols	length-2 colors for AMP and DEL.
plot_ideogram	default is TRUE, show ideogram.
track_height	track height in mm unit.
ideogram_height	ideogram height in mm unit.
...	other parameters passing to <a href="#">circize::circos.genomicLines</a> .

**Value**

Nothing.

**Examples**

```
load(system.file("extdata", "toy_copynumber.RData",
  package = "sigminer", mustWork = TRUE
))

show_cn_freq_circos(cn)
ss <- unique(cn@data$sample)
show_cn_freq_circos(cn, groups = list(a = ss[1:5], b = ss[6:10]), cols = c("red", "green"))
```

---

show\_cn\_group\_profile *Show Summary Copy Number Profile for Sample Groups*

---

**Description**

Show Summary Copy Number Profile for Sample Groups

**Usage**

```
show_cn_group_profile(
  data,
  groups = NULL,
  fill_area = TRUE,
  cols = NULL,
  chrs = paste0("chr", c(1:22, "X")),
  genome_build = c("hg19", "hg38", "mm10", "mm9"),
  cutoff = 2L,
  resolution_factor = 1L,
  force_y_limit = TRUE,
  highlight_genes = NULL,
  repel = FALSE,
  nrow = NULL,
  ncol = NULL,
  return_plotlist = FALSE
)
```

**Arguments**

data	a CopyNumber object or a data.frame containing at least 'chromosome', 'start', 'end', 'segVal', 'sample' these columns.
groups	a named list or a column name for specifying groups.
fill_area	default is TRUE, fill area with colors.

cols	length-2 colors for AMP and DEL.
chrs	chromosomes start with 'chr'.
genome_build	genome build version, used when data is a data.frame, should be 'hg19' or 'hg38'.
cutoff	copy number value cutoff for splitting data into AMP and DEL. The values equal to cutoff are discarded. Default is 2, you can also set a length-2 vector, e.g. c(2, 2).
resolution_factor	an integer to control the resolution. When it is 1 (default), compute frequency in each cytoband. When it is 2, use compute frequency in each half cytoband.
force_y_limit	default is TRUE, force multiple plots
highlight_genes	gene list to highlight. have same y ranges. You can also set a length-2 numeric value.
repel	if TRUE (default is FALSE), repel highlight genes to avoid overlap.
nrow	number of rows in the plot grid when multiple samples are selected.
ncol	number of columns in the plot grid when multiple samples are selected.
return_plotlist	default is FALSE, if TRUE, return a plot list instead of a combined plot.

### Value

a (list of) ggplot object.

### Examples

```
load(system.file("extdata", "toy_copynumber.RData",
  package = "sigminer", mustWork = TRUE
))

p1 <- show_cn_group_profile(cn)
p1

ss <- unique(cn@data$sample)
p2 <- show_cn_group_profile(cn, groups = list(a = ss[1:5], b = ss[6:10]))
p2
p3 <- show_cn_group_profile(cn,
  groups = list(g1 = ss[1:5], g2 = ss[6:10]),
  force_y_limit = c(-1, 1), nrow = 2
)
p3

## Set custom cutoff for custom data
data <- cn@data
data$segVal <- data$segVal - 2L
p4 <- show_cn_group_profile(data,
  groups = list(g1 = ss[1:5], g2 = ss[6:10]),
  force_y_limit = c(-1, 1), nrow = 2,
```

```

    cutoff = c(0, 0)
  )
  p4

  ## Add highlight gene
  p5 <- show_cn_group_profile(cn, highlight_genes = c("TP53", "EGFR"))
  p5

```

---

show\_cn\_profile

*Show Sample Copy Number Profile*


---

## Description

Sometimes it is very useful to check details about copy number profile for one or multiple samples. This function is designed to do this job and can be further modified by **ggplot2** related packages.

## Usage

```

show_cn_profile(
  data,
  samples = NULL,
  show_n = NULL,
  show_title = FALSE,
  show_labels = NULL,
  chrs = paste0("chr", 1:22),
  position = NULL,
  genome_build = c("hg19", "hg38", "mm10", "mm9"),
  ylim = NULL,
  nrow = NULL,
  ncol = NULL,
  return_plotlist = FALSE
)

```

## Arguments

data	a <a href="#">CopyNumber</a> object or a data.frame containing at least 'chromosome', 'start', 'end', 'segVal' these columns.
samples	default is NULL, can be a character vector representing multiple samples. If data argument is a data.frame, a column called sample must exist.
show_n	number of samples to show, this is used for checking.
show_title	if TRUE, show title for multiple samples.
show_labels	one of NULL, "s" (for labelling short segments < 1e7) or "a" (all segments).
chrs	chromosomes start with 'chr'.
position	a position range, e.g. "chr1:3218923-116319008". Only data overlaps with this range will be shown.

genome\_build genome build version, used when data is a data.frame, should be 'hg19' or 'hg38'.

ylim limites for y axis.

nrow number of rows in the plot grid when multiple samples are selected.

ncol number of columns in the plot grid when multiple samples are selected.

return\_plotlist default is FALSE, if TRUE, return a plot list instead of a combined plot.

**Value**

a ggplot object or a list

**Examples**

```
# Load copy number object
load(system.file("extdata", "toy_copynumber.RData",
  package = "sigminer", mustWork = TRUE
))

p <- show_cn_profile(cn, nrow = 2, ncol = 1)
p

p2 <- show_cn_profile(cn,
  nrow = 2, ncol = 1,
  position = "chr1:3218923-116319008"
)
p2
```

---

 show\_cor

*A Simple and General Way for Association Analysis*


---

**Description**

All variables must be continuous. The matrix will be returned as an element of ggplot object. This is basically a wrapper of R package [ggcorrplot](#).

**Usage**

```
show_cor(
  data,
  x_vars = colnames(data),
  y_vars = x_vars,
  cor_method = "spearman",
  vis_method = "square",
  lab = TRUE,
  test = TRUE,
```

```

    hc_order = FALSE,
    p_adj = NULL,
    ...
  )

```

### Arguments

data	a data.frame.
x_vars	variables/column names shown in x axis.
y_vars	variables/column names shown in y axis.
cor_method	method for correlation, default is 'spearman'.
vis_method	visualization method, default is 'square', can also be 'circle'.
lab	logical value. If TRUE, add correlation coefficient on the plot.
test	if TRUE, run test for correlation and mark significance.
hc_order	logical value. If TRUE, correlation matrix will be hc.ordered using hclust function.
p_adj	p adjust method, see <a href="#">stats::p.adjust</a> for details.
...	other parameters passing to <code>ggcorrplot::ggcorrplot()</code> .

### Value

a ggplot object

### See Also

[show\\_sig\\_feature\\_corrplot](#) for specific and more powerful association analysis and visualization.

### Examples

```

data("mtcars")
p1 <- show_cor(mtcars)
p2 <- show_cor(mtcars,
  x_vars = colnames(mtcars)[1:4],
  y_vars = colnames(mtcars)[5:8]
)
p3 <- show_cor(mtcars, vis_method = "circle", p_adj = "fdr")
p1
p1$cor
p2
p3

## Auto detect problem variables
mtcars$xx <- 0L
p4 <- show_cor(mtcars)
p4

```

---

`show_cosmic`*Show Signature Information in Web Browser*

---

**Description**

Show Signature Information in Web Browser

**Usage**

```
show_cosmic(x = "home")
```

**Arguments**

x a string indicating location ("home" for COSMIC signature home, "legacy" for COSMIC v2 signatures, "SBS" for COSMIC v3 SBS signatures, "DBS" for COSMIC v3 DBS signatures, "ID" for COSMIC v3 INDEL signatures) or signature index (e.g. "SBS1", "DBS2", "ID3").

**Value**

Nothing.

**Examples**

```
## Not run:
show_cosmic()
show_cosmic("legacy")
show_cosmic("SBS")
show_cosmic("DBS")
show_cosmic("ID")
show_cosmic("SBS1")
show_cosmic("DBS2")
show_cosmic("ID3")

## End(Not run)
```

---

`show_cosmic_sig_profile`*Plot Reference (Mainly COSMIC) Signature Profile*

---

**Description**

Plot Reference (Mainly COSMIC) Signature Profile



**Usage**

```
show_cosmic_sig_profile(
  sig_index = NULL,
  show_index = TRUE,
  sig_db = "legacy",
  ...
)
```

**Arguments**

sig_index	a vector for signature index. "ALL" for all signatures.
show_index	if TRUE, show valid indices.
sig_db	default 'legacy', it can be 'legacy' (for <b>COSMIC v2 'SBS'</b> ), 'SBS', 'DBS', 'ID' and 'TSB' (for <b>COSMIV v3.1 signatures</b> ) for small scale mutations. For more specific details, it can also be 'SBS_hg19', 'SBS_hg38', 'SBS_mm9', 'SBS_mm10', 'DBS_hg19', 'DBS_hg38', 'DBS_mm9', 'DBS_mm10' to use COSMIC v3 reference signatures from Alexandrov, Ludmil B., et al. (2020) (reference #1). In addition, it can be one of "SBS_Nik_lab_Organ", "RS_Nik_lab_Organ", "SBS_Nik_lab", "RS_Nik_lab" to refer reference signatures from Degasperi, Andrea, et al. (2020) (reference #2); "RS_BRCA560", "RS_USARC" to reference signatures from BRCA560 and USARC cohorts; "CNS_USARC" (40 categories), "CNS_TCGA" (48 categories) to reference copy number signatures from USARC cohort and TCGA. <b>UPDATE</b> , the latest version of reference version can be automatically downloaded and loaded from <a href="https://cancer.sanger.ac.uk/signatures/downloads/">https://cancer.sanger.ac.uk/signatures/downloads/</a> when a option with latest_ prefix is specified (e.g. "latest_SBS_GRCh37"). <b>Note</b> : the signature profile for different genome builds are basically same. And specific database (e.g. 'SBS_mm10') contains less signatures than all COSMIC signatures (because some signatures are not detected from Alexandrov, Ludmil B., et al. (2020)). For all available options, check the parameter setting.
...	other arguments passing to <a href="#">show_sig_profile</a> .

**Value**

a ggplot object

**Author(s)**

Shixiang Wang [w\\_shixiang@163.com](mailto:w_shixiang@163.com)

**Examples**

```
show_cosmic_sig_profile()
show_cosmic_sig_profile(sig_db = "SBS")
show_cosmic_sig_profile(sig_index = 1:5)
show_cosmic_sig_profile(sig_db = "SBS", sig_index = c("10a", "17a"))

gg <- show_cosmic_sig_profile(sig_index = 1:5)
gg$aetiology
```

---

show_groups	<i>Show Signature Contribution in Clusters</i>
-------------	--

---

### Description

See example section in [sig\\_fit\(\)](#) for an examples.

### Usage

```
show_groups(grp_dt, ...)
```

### Arguments

grp_dt	a result data.table from <a href="#">get_groups</a> .
...	parameters passing to <a href="#">legend()</a> , e.g. x = "topleft".

### Value

nothing.

### See Also

[get\\_groups](#), [sig\\_fit](#).

---

show_group_comparison	<i>Plot Group Comparison Result</i>
-----------------------	-------------------------------------

---

### Description

Using result data from [get\\_group\\_comparison](#), this function plots genotypes/phenotypes comparison between signature groups using **ggplot2** package and return a list of ggplot object contains individual and combined plots. The combined plot is easily saved to local using [cowplot::save\\_plot\(\)](#). Of note, default fisher test p values are shown for categorical data and fdr values are shown for continuous data.

### Usage

```
show_group_comparison(
  group_comparison,
  xlab = "group",
  ylab_co = NA,
  legend_title_ca = NA,
  legend_position_ca = "bottom",
  set_ca_sig_yaxis = FALSE,
  set_ca_custom_xlab = FALSE,
```

```

    show_pvalue = TRUE,
    ca_p_threshold = 0.01,
    method = "wilcox.test",
    p.adjust.method = "fdr",
    base_size = 12,
    font_size_x = 12,
    text_angle_x = 30,
    text_hjust_x = 0.2,
    ...
)

```

## Arguments

<code>group_comparison</code>	a list from result of <a href="#">get_group_comparison</a> function.
<code>xlab</code>	lab name of x axis for all plots. if it is NA, remove title for x axis.
<code>ylab_co</code>	lab name of y axis for plots of continuous type data. Of note, this argument should be a character vector has same length as <code>group_comparison</code> , the location for categorical type data should mark with NA.
<code>legend_title_ca</code>	legend title for plots of categorical type data.
<code>legend_position_ca</code>	legend position for plots of categorical type data. Of note, this argument should be a character vector has same length as <code>group_comparison</code> , the location for continuous type data should mark with NA.
<code>set_ca_sig_yaxis</code>	if TRUE, use y axis to show signature proportion instead of variable proportion.
<code>set_ca_custom_xlab</code>	only works when <code>set_ca_sig_yaxis</code> is TRUE. If TRUE, set x labels using input <code>xlab</code> , otherwise variable names will be used.
<code>show_pvalue</code>	if TRUE, show p values.
<code>ca_p_threshold</code>	a p threshold for categorical variables, default is 0.01. A p value less than 0.01 will be shown as $P < 0.01$ .
<code>method</code>	a character string indicating which method to be used for comparing means. It can be 't.test', 'wilcox.test' etc..
<code>p.adjust.method</code>	correction method, default is 'fdr'. Run <code>p.adjust.methods</code> to see all available options.
<code>base_size</code>	overall font size.
<code>font_size_x</code>	font size for x.
<code>text_angle_x</code>	text angle for x.
<code>text_hjust_x</code>	adjust x axis text
<code>...</code>	other paramters pass to <code>ggpubr::compare_means()</code> or <code>ggpubr::stat_compare_means()</code> according to the specified method.

**Value**

a list of ggplot objects.

**Author(s)**

Shixiang Wang [w\\_shixiang@163.com](mailto:w_shixiang@163.com)

**Examples**

```
load(system.file("extdata", "toy_copynumber_signature_by_W.RData",
  package = "sigminer", mustWork = TRUE
))

# Assign samples to clusters
groups <- get_groups(sig, method = "k-means")

set.seed(1234)

groups$prob <- rnorm(10)
groups$new_group <- sample(c("1", "2", "3", "4", NA), size = nrow(groups), replace = TRUE)

# Compare groups (filter NAs for categorical columns)
groups.cmp <- get_group_comparison(groups[, -1],
  col_group = "group",
  cols_to_compare = c("prob", "new_group"),
  type = c("co", "ca"), verbose = TRUE
)

# Compare groups (Set NAs of categorical columns to 'Rest')
groups.cmp2 <- get_group_comparison(groups[, -1],
  col_group = "group",
  cols_to_compare = c("prob", "new_group"),
  type = c("co", "ca"), NAs = "Rest", verbose = TRUE
)

show_group_comparison(groups.cmp)

ggcomp <- show_group_comparison(groups.cmp2)
ggcomp$co_comb
ggcomp$ca_comb
```

---

show\_group\_distribution

*Show Grouped Variable Distribution*

---

**Description**

This is a general function, it can be used in any proper analysis.

**Usage**

```

show_group_distribution(
  data,
  gvar,
  dvar,
  fun = stats::median,
  order_by_fun = FALSE,
  alpha = 0.8,
  g_label = "label",
  g_angle = 0,
  g_position = "top",
  point_size = 1L,
  segment_size = 1L,
  segment_color = "red",
  xlab = NULL,
  ylab = NULL,
  nrow = 1L,
  background_color = c("#DCDCDC", "#F5F5F5")
)

```

**Arguments**

data	a data.frame.
gvar	a group variable name/index.
dvar	a distribution variable name/index.
fun	a function to summarize, default is <a href="#">stats::median</a> , can also be <a href="#">mean</a> .
order_by_fun	if TRUE, reorder the groups by summary measure computed by argument fun.
alpha	alpha for points, range from 0 to 1.
g_label	a string 'label' (default) for labeling with sample size, or 'norm' to show just group name, or a named vector to set facet labels.
g_angle	angle for facet labels, default is 0.
g_position	position for facet labels, default is 'top', can also be 'bottom'.
point_size	size of point.
segment_size	size of segment.
segment_color	color of segment.
xlab	title for x axis.
ylab	title for y axis.
nrow	number of row.
background_color	background color for plot panel.

**Value**

a ggplot object.

**Author(s)**

Shixiang Wang [w\\_shixiang@163.com](mailto:w_shixiang@163.com)

**Examples**

```
set.seed(1234)
data <- data.frame(
  yval = rnorm(120),
  gr = c(rep("A", 50), rep("B", 40), rep("C", 30))
)
p <- show_group_distribution(data,
  gvar = 2, dvar = 1,
  g_label = "norm",
  background_color = "grey"
)
p
p2 <- show_group_distribution(data,
  gvar = "gr", dvar = "yval",
  g_position = "bottom",
  order_by_fun = TRUE,
  alpha = 0.3
)
p2

# Set custom group names
p3 <- show_group_distribution(data,
  gvar = 2, dvar = 1,
  g_label = c("A" = "X", "B" = "Y", "C" = "Z")
)
p3
```

---

show\_group\_enrichment *Show Group Enrichment Result*

---

**Description**

See [group\\_enrichment](#) for examples. NOTE the box fill and the box text have different meanings.

**Usage**

```
show_group_enrichment(
  df_enrich,
  return_list = FALSE,
  scales = "free",
  add_text_annotation = TRUE,
  fill_by_p_value = TRUE,
  use_fdr = TRUE,
  cut_p_value = FALSE,
  cut_breaks = c(-Inf, -5, log10(0.05), -log10(0.05), 5, Inf),
```

```

  cut_labels = c("↓ 1e-5", "↓ 0.05", "non-significant", "↑ 0.05", "↑ 1e-5"),
  fill_scale = scale_fill_gradient2(low = "#08A76B", mid = "white", high = "red",
    midpoint = ifelse(fill_by_p_value, 0, 1)),
  cluster_row = FALSE,
  ...
)

```

### Arguments

<code>df_enrich</code>	result data.frame from <a href="#">group_enrichment</a> .
<code>return_list</code>	if TRUE, return a list of ggplot object so user can combine multiple plots by other R packages like patchwork.
<code>scales</code>	Should scales be fixed ("fixed", the default), free ("free"), or free in one dimension ("free_x", "free_y")?
<code>add_text_annotation</code>	if TRUE, add text annotation in box. When show p value with filled color, the text indicates relative change; when show relative change with filled color, the text indicates p value.
<code>fill_by_p_value</code>	if TRUE, show log10 based p values with filled color. The +/- of p values indicates change direction. If p vlaues is mapped to fill, then text shows effect size, and vice versa.
<code>use_fdr</code>	if TRUE, show FDR values instead of raw p-values.
<code>cut_p_value</code>	if TRUE, cut p values into 5 regions for better visualization. Only works when <code>fill_by_p_value = TRUE</code> .
<code>cut_breaks</code>	when <code>cut_p_value</code> is TRUE, this option set the (log10 based) breaks.
<code>cut_labels</code>	when <code>cut_p_value</code> is TRUE, this option set the labels.
<code>fill_scale</code>	a Scale object generated by ggplot2 package to set color for continuous values.
<code>cluster_row</code>	if TRUE, cluster rows with Hierarchical Clustering ('complete' method).
<code>...</code>	other parameters passing to <a href="#">ggplot2::facet_wrap</a> , only used when <code>return_list</code> is FALSE.

### Value

a (list of) ggplot object.

---

show\_group\_mapping      *Map Groups using Sankey*

---

### Description

This feature is designed for signature analysis. However, users can also use it in other similar situations.

**Usage**

```
show_group_mapping(  
  data,  
  col_to_flow,  
  cols_to_map,  
  include_sig = FALSE,  
  fill_na = FALSE,  
  title = NULL,  
  xlab = NULL,  
  ylab = NULL,  
  custom_theme = cowplot::theme_minimal_hgrid()  
)
```

**Arguments**

<code>data</code>	a data.frame containing signature group and other categorical groups.
<code>col_to_flow</code>	length-1 character showing the column to flow, typically a signature group.
<code>cols_to_map</code>	character vector showing colnames of other groups.
<code>include_sig</code>	default if FALSE, if TRUE, showing signature group.
<code>fill_na</code>	length-1 string to fill NA, default is FALSE.
<code>title</code>	the title.
<code>xlab</code>	label for x axis.
<code>ylab</code>	label for y axis.
<code>custom_theme</code>	theme for plotting, default is <code>cowplot::theme_minimal_hgrid()</code> .

**Value**

a ggplot object

**Examples**

```
data <- dplyr::tibble(  
  Group1 = rep(LETTERS[1:5], each = 10),  
  Group2 = rep(LETTERS[6:15], each = 5),  
  zzzz = c(rep("xx", 20), rep("yy", 20), rep(NA, 10))  
)  
p1 <- show_group_mapping(data, col_to_flow = "Group1", cols_to_map = colnames(data)[-1])  
p1  
  
p2 <- show_group_mapping(data,  
  col_to_flow = "Group1", cols_to_map = colnames(data)[-1],  
  include_sig = TRUE  
)  
p2
```



---

show\_sig\_bootstrap      *Show Signature Bootstrap Analysis Results*

---

### Description

See details for description.

### Usage

```
show_sig_bootstrap_exposure(  
  bt_result,  
  sample = NULL,  
  signatures = NULL,  
  methods = "QP",  
  plot_fun = c("boxplot", "violin"),  
  agg_fun = c("mean", "median", "min", "max"),  
  highlight = "auto",  
  highlight_size = 4,  
  palette = "aaas",  
  title = NULL,  
  xlab = FALSE,  
  ylab = "Signature exposure",  
  width = 0.3,  
  dodge_width = 0.8,  
  outlier.shape = NA,  
  add = "jitter",  
  add.params = list(alpha = 0.3),  
  ...  
)
```

```
show_sig_bootstrap_error(  
  bt_result,  
  sample = NULL,  
  methods = "QP",  
  plot_fun = c("boxplot", "violin"),  
  agg_fun = c("mean", "median"),  
  highlight = "auto",  
  highlight_size = 4,  
  palette = "aaas",  
  title = NULL,  
  xlab = FALSE,  
  ylab = "Reconstruction error (L2 norm)",  
  width = 0.3,  
  dodge_width = 0.8,  
  outlier.shape = NA,  
  add = "jitter",  
  add.params = list(alpha = 0.3),
```

```

    legend = "none",
    ...
)

show_sig_bootstrap_stability(
  bt_result,
  signatures = NULL,
  measure = c("RMSE", "CV", "MAE", "AbsDiff"),
  methods = "QP",
  plot_fun = c("boxplot", "violin"),
  palette = "aaas",
  title = NULL,
  xlab = FALSE,
  ylab = "Signature instability",
  width = 0.3,
  outlier.shape = NA,
  add = "jitter",
  add.params = list(alpha = 0.3),
  ...
)

```

### Arguments

<code>bt_result</code>	result object from <a href="#">sig_fit_bootstrap_batch</a> .
<code>sample</code>	a sample id.
<code>signatures</code>	signatures to show.
<code>methods</code>	a subset of <code>c("NLS", "QP", "SA")</code> .
<code>plot_fun</code>	set the plot function.
<code>agg_fun</code>	set the aggregation function when <code>sample</code> is <code>NULL</code> .
<code>highlight</code>	set the color for optimal solution. Default is "auto", which use the same color as bootstrap results, you can set it to color like "red", "gold", etc.
<code>highlight_size</code>	size for highlighting triangle, default is 4.
<code>palette</code>	the color palette to be used for coloring or filling by groups. Allowed values include "grey" for grey color palettes; brewer palettes e.g. "RdBu", "Blues", ...; or custom color palette e.g. <code>c("blue", "red")</code> ; and scientific journal palettes from ggsci R package, e.g.: "npg", "aaas", "lancet", "jco", "ucscgb", "uchicago", "simpsons" and "rickandmorty".
<code>title</code>	plot main title.
<code>xlab</code>	character vector specifying x axis labels. Use <code>xlab = FALSE</code> to hide xlab.
<code>ylab</code>	character vector specifying y axis labels. Use <code>ylab = FALSE</code> to hide ylab.
<code>width</code>	numeric value between 0 and 1 specifying box width.
<code>dodge_width</code>	dodge width.
<code>outlier.shape</code>	point shape of outlier. Default is 19. To hide outlier, specify <code>outlier.shape = NA</code> . When jitter is added, then outliers will be automatically hidden.

add	character vector for adding another plot element (e.g.: dot plot or error bars). Allowed values are one or the combination of: "none", "dotplot", "jitter", "boxplot", "point", "mean", "mean_se", "mean_sd", "mean_ci", "mean_range", "median", "median_iqr", "median_hilow", "median_q1q3", "median_mad", "median_range"; see ?desc_statby for more details.
add.params	parameters (color, shape, size, fill, linetype) for the argument 'add'; e.g.: add.params = list(color = "red").
...	other parameters passing to <code>ggpubr::ggboxplot</code> or <code>ggpubr::ggviolin</code> .
legend	character specifying legend position. Allowed values are one of c("top", "bottom", "left", "right", "none"). To remove the legend use legend = "none". Legend position can be also specified using a numeric vector c(x, y); see details section.
measure	measure to estimate the exposure instability, can be one of 'RMSE', 'CV', 'MAE' and 'AbsDiff'.

## Details

### Functions:

- [show\\_sig\\_bootstrap\\_exposure](#) - this function plots exposures from bootstrap samples with both dotted boxplot. The optimal exposure (the exposure from original input) is shown as triangle point. **Only one sample can be plotted.**
- [show\\_sig\\_bootstrap\\_error](#) - this function plots decomposition errors from bootstrap samples with both dotted boxplot. The error from optimal solution (the decomposition error from original input) is shown as triangle point. **Only one sample can be plotted.**
- [show\\_sig\\_bootstrap\\_stability](#) - this function plots the signature exposure instability for specified signatures. Currently, the instability measure supports 3 types:
  - 'RMSE' for Mean Root Squared Error (default) of bootstrap exposures and original exposures for each sample.
  - 'CV' for Coefficient of Variation (CV) based on RMSE (i.e.  $RMSE / btExposure\_mean$ ).
  - 'MAE' for Mean Absolute Error of bootstrap exposures and original exposures for each sample.
  - 'AbsDiff' for Absolute Difference between mean bootstram exposure and original exposure.

## Value

a ggplot object

## References

Huang X, Wojtowicz D, Przytycka TM. Detecting presence of mutational signatures in cancer with confidence. *Bioinformatics*. 2018;34(2):330–337. doi:10.1093/bioinformatics/btx604

## See Also

[sig\\_fit\\_bootstrap\\_batch](#), [sig\\_fit](#), [sig\\_fit\\_bootstrap](#)

**Examples**

```

if (require("BSgenome.Hsapiens.UCSC.hg19")) {
  lam1.maf <- system.file("extdata", "tcga_lam1.maf.gz", package = "maftools")
  lam1 <- read_maf(maf = lam1.maf)
  mt_tally <- sig_tally(
    lam1,
    ref_genome = "BSgenome.Hsapiens.UCSC.hg19",
    use_syn = TRUE
  )

  library(NMF)
  mt_sig <- sig_extract(mt_tally$nmf_matrix,
    n_sig = 3,
    nrun = 2,
    cores = 1
  )

  mat <- t(mt_tally$nmf_matrix)
  mat <- mat[, colSums(mat) > 0]
  bt_result <- sig_fit_bootstrap_batch(mat, sig = mt_sig, n = 10)
  ## Parallel computation
  ## bt_result = sig_fit_bootstrap_batch(mat, sig = mt_sig, n = 10, use_parallel = TRUE)

  ## At default, mean bootstrap exposure for each sample has been calculated
  p <- show_sig_bootstrap_exposure(bt_result, methods = c("QP"))
  ## Show bootstrap exposure (optimal exposure is shown as triangle)
  p1 <- show_sig_bootstrap_exposure(bt_result, methods = c("QP"), sample = "TCGA-AB-2802")
  p1
  p2 <- show_sig_bootstrap_exposure(bt_result,
    methods = c("QP"),
    sample = "TCGA-AB-3012",
    signatures = c("Sig1", "Sig2")
  )
  p2

  ## Show bootstrap error
  ## Similar to exposure above
  p <- show_sig_bootstrap_error(bt_result, methods = c("QP"))
  p
  p3 <- show_sig_bootstrap_error(bt_result, methods = c("QP"), sample = "TCGA-AB-2802")
  p3

  ## Show exposure (in)stability
  p4 <- show_sig_bootstrap_stability(bt_result, methods = c("QP"))
  p4
  p5 <- show_sig_bootstrap_stability(bt_result, methods = c("QP"), measure = "MAE")
  p5
  p6 <- show_sig_bootstrap_stability(bt_result, methods = c("QP"), measure = "AbsDiff")
  p6
  p7 <- show_sig_bootstrap_stability(bt_result, methods = c("QP"), measure = "CV")
  p7

```

```

} else {
  message("Please install package 'BSgenome.Hsapiens.UCSC.hg19' firstly!")
}

```

---

show\_sig\_consensusmap *Show Signature Consensus Map*

---

## Description

This function is a wrapper of `NMF::consensusmap()`.

## Usage

```

show_sig_consensusmap(
  sig,
  main = "Consensus matrix",
  tracks = c("consensus:", "silhouette:"),
  lab_row = NA,
  lab_col = NA,
  ...
)

```

## Arguments

<code>sig</code>	a Signature object obtained from <a href="#">sig_extract</a> .
<code>main</code>	Main title as a character string or a grob.
<code>tracks</code>	Special additional annotation tracks to highlight associations between basis components and sample clusters:  <b>basis</b> matches each row (resp. column) to the most contributing basis component in <code>basismap</code> (resp. <code>coefmap</code> ). In <code>basismap</code> (resp. <code>coefmap</code> ), adding a track <code>:basis</code> to <code>annCol</code> (resp. <code>annRow</code> ) makes the column (resp. row) corresponding to the component being also highlighted using the matching colours.
<code>lab_row</code>	labels for the rows.
<code>lab_col</code>	labels for the columns.
<code>...</code>	other parameters passing to <code>NMF::consensusmap()</code> .

## Value

nothing

---

show\_sig\_exposure      *Plot Signature Exposure*

---

## Description

Currently support copy number signatures and mutational signatures.

## Usage

```
show_sig_exposure(
  Signature,
  sig_names = NULL,
  groups = NULL,
  grp_order = NULL,
  grp_size = NULL,
  cutoff = NULL,
  style = c("default", "cosmic"),
  palette = use_color_style(style),
  base_size = 12,
  font_scale = 1,
  rm_space = FALSE,
  rm_grid_line = TRUE,
  rm_panel_border = FALSE,
  hide_samps = TRUE,
  legend_position = "top"
)
```

## Arguments

Signature	a Signature object obtained either from <a href="#">sig_extract</a> or <a href="#">sig_auto_extract</a> , or just a raw <b>absolute</b> exposure matrix with column representing samples (patients) and row representing signatures (signature names must end with different digital numbers, e.g. Sig1, Sig10, x12). If you named signatures with letters, you can specify them by sig_names parameter.
sig_names	set name of signatures, can be a character vector.
groups	sample groups, default is NULL.
grp_order	order of groups, default is NULL.
grp_size	font size of groups.
cutoff	a cutoff value to remove hyper-mutated samples.
style	plot style, one of 'default' and 'cosmic', works when parameter set_gradient_color is FALSE.
palette	palette used to plot, default use a built-in palette according to parameter style.
base_size	overall font size.
font_scale	a number used to set font scale.

rm\_space            default is FALSE. If TRUE, it will remove border color and expand the bar width to 1. This is useful when the sample size is big.

rm\_grid\_line        default is FALSE, if TRUE, remove grid lines of plot.

rm\_panel\_border    default is TRUE for style 'cosmic', remove panel border to keep plot tight.

hide\_samps         if TRUE, hide sample names.

legend\_position    position of legend, default is 'top'.

**Value**

a ggplot object

**Author(s)**

Shixiang Wang

**Examples**

```
# Load mutational signature
load(system.file("extdata", "toy_mutational_signature.RData",
  package = "sigminer", mustWork = TRUE
))
# Show signature exposure
p1 <- show_sig_exposure(sig2)
p1

# Load copy number signature
load(system.file("extdata", "toy_copynumber_signature_by_W.RData",
  package = "sigminer", mustWork = TRUE
))
# Show signature exposure
p2 <- show_sig_exposure(sig)
p2
```

---

show\_sig\_feature\_corrplot

*Draw Corrplot for Signature Exposures and Other Features*

---

**Description**

This function is for association visualization. Of note, the parameters `p_val` and `drop` will affect the visualization of association results under p value threshold.

**Usage**

```

show_sig_feature_corrplot(
  tidy_cor,
  feature_list,
  sort_features = FALSE,
  sig_orders = NULL,
  drop = TRUE,
  return_plotlist = FALSE,
  p_val = 0.05,
  xlab = "Signatures",
  ylab = "Features",
  co_gradient_colors = scale_color_gradient2(low = "blue", mid = "white", high = "red",
  midpoint = 0),
  ca_gradient_colors = co_gradient_colors,
  plot_ratio = "auto",
  breaks_count = NULL
)

```

**Arguments**

<code>tidy_cor</code>	data returned by <a href="#">get_tidy_association</a> .
<code>feature_list</code>	a character vector contains features want to be plotted. If missing, all features will be used.
<code>sort_features</code>	default is FALSE, use feature order obtained from the previous step. If TRUE, sort features as <code>feature_list</code> .
<code>sig_orders</code>	signature levels for ordering.
<code>drop</code>	if TRUE, when a feature has no association with all signatures (p value larger than threshold set by <code>p_val</code> ), this feature will be removed from the plot. Otherwise, this feature (a row) will keep with all blank white.
<code>return_plotlist</code>	if TRUE, return as a list of ggplot objects.
<code>p_val</code>	p value threshold. If p value larger than this threshold, the result becomes blank white.
<code>xlab</code>	label for x axis.
<code>ylab</code>	label for y axis.
<code>co_gradient_colors</code>	a Scale object representing gradient colors used to plot for continuous features.
<code>ca_gradient_colors</code>	a Scale object representing gradient colors used to plot for categorical features.
<code>plot_ratio</code>	a length-2 numeric vector to set the height/width ratio.
<code>breaks_count</code>	breaks for sample count. If set it to NULL, ggplot bin scale will be used to automatically determine the breaks. If set it to NA, aes for sample will be not used.



**Value**

a ggplot2 object

**See Also**

[get\\_tidy\\_association](#) and [get\\_sig\\_feature\\_association](#)

**Examples**

```
# The data is generated from Wang, Shixiang et al.
load(system.file("extdata", "asso_data.RData",
  package = "sigminer", mustWork = TRUE
))

p <- show_sig_feature_corrplot(
  tidy_data.seqz.feature,
  p_val = 0.05,
  breaks_count = c(0L,200L, 400L, 600L, 800L, 1020L))
p
```

---

show\_sig\_fit

*Show Signature Fit Result*

---

**Description**

See [sig\\_fit](#) for examples.

**Usage**

```
show_sig_fit(
  fit_result,
  samples = NULL,
  signatures = NULL,
  plot_fun = c("boxplot", "violin", "scatter"),
  palette = "aaas",
  title = NULL,
  xlab = FALSE,
  ylab = "Signature exposure",
  legend = "none",
  width = 0.3,
  outlier.shape = NA,
  add = "jitter",
  add.params = list(alpha = 0.3),
  ...
)
```

**Arguments**

<code>fit_result</code>	result object from <a href="#">sig_fit</a> .
<code>samples</code>	samples to show, if NULL, all samples are used.
<code>signatures</code>	signatures to show.
<code>plot_fun</code>	set the plot function.
<code>palette</code>	the color palette to be used for coloring or filling by groups. Allowed values include "grey" for grey color palettes; brewer palettes e.g. "RdBu", "Blues", ...; or custom color palette e.g. <code>c("blue", "red")</code> ; and scientific journal palettes from ggsci R package, e.g.: "npg", "aaas", "lancet", "jco", "ucscgb", "uchicago", "simpsons" and "rickandmorty".
<code>title</code>	plot main title.
<code>xlab</code>	character vector specifying x axis labels. Use <code>xlab = FALSE</code> to hide xlab.
<code>ylab</code>	character vector specifying y axis labels. Use <code>ylab = FALSE</code> to hide ylab.
<code>legend</code>	character specifying legend position. Allowed values are one of <code>c("top", "bottom", "left", "right", "none")</code> . To remove the legend use <code>legend = "none"</code> . Legend position can be also specified using a numeric vector <code>c(x, y)</code> ; see details section.
<code>width</code>	numeric value between 0 and 1 specifying box width.
<code>outlier.shape</code>	point shape of outlier. Default is 19. To hide outlier, specify <code>outlier.shape = NA</code> . When jitter is added, then outliers will be automatically hidden.
<code>add</code>	character vector for adding another plot element (e.g.: dot plot or error bars). Allowed values are one or the combination of: "none", "dotplot", "jitter", "boxplot", "point", "mean", "mean_se", "mean_sd", "mean_ci", "mean_range", "median", "median_iqr", "median_hilow", "median_q1q3", "median_mad", "median_range"; see <code>?desc_statby</code> for more details.
<code>add.params</code>	parameters (color, shape, size, fill, linetype) for the argument 'add'; e.g.: <code>add.params = list(color = "red")</code> .
<code>...</code>	other arguments to be passed to <a href="#">geom_boxplot</a> , <a href="#">ggpar</a> and <a href="#">facet</a> .

**Value**

a ggplot object.

**See Also**

[sig\\_fit](#), [show\\_sig\\_bootstrap\\_exposure](#), [sig\\_fit\\_bootstrap](#), [sig\\_fit\\_bootstrap\\_batch](#)

---

show_sig_profile	<i>Show Signature Profile</i>
------------------	-------------------------------

---

## Description

Who don't like to show a barplot for signature profile? This is for it.

## Usage

```
show_sig_profile(
  Signature,
  mode = c("SBS", "copynumber", "DBS", "ID", "RS"),
  method = "Wang",
  by_context = FALSE,
  normalize = c("row", "column", "raw", "feature"),
  y_tr = NULL,
  filters = NULL,
  feature_setting = sigminer::CN.features,
  style = c("default", "cosmic"),
  palette = use_color_style(style, ifelse(by_context, "SBS", mode), method),
  set_gradient_color = FALSE,
  free_space = "free_x",
  rm_panel_border = style == "cosmic",
  rm_grid_line = style == "cosmic",
  rm_axis_text = FALSE,
  bar_border_color = ifelse(style == "default", "grey50", "white"),
  bar_width = 0.7,
  paint_axis_text = TRUE,
  x_label_angle = ifelse(mode == "copynumber" & !(startsWith(method, "T") | method ==
    "X"), 60, 90),
  x_label_vjust = ifelse(mode == "copynumber" & !(startsWith(method, "T") | method ==
    "X"), 1, 0.5),
  x_label_hjust = 1,
  x_lab = "Components",
  y_lab = "auto",
  y_limits = NULL,
  params = NULL,
  show_cv = FALSE,
  params_label_size = 3,
  params_label_angle = 60,
  y_expand = 1,
  digits = 2,
  base_size = 12,
  font_scale = 1,
  sig_names = NULL,
  sig_orders = NULL,
  check_sig_names = TRUE
```

)

**Arguments**

Signature	a Signature object obtained either from <code>sig_extract</code> or <code>sig_auto_extract</code> , or just a raw signature matrix with row representing components (motifs) and column representing signatures (column names must start with 'Sig').
mode	signature type for plotting, now supports 'copynumber', 'SBS', 'DBS', 'ID' and 'RS' (genome rearrangement signature).
method	method for copy number feature classification in <code>sig_tally</code> , can be one of "Wang" ("W"), "S".
by_context	for specific use.
normalize	one of 'row', 'column', 'raw' and "feature", for row normalization (signature), column normalization (component), raw data, row normalization by feature, respectively. Of note, 'feature' only works when the mode is 'copynumber'.
y_tr	a function (e.g. $\log_{10}$ ) to transform y axis before plotting.
filters	a pattern used to select components to plot.
feature_setting	a data.frame used for classification. <b>Only used when method is "Wang" ("W").</b> Default is <code>CN.features</code> . Users can also set custom input with "feature", "min" and "max" columns available. Valid features can be printed by <code>unique(CN.features\$feature)</code> .
style	plot style, one of 'default' and 'cosmic', works when parameter <code>set_gradient_color</code> is FALSE.
palette	palette used to plot when <code>set_gradient_color</code> is FALSE, default use a built-in palette according to parameter style.
set_gradient_color	default is FALSE, if TRUE, use gradient colors to fill bars.
free_space	default is 'free_x'. If "fixed", all panels have the same size. If "free_y" their height will be proportional to the length of the y scale; if "free_x" their width will be proportional to the length of the x scale; or if "free" both height and width will vary. This setting has no effect unless the appropriate scales also vary.
rm_panel_border	default is TRUE for style 'cosmic', remove panel border to keep plot tight.
rm_grid_line	default is FALSE, if TRUE, remove grid lines of plot.
rm_axis_text	default is FALSE, if TRUE, remove component texts. This is useful when multiple signature profiles are plotted together.
bar_border_color	the color of bar border.
bar_width	bar width. By default, set to 70% of the resolution of the data.
paint_axis_text	if TRUE, color on text of x axis.
x_label_angle	font angle for x label.

x_label_vjust	font vjust for x label.
x_label_hjust	font hjust for x label.
x_lab	x axis lab.
y_lab	y axis lab.
y_limits	limits to expand in y axis. e.g., 0.2, c(0, 0.3).
params	params data. frame of components, obtained from <a href="#">sig_tally</a> .
show_cv	default is FALSE, if TRUE, show coefficient of variation when params is not NULL.
params_label_size	font size for params label.
params_label_angle	font angle for params label.
y_expand	y expand height for plotting params of copy number signatures.
digits	digits for plotting params of copy number signatures.
base_size	overall font size.
font_scale	a number used to set font scale.
sig_names	subset signatures or set name of signatures, can be a character vector. Default is NULL, prefix 'Sig' plus number is used.
sig_orders	set order of signatures, can be a character vector. Default is NULL, the signatures are ordered by alphabetical order. If an integer vector set, only specified signatures are plotted.
check_sig_names	if TRUE, check signature names when input is a matrix, i.e., all signatures (col-names) must start with 'Sig'.

**Value**

a ggplot object

**Author(s)**

Shixiang Wang

**See Also**

[show\\_sig\\_profile\\_loop](#), [show\\_sig\\_profile\\_heatmap](#)

**Examples**

```
# Load SBS signature
load(system.file("extdata", "toy_mutational_signature.RData",
  package = "sigminer", mustWork = TRUE
))
# Show signature profile
p1 <- show_sig_profile(sig2, mode = "SBS")
p1
```

```

# Use 'y_tr' option to transform values in y axis
p11 <- show_sig_profile(sig2, mode = "SBS", y_tr = function(x) x * 100)
p11

# Load copy number signature from method "W"
load(system.file("extdata", "toy_copynumber_signature_by_W.RData",
  package = "sigminer", mustWork = TRUE
))
# Show signature profile
p2 <- show_sig_profile(sig,
  style = "cosmic",
  mode = "copynumber",
  method = "W",
  normalize = "feature"
)
p2

# Visualize rearrangement signatures
s <- get_sig_db("RS_Nik_lab")
ss <- s$db[, 1:3]
colnames(ss) <- c("Sig1", "Sig2", "Sig3")
p3 <- show_sig_profile(ss, mode = "RS", style = "cosmic")
p3

```

---

```
show_sig_profile_heatmap
```

*Show Signature Profile with Heatmap*

---

## Description

This is a complementary function to [show\\_sig\\_profile\(\)](#), it is used for visualizing some big signatures, i.e. SBS-1536, not all signatures are supported. See details for current supported signatures.

## Usage

```

show_sig_profile_heatmap(
  Signature,
  mode = c("SBS", "DBS"),
  normalize = c("row", "column", "raw"),
  filters = NULL,
  x_lab = NULL,
  y_lab = NULL,
  legend_name = "auto",
  palette = "red",
  x_label_angle = 90,
  x_label_vjust = 1,
  x_label_hjust = 0.5,
  y_label_angle = 0,

```

```

    y_label_vjust = 0.5,
    y_label_hjust = 1,
    flip_xy = FALSE,
    sig_names = NULL,
    sig_orders = NULL,
    check_sig_names = TRUE
)

```

## Arguments

Signature	a Signature object obtained either from <a href="#">sig_extract</a> or <a href="#">sig_auto_extract</a> , or just a raw signature matrix with row representing components (motifs) and column representing signatures (column names must start with 'Sig').
mode	one of "SBS" and "DBS".
normalize	one of 'row', 'column', 'raw' and "feature", for row normalization (signature), column normalization (component), raw data, row normalization by feature, respectively. Of note, 'feature' only works when the mode is 'copynumber'.
filters	a pattern used to select components to plot.
x_lab	x label.
y_lab	y label.
legend_name	name of figure legend.
palette	color for value.
x_label_angle	angle for x axis text.
x_label_vjust	vjust for x axis text.
x_label_hjust	hjust for x axis text.
y_label_angle	angle for y axis text.
y_label_vjust	vjust for y axis text.
y_label_hjust	hjust for y axis text.
flip_xy	if TRUE, flip x axis and y axis.
sig_names	subset signatures or set name of signatures, can be a character vector. Default is NULL, prefix 'Sig' plus number is used.
sig_orders	set order of signatures, can be a character vector. Default is NULL, the signatures are ordered by alphabetical order. If an integer vector set, only specified signatures are plotted.
check_sig_names	if TRUE, check signature names when input is a matrix, i.e., all signatures (column names) must start with 'Sig'.

## Details

Support:

- SBS-24
- SBS-96

- SBS-384
- SBS-1536
- SBS-6144
- DBS-78
- DBS-186

### Value

a ggplot object.

### Examples

```
# Load SBS signature
load(system.file("extdata", "toy_mutational_signature.RData",
  package = "sigminer", mustWork = TRUE
))
# Show signature profile
p1 <- show_sig_profile_heatmap(sig2, mode = "SBS")
p1
```

---

show\_sig\_profile\_loop *Show Signature Profile with Loop Way*

---

### Description

Show Signature Profile with Loop Way

### Usage

```
show_sig_profile_loop(
  Signature,
  sig_names = NULL,
  ncol = 1,
  nrow = NULL,
  x_lab = "Components",
  ...
)
```

### Arguments

Signature	a Signature object obtained either from <a href="#">sig_extract</a> or <a href="#">sig_auto_extract</a> , or just a raw signature matrix with row representing components (motifs) and column representing signatures (column names must start with 'Sig').
sig_names	subset signatures or set name of signatures, can be a character vector. Default is NULL, prefix 'Sig' plus number is used.
ncol	(optional) Number of columns in the plot grid.



nrow (optional) Number of rows in the plot grid.  
x\_lab x axis lab.  
... other parameters but sig\_order passing to [show\\_sig\\_profile](#).

### Value

a ggplot result from `cowplot::plot_grid()`.

### See Also

[show\\_sig\\_profile](#)

### Examples

```
load(system.file("extdata", "toy_mutational_signature.RData",
  package = "sigminer", mustWork = TRUE
))
# Show signature profile
p1 <- show_sig_profile_loop(sig2, mode = "SBS")
p1
p2 <- show_sig_profile_loop(sig2, mode = "SBS", style = "cosmic", sig_names = c("A", "B", "C"))
p2
```

---

sigminer

*sigminer: Extract, Analyze and Visualize Signatures for Genomic Variations*

---

### Description

- Author: [Shixiang Wang \(w\\_shixiang@163.com\)](mailto:w_shixiang@163.com)
- Please go to <https://shixiangwang.github.io/sigminer-doc/> for full vignette.
- Please go to <https://shixiangwang.github.io/sigminer/reference/index.html> for organized documentation of functions and datasets.
- Result visualization for [MAF](#) is provide by [maftools](#) package, please read its [vignette](#).

sigprofiler

*Extract Signatures with SigProfiler***Description**

This function provides an interface to software SigProfiler. More please see <https://github.com/AlexandrovLab/SigProfilerExtractor>. Typically, a reference genome is not required because the input is a matrix (my understanding).

**Usage**

```
sigprofiler_extract(
  nmf_matrix,
  output,
  range = 2:5,
  nrun = 10L,
  refit = FALSE,
  refit_plot = FALSE,
  is_exome = FALSE,
  init_method = c("nndsvd_min", "random", "alexandrov-lab-custom", "nndsvd", "nndsvda",
    "nndsvdar"),
  cores = -1L,
  genome_build = c("hg19", "hg38", "mm10", "mm9"),
  use_conda = FALSE,
  py_path = NULL,
  sigprofiler_version = "1.1.3"
)

sigprofiler_import(
  output,
  order_by_expo = FALSE,
  type = c("suggest", "refit", "all")
)
```

**Arguments**

nmf_matrix	a matrix used for NMF decomposition with rows indicate samples and columns indicate components.
output	output directory.
range	signature number range, i.e. 2:5.
nrun	the number of iteration to be performed to extract each signature number.
refit	if TRUE, then refit the denovo signatures with nnls. Same meaning as optimize option in <a href="#">sig_extract</a> or <a href="#">sig_auto_extract</a> .
refit_plot	if TRUE, SigProfiler will make denovo to COSMIC sigantures decomposition plots. However, this may fail due to some matrix cannot be identified by Sig-Profiler plot program.

is_exome	if TRUE, the exomes will be extracted.
init_method	the initialization algorithm for W and H matrix of NMF. Options are 'random', 'nndsvd', 'nndsvda', 'nndsvdar', 'alexandrov-lab-custom' and 'nndsvd_min'.
cores	number of cores used for computation.
genome_build	I think this option is useless when input is matrix, keep it in case it is useful.
use_conda	if TRUE, create an independent conda environment to run SigProfiler.
py_path	path to Python executable file, e.g. '/Users/wsx/anaconda3/bin/python'.
sigprofiler_version	version of SigProfilerExtractor. If this package is not installed, the specified package will be installed. If this package is installed, this option is useless.
order_by_expo	if TRUE, order the import signatures by their exposures, e.g. the signature contributed the most exposure in all samples will be named as Sig1.
type	one of 'suggest' (for suggested solution), 'refit' (for refit solution) or 'all' (for all solutions).

### Value

For sigprofiler\_extract(), returns nothing. See output directory.

For sigprofiler\_import(), a list containing Signature object.

### Examples

```
if (FALSE) {
  load(system.file("extdata", "toy_copynumber_tally_W.RData",
    package = "sigminer", mustWork = TRUE
  ))

  reticulate::conda_list()

  sigprofiler_extract(cn_tally_W$nmf_matrix, "~/test/test_sigminer",
    use_conda = TRUE
  )

  sigprofiler_extract(cn_tally_W$nmf_matrix, "~/test/test_sigminer",
    use_conda = FALSE, py_path = "/Users/wsx/anaconda3/bin/python"
  )
}
```

## Description

A bayesian variant of NMF algorithm to enable optimal inferences for the number of signatures through the automatic relevance determination technique. This functions delivers highly interpretable and sparse representations for both signature profiles and attributions at a balance between data fitting and model complexity (this method may introduce more signatures than expected, especially for copy number signatures (thus **I don't recommend you to use this feature to extract copy number signatures**)). See detail part and references for more.

## Usage

```
sig_auto_extract(
  nmf_matrix = NULL,
  result_prefix = "BayesNMF",
  destdir = tempdir(),
  method = c("L1W.L2H", "L1KL", "L2KL"),
  strategy = c("stable", "optimal", "ms"),
  ref_sigs = NULL,
  K0 = 25,
  nrun = 10,
  niter = 2e+05,
  tol = 1e-07,
  cores = 1,
  optimize = FALSE,
  skip = FALSE,
  recover = FALSE
)
```

## Arguments

nmf_matrix	a matrix used for NMF decomposition with rows indicate samples and columns indicate components.
result_prefix	prefix for result data files.
destdir	path to save data runs, default is tempdir().
method	default is "L1W.L2H", which uses an exponential prior for W and a half-normal prior for H (This method is used by PCAWG project, see reference #3). You can also use "L1KL" to set expoential priors for both W and H, and "L2KL" to set half-normal priors for both W and H. The latter two methods are originally implemented by <a href="#">SignatureAnalyzer software</a> .
strategy	the selection strategy for returned data. Set 'stable' for getting optimal result from the most frequent K. Set 'optimal' for getting optimal result from all Ks. Set 'ms' for getting result with maximum mean cosine similarity with provided reference signatures. See ref_sigs option for details. If you want select other solution, please check <a href="#">get_bayesian_result</a> .
ref_sigs	A Signature object or matrix or string for specifying reference signatures, only used when strategy = 'ms'. See Signature and sig_db options in <a href="#">get_sig_similarity</a> for details.
K0	number of initial signatures.

nrun	number of independent simulations.
niter	the maximum number of iterations.
tol	tolerance for convergence.
cores	number of cpu cores to run NMF.
optimize	if TRUE, then refit the denovo signatures with QP method, see <a href="#">sig_fit</a> .
skip	if TRUE, it will skip running a previous stored result. This can be used to extend run times, e.g. you try running 10 times firstly and then you want to extend it to 20 times.
recover	if TRUE, try to recover result from previous runs based on input result_prefix, destdir and nrun. This is pretty useful for reproducing result. Please use skip if you want to recover an unfinished job.

### Details

There are three methods available in this function: "L1W.L2H", "L1KL" and "L2KL". They use different priors for the bayesian variant of NMF algorithm (see method parameter) written by reference #1 and implemented in [SignatureAnalyzer software](#) (reference #2).

I copied source code for the three methods from Broad Institute and supplementary files of reference #3, and wrote this higher function. It is more friendly for users to extract, visualize and analyze signatures by combining with other powerful functions in **sigminer** package. Besides, I implemented parallel computation to speed up the calculation process and a similar input and output structure like [sig\\_extract\(\)](#).

### Value

a list with Signature class.

### Author(s)

Shixiang Wang

### References

Tan, Vincent YF, and Cédric Févotte. "Automatic relevance determination in nonnegative matrix factorization with the/spl beta/-divergence." *IEEE Transactions on Pattern Analysis and Machine Intelligence* 35.7 (2012): 1592-1605.

Kim, Jaegil, et al. "Somatic ERCC2 mutations are associated with a distinct genomic signature in urothelial tumors." *Nature genetics* 48.6 (2016): 600.

Alexandrov, Ludmil, et al. "The repertoire of mutational signatures in human cancer." *BioRxiv* (2018): 322859.

### See Also

[sig\\_tally](#) for getting variation matrix, [sig\\_extract](#) for extracting signatures using NMF package, [sig\\_estimate](#) for estimating signature number for [sig\\_extract](#).

**Examples**

```

load(system.file("extdata", "toy_copynumber_tally_W.RData",
  package = "sigminer", mustWork = TRUE
))
res <- sig_auto_extract(cn_tally_W$nmf_matrix, result_prefix = "Test_copynumber", nrun = 1)
# At default, all run files are stored in tempdir()
dir(tempdir(), pattern = "Test_copynumber")

laml.maf <- system.file("extdata", "tcga_laml.maf.gz", package = "maftools")
laml <- read_maf(maf = laml.maf)
mt_tally <- sig_tally(
  laml,
  ref_genome = "BSgenome.Hsapiens.UCSC.hg19",
  use_syn = TRUE
)

x <- sig_auto_extract(mt_tally$nmf_matrix,
  strategy = "ms", nrun = 3, ref_sigs = "legacy"
)
x

```

sig\_convert

*Convert Signatures between different Genomic Distribution of Components***Description**

Converts signatures between two representations relative to different sets of mutational opportunities. Currently, only SBS signature is supported.

**Usage**

```
sig_convert(sig, from = "human-genome", to = "human-exome")
```

**Arguments**

sig	a Signature object obtained either from <code>sig_extract</code> or <code>sig_auto_extract</code> , or just a raw signature matrix/data.frame with row representing components (motifs) and column representing signatures.
from	either one of "human-genome" and "human-exome" or an opportunity matrix (repeated n columns with each row represents the total number of mutations for a component, n is the number of signature).
to	same as from.

**Details**

The default opportunity matrix for "human-genome" and "human-exome" comes from COSMIC signature database v2 and v3.

**Value**

a matrix.

**References**

convert\_signatures function from sigfit package.

**Examples**

```
# Load SBS signature
load(system.file("extdata", "toy_mutational_signature.RData",
  package = "sigminer", mustWork = TRUE
))
# Exome-relative to Genome-relative
sig_converted <- sig_convert(sig2,
  from = "human-exome",
  to = "human-genome"
)
sig_converted

show_sig_profile(sig2, style = "cosmic")
show_sig_profile(sig_converted, style = "cosmic")
```

---

sig\_estimate

*Estimate Signature Number*


---

**Description**

Use **NMF** package to evaluate the optimal number of signatures. This is used along with [sig\\_extract](#). Users should `library(NMF)` firstly. If **NMF** objects are returned, the result can be further visualized by **NMF** plot methods like `NMF::consensusmap()` and `NMF::basismap()`.

`sig_estimate()` shows comprehensive rank survey generated by **NMF** package, sometimes it is hard to consider all measures. `show_sig_number_survey()` provides a one or two y-axis visualization method to help users determine the optimal signature number (showing both stability ("cophe-netic") and error (RSS) at default). Users can also set custom measures to show.

`show_sig_number_survey2()` is modified from **NMF** package to better help users to explore survey of signature number.

**Usage**

```
sig_estimate(
  nmf_matrix,
  range = 2:5,
  nrun = 10,
  use_random = FALSE,
  method = "brunet",
  seed = 123456,
```

```

cores = 1,
keep_nmfObj = FALSE,
save_plots = FALSE,
plot_basename = file.path(tempdir(), "nmf"),
what = "all",
verbose = FALSE
)

show_sig_number_survey(
  object,
  x = "rank",
  left_y = "cophenetic",
  right_y = "rss",
  left_name = left_y,
  right_name = toupper(right_y),
  left_color = "black",
  right_color = "red",
  left_shape = 16,
  right_shape = 18,
  shape_size = 4,
  highlight = NULL
)

show_sig_number_survey2(
  x,
  y = NULL,
  what = c("all", "cophenetic", "rss", "residuals", "dispersion", "evar", "sparseness",
    "sparseness.basis", "sparseness.coef", "silhouette", "silhouette.coef",
    "silhouette.basis", "silhouette.consensus"),
  na.rm = FALSE,
  xlab = "Total signatures",
  ylab = "",
  main = "Signature number survey using NMF package"
)

```

### Arguments

<code>nmf_matrix</code>	a matrix used for NMF decomposition with rows indicate samples and columns indicate components.
<code>range</code>	a numeric vector containing the ranks of factorization to try. Note that duplicates are removed and values are sorted in increasing order. The results are notably returned in this order.
<code>nrun</code>	a numeric giving the number of run to perform for each value in range, nrun set to 30~50 is enough to achieve robust result.
<code>use_random</code>	Should generate random data from input to test measurements. Default is TRUE.
<code>method</code>	specification of the NMF algorithm. Use 'brunet' as default. Available methods for NMF decompositions are 'brunet', 'lee', 'ls-nmf', 'nsNMF', 'offset'.



seed	specification of the starting point or seeding method, which will compute a starting point, usually using data from the target matrix in order to provide a good guess.
cores	number of cpu cores to run NMF.
keep_nmfObj	default is FALSE, if TRUE, keep NMF objects from runs, and the result may be huge.
save_plots	if TRUE, save signature number survey plot to local machine.
plot_basename	when save plots, set custom basename for file path.
what	a character vector whose elements partially match one of the following item, which correspond to the measures computed by <code>summary()</code> on each – multi-run – NMF result: 'all', 'cophenetic', 'rss', 'residuals', 'dispersion', 'evar', 'silhouette' (and more specific <code>*.coef</code> , <code>*.basis</code> , <code>*.consensus</code> ), 'sparseness' (and more specific <code>*.coef</code> , <code>*.basis</code> ). It specifies which measure must be plotted (what='all' plots all the measures).
verbose	if TRUE, print extra message.
object	a Survey object generated from <code>sig_estimate</code> , or a <code>data.frame</code> contains at least rank columns and columns for one measure.
x	a <code>data.frame</code> or <code>NMF.rank</code> object obtained from <code>sig_estimate()</code> .
left_y	column name for left y axis.
right_y	column name for right y axis.
left_name	label name for left y axis.
right_name	label name for right y axis.
left_color	color for left axis.
right_color	color for right axis.
left_shape, right_shape, shape_size	shape setting.
highlight	a integer to highlight a x.
y	for random simulation, a <code>data.frame</code> or <code>NMF.rank</code> object obtained from <code>sig_estimate()</code> .
na.rm	single logical that specifies if the rank for which the measures are NA values should be removed from the graph or not (default to FALSE). This is useful when plotting results which include NAs due to error during the estimation process. See argument <code>stop</code> for <code>nmfEstimateRank</code> .
xlab	x-axis label
ylab	y-axis label
main	main title

## Details

The most common approach is to choose the smallest rank for which cophenetic correlation coefficient starts decreasing (Used by this function). Another approach is to choose the rank for which the plot of the residual sum of squares (RSS) between the input matrix and its estimate shows an inflection point. More custom features please directly use `NMF::nmfEstimateRank`.

**Value**

- sig\_estimate: a list contains information of NMF run and rank survey.
- show\_sig\_number\_survey: a ggplot object
- show\_sig\_number\_survey2: a ggplot object

**Author(s)**

Shixiang Wang

**References**

Gaujoux, Renaud, and Cathal Seoighe. "A flexible R package for nonnegative matrix factorization." BMC bioinformatics 11.1 (2010): 367.

**See Also**

[sig\\_extract](#) for extracting signatures using NMF package, [sig\\_auto\\_extract](#) for extracting signatures using automatic relevance determination technique.

[sig\\_estimate](#) for estimating signature number for [sig\\_extract](#), [show\\_sig\\_number\\_survey2](#) for more visualization method.

**Examples**

```
load(system.file("extdata", "toy_copynumber_tally_W.RData",
  package = "sigminer", mustWork = TRUE
))
library(NMF)
cn_estimate <- sig_estimate(cn_tally_W$nmf_matrix,
  cores = 1, nrun = 5,
  verbose = TRUE
)

p <- show_sig_number_survey2(cn_estimate$survey)
p

# Show two measures
show_sig_number_survey(cn_estimate)
# Show one measure
p1 <- show_sig_number_survey(cn_estimate, right_y = NULL)
p1
p2 <- add_h_arrow(p, x = 4.1, y = 0.953, label = "selected number")
p2

# Show data from a data.frame
p3 <- show_sig_number_survey(cn_estimate$survey)
p3
# Show other measures
head(cn_estimate$survey)
```

```
p4 <- show_sig_number_survey(cn_estimate$survey,
  right_y = "dispersion",
  right_name = "dispersion"
)
p4
p5 <- show_sig_number_survey(cn_estimate$survey,
  right_y = "evar",
  right_name = "evar"
)
p5
```

---

sig\_extract

*Extract Signatures through NMF*

---

## Description

Do NMF de-composition and then extract signatures.

## Usage

```
sig_extract(
  nmf_matrix,
  n_sig,
  nrun = 10,
  cores = 1,
  method = "brunet",
  optimize = FALSE,
  pynmf = FALSE,
  use_conda = TRUE,
  py_path = "/Users/wsx/anaconda3/bin/python",
  seed = 123456,
  ...
)
```

## Arguments

nmf_matrix	a matrix used for NMF decomposition with rows indicate samples and columns indicate components.
n_sig	number of signature. Please run <a href="#">sig_estimate</a> to select a suitable value.
nrun	a numeric giving the number of run to perform for each value in range, nrun set to 30~50 is enough to achieve robust result.
cores	number of cpu cores to run NMF.
method	specification of the NMF algorithm. Use 'brunet' as default. Available methods for NMF decompositions are 'brunet', 'lee', 'ls-nmf', 'nsNMF', 'offset'.
optimize	if TRUE, then refit the denovo signatures with QP method, see <a href="#">sig_fit</a> .

pynmf	if TRUE, use Python NMF driver <b>Nimfa</b> . The seed currently is not used by this implementation.
use_conda	if TRUE, create an independent conda environment to run NMF.
py_path	path to Python executable file, e.g. <code>'/Users/wsx/anaconda3/bin/python'</code> . In my test, it is more stable than <code>use_conda=TRUE</code> . You can install the Nimfa package by yourself or set <code>use_conda</code> to TRUE to install required Python environment, and then set this option.
seed	specification of the starting point or seeding method, which will compute a starting point, usually using data from the target matrix in order to provide a good guess.
...	other arguments passed to <code>NMF::nmf()</code> .

### Value

a list with Signature class.

### Author(s)

Shixiang Wang

### References

Gaujoux, Renaud, and Cathal Seoighe. "A flexible R package for nonnegative matrix factorization." *BMC bioinformatics* 11.1 (2010): 367.

Mayakonda, Anand, et al. "Maftools: efficient and comprehensive analysis of somatic variants in cancer." *Genome research* 28.11 (2018): 1747-1756.

### See Also

[sig\\_tally](#) for getting variation matrix, [sig\\_estimate](#) for estimating signature number for [sig\\_extract](#), [sig\\_auto\\_extract](#) for extracting signatures using automatic relevance determination technique.

### Examples

```
load(system.file("extdata", "toy_copynumber_tally_W.RData",
  package = "sigminer", mustWork = TRUE
))
# Extract copy number signatures
res <- sig_extract(cn_tally_W$nmf_matrix, 2, nrun = 1)
```

sig\_fit

*Fit Signature Exposures with Linear Combination Decomposition***Description**

The function performs a signatures decomposition of a given mutational catalogue  $V$  with known signatures  $W$  by solving the minimization problem  $\min(\|W*H - V\|)$  where  $W$  and  $V$  are known.

**Usage**

```
sig_fit(
  catalogue_matrix,
  sig,
  sig_index = NULL,
  sig_db = c("legacy", "SBS", "DBS", "ID", "TSB", "SBS_Nik_lab", "RS_Nik_lab",
    "RS_BRCA560", "RS_USARC", "CNS_USARC", "CNS_TCGA", "SBS_hg19", "SBS_hg38", "SBS_mm9",
    "SBS_mm10", "DBS_hg19", "DBS_hg38", "DBS_mm9", "DBS_mm10", "SBS_Nik_lab_Organ",
    "RS_Nik_lab_Organ", "latest_SBS_GRCh37", "latest_DBS_GRCh37", "latest_ID_GRCh37",
    "latest_SBS_GRCh38", "latest_DBS_GRCh38", "latest_SBS_mm9", "latest_DBS_mm9",
    "latest_SBS_mm10", "latest_DBS_mm10", "latest_SBS_rn6", "latest_DBS_rn6"),
  db_type = c("", "human-exome", "human-genome"),
  show_index = TRUE,
  method = c("QP", "NNLS", "SA"),
  auto_reduce = FALSE,
  type = c("absolute", "relative"),
  return_class = c("matrix", "data.table"),
  return_error = FALSE,
  rel_threshold = 0,
  mode = c("SBS", "DBS", "ID", "copynumber"),
  true_catalog = NULL,
  ...
)
```

**Arguments**

catalogue_matrix	a numeric matrix $V$ with row representing components and columns representing samples, typically you can get <code>nmf_matrix</code> from <code>sig_tally()</code> and transpose it by <code>t()</code> .
sig	a Signature object obtained either from <code>sig_extract</code> or <code>sig_auto_extract</code> , or just a raw signature matrix/data.frame with row representing components (motifs) and column representing signatures.
sig_index	a vector for signature index. "ALL" for all signatures.
sig_db	default 'legacy', it can be 'legacy' (for <b>COSMIC v2 'SBS'</b> ), 'SBS', 'DBS', 'ID' and 'TSB' (for <b>COSMIV v3.1 signatures</b> ) for small scale mutations. For more specific details, it can also be 'SBS_hg19', 'SBS_hg38', 'SBS_mm9',

'SBS\_mm10', 'DBS\_hg19', 'DBS\_hg38', 'DBS\_mm9', 'DBS\_mm10' to use COSMIC v3 reference signatures from Alexandrov, Ludmil B., et al. (2020) (reference #1). In addition, it can be one of "SBS\_Nik\_lab\_Organ", "RS\_Nik\_lab\_Organ", "SBS\_Nik\_lab", "RS\_Nik\_lab" to refer reference signatures from Degasperi, Andrea, et al. (2020) (reference #2); "RS\_BRCA560", "RS\_USARC" to reference signatures from BRCA560 and USARC cohorts; "CNS\_USARC" (40 categories), "CNS\_TCGA" (48 categories) to reference copy number signatures from USARC cohort and TCGA. **UPDATE**, the latest version of reference version can be automatically downloaded and loaded from <https://cancer.sanger.ac.uk/signatures/downloads/> when a option with latest\_prefix is specified (e.g. "latest\_SBS\_GRCh37"). **Note**: the signature profile for different genome builds are basically same. And specific database (e.g. 'SBS\_mm10') contains less signatures than all COSMIC signatures (because some signatures are not detected from Alexandrov, Ludmil B., et al. (2020)). For all available options, check the parameter setting.

db_type	only used when sig_db is enabled. "" for keeping default, "human-exome" for transforming to exome frequency of component, and "human-genome" for transforming to whole genome frequency of component. Currently only works for 'SBS'.
show_index	if TRUE, show valid indices.
method	method to solve the minimization problem. 'NNLS' for non-negative least square; 'QP' for quadratic programming; 'SA' for simulated annealing.
auto_reduce	if TRUE, try reducing the input reference signatures to increase the cosine similarity of reconstructed profile to observed profile.
type	'absolute' for signature exposure and 'relative' for signature relative exposure.
return_class	string, 'matrix' or 'data.table'.
return_error	if TRUE, also return sample error (Frobenius norm) and cosine similarity between observed sample profile (asa. spectrum) and reconstructed profile. NOTE: it is better to obtain the error when the type is 'absolute', because the error is affected by relative exposure accuracy.
rel_threshold	numeric vector, a signature with relative exposure lower than (equal is included, i.e. <=) this value will be set to 0 (both absolute exposure and relative exposure). In this case, sum of signature contribution may not equal to 1.
mode	signature type for plotting, now supports 'copynumber', 'SBS', 'DBS', 'ID' and 'RS' (genome rearrangement signature).
true_catalog	used by <a href="#">sig_fit_bootstrap</a> , user never use it.
...	control parameters passing to argument control in GenSA function when use method 'SA'.

## Details

The method 'NNLS' solves the minimization problem with nonnegative least-squares constraints. The method 'QP' and 'SA' are modified from SignatureEstimation package. See references for details. Of note, when fitting exposures for copy number signatures, only components of feature CN is used.

**Value**

The exposure result either in `matrix` or `data.table` format. If `return_error` set `TRUE`, a list is returned.

**References**

Daniel Huebschmann, Zuguang Gu and Matthias Schlesner (2019). YAPSA: Yet Another Package for Signature Analysis. R package version 1.12.0.

Huang X, Wojtowicz D, Przytycka TM. Detecting presence of mutational signatures in cancer with confidence. *Bioinformatics*. 2018;34(2):330–337. doi:10.1093/bioinformatics/btx604

Kim, Jaegil, et al. "Somatic ERCC2 mutations are associated with a distinct genomic signature in urothelial tumors." *Nature genetics* 48.6 (2016): 600.

**See Also**

[sig\\_extract](#), [sig\\_auto\\_extract](#), [sig\\_fit\\_bootstrap](#), [sig\\_fit\\_bootstrap\\_batch](#)

**Examples**

```
W <- matrix(c(1, 2, 3, 4, 5, 6), ncol = 2)
colnames(W) <- c("sig1", "sig2")
W <- apply(W, 2, function(x) x / sum(x))

H <- matrix(c(2, 5, 3, 6, 1, 9, 1, 2), ncol = 4)
colnames(H) <- paste0("samp", 1:4)

V <- W %*% H
V

if (requireNamespace("quadprog", quietly = TRUE)) {
  H_infer <- sig_fit(V, W, method = "QP")
  H_infer
  H

  H_dt <- sig_fit(V, W, method = "QP", auto_reduce = TRUE, return_class = "data.table")
  H_dt

  ## Show results
  show_sig_fit(H_infer)
  show_sig_fit(H_dt)

  ## Get clusters/groups
  H_dt_rel <- sig_fit(V, W, return_class = "data.table", type = "relative")
  z <- get_groups(H_dt_rel, method = "k-means")
  show_groups(z)
}

# if (requireNamespace("GenSA", quietly = TRUE)) {
#   H_infer <- sig_fit(V, W, method = "SA")
#   H_infer
#   H
# }
```

```

#
# H_dt <- sig_fit(V, W, method = "SA", return_class = "data.table")
# H_dt
#
# ## Modify arguments to method
# sig_fit(V, W, method = "SA", maxit = 10, temperature = 100)
#
# ## Show results
# show_sig_fit(H_infer)
# show_sig_fit(H_dt)
# }

```

---

sig_fit_bootstrap	<i>Obtain Bootstrap Distribution of Signature Exposures of a Certain Tumor Sample</i>
-------------------	---

---

### Description

This can be used to obtain the confidence of signature exposures or search the suboptimal decomposition solution.

### Usage

```

sig_fit_bootstrap(
  catalog,
  sig,
  n = 100L,
  sig_index = NULL,
  sig_db = "legacy",
  db_type = c("", "human-exome", "human-genome"),
  show_index = TRUE,
  method = c("QP", "NNLS", "SA"),
  auto_reduce = FALSE,
  SA_not_bootstrap = FALSE,
  type = c("absolute", "relative"),
  rel_threshold = 0,
  mode = c("SBS", "DBS", "ID", "copynumber"),
  find_suboptimal = FALSE,
  suboptimal_ref_error = NULL,
  suboptimal_factor = 1.05,
  ...
)

```

### Arguments

catalog	a named numeric vector or a numeric matrix with dimension Nx1. N is the number of component, 1 is the sample.
---------	---



sig	a Signature object obtained either from <code>sig_extract</code> or <code>sig_auto_extract</code> , or just a raw signature matrix/data.frame with row representing components (motifs) and column representing signatures.
n	the number of bootstrap replicates.
sig_index	a vector for signature index. "ALL" for all signatures.
sig_db	default 'legacy', it can be 'legacy' (for <b>COSMIC v2 'SBS'</b> ), 'SBS', 'DBS', 'ID' and 'TSB' (for <b>COSMIV v3.1 signatures</b> ) for small scale mutations. For more specific details, it can also be 'SBS_hg19', 'SBS_hg38', 'SBS_mm9', 'SBS_mm10', 'DBS_hg19', 'DBS_hg38', 'DBS_mm9', 'DBS_mm10' to use COSMIC v3 reference signatures from Alexandrov, Ludmil B., et al. (2020) (reference #1). In addition, it can be one of "SBS_Nik_lab_Organ", "RS_Nik_lab_Organ", "SBS_Nik_lab", "RS_Nik_lab" to refer reference signatures from Degasperi, Andrea, et al. (2020) (reference #2); "RS_BRCA560", "RS_USARC" to reference signatures from BRCA560 and USARC cohorts; "CNS_USARC" (40 categories), "CNS_TCGA" (48 categories) to reference copy number signatures from USARC cohort and TCGA. <b>UPDATE</b> , the latest version of reference version can be automatically downloaded and loaded from <a href="https://cancer.sanger.ac.uk/signatures/downloads/">https://cancer.sanger.ac.uk/signatures/downloads/</a> when a option with latest_prefix is specified (e.g. "latest_SBS_GRCh37"). <b>Note</b> : the signature profile for different genome builds are basically same. And specific database (e.g. 'SBS_mm10') contains less signatures than all COSMIC signatures (because some signatures are not detected from Alexandrov, Ludmil B., et al. (2020)). For all available options, check the parameter setting.
db_type	only used when sig_db is enabled. "" for keeping default, "human-exome" for transforming to exome frequency of component, and "human-genome" for transforming to whole genome frequency of component. Currently only works for 'SBS'.
show_index	if TRUE, show valid indices.
method	method to solve the minimization problem. 'NNLS' for non-negative least square; 'QP' for quadratic programming; 'SA' for simulated annealing.
auto_reduce	if TRUE, try reducing the input reference signatures to increase the cosine similarity of reconstructed profile to observed profile.
SA_not_bootstrap	if TRUE, directly run 'SA' multiple times with original input instead of bootstrap samples.
type	'absolute' for signature exposure and 'relative' for signature relative exposure.
rel_threshold	numeric vector, a signature with relative exposure lower than (equal is included, i.e. <=) this value will be set to 0 (both absolute exposure and relative exposure). In this case, sum of signature contribution may not equal to 1.
mode	signature type for plotting, now supports 'copynumber', 'SBS', 'DBS', 'ID' and 'RS' (genome rearrangement signature).
find_suboptimal	logical, if TRUE, find suboptimal decomposition with slightly higher error than the optimal solution by method 'SA'. This is useful to explore hidden dependencies between signatures. More see reference.

suboptimal\_ref\_error  
 baseline error used for finding suboptimal solution. if it is NULL, then use 'SA' method to obtain the optimal error.

suboptimal\_factor  
 suboptimal factor to get suboptimal error, default is 1.05, i.e., suboptimal error is 1.05 times baseline error.

...  
 control parameters passing to argument control in GenSA function when use method 'SA'.

### Value

a list

### References

Huang X, Wojtowicz D, Przytycka TM. Detecting presence of mutational signatures in cancer with confidence. *Bioinformatics*. 2018;34(2):330–337. doi:10.1093/bioinformatics/btx604

### See Also

[report\\_bootstrap\\_p\\_value](#), [sig\\_fit](#), [sig\\_fit\\_bootstrap\\_batch](#)

### Examples

```
W <- matrix(c(1, 2, 3, 4, 5, 6), ncol = 2)
colnames(W) <- c("sig1", "sig2")
W <- apply(W, 2, function(x) x / sum(x))

H <- matrix(c(2, 5, 3, 6, 1, 9, 1, 2), ncol = 4)
colnames(H) <- paste0("samp", 1:4)

V <- W %*% H
V

if (requireNamespace("quadprog", quietly = TRUE)) {
  H_bootstrap <- sig_fit_bootstrap(V[, 1], W, n = 10, type = "absolute")
  ## Typically, you have to run many times to get close to the answer
  boxplot(t(H_bootstrap$expo))
  H[, 1]

  ## Return P values
  ## In practice, run times >= 100
  ## is recommended
  report_bootstrap_p_value(H_bootstrap)
  ## For multiple samples
  ## Input a list
  report_bootstrap_p_value(list(samp1 = H_bootstrap, samp2 = H_bootstrap))

  # ## Find suboptimal decomposition
  # H_suboptimal <- sig_fit_bootstrap(V[, 1], W,
  #   n = 10,
```

```

#   type = "absolute",
#   method = "SA",
#   find_suboptimal = TRUE
# )
}

```

---

sig\_fit\_bootstrap\_batch

*Exposure Instability Analysis of Signature Exposures with Bootstrapping*

---

## Description

Read [sig\\_fit\\_bootstrap](#) for more option setting.

## Usage

```

sig_fit_bootstrap_batch(
  catalogue_matrix,
  methods = c("QP"),
  n = 100L,
  min_count = 1L,
  p_val_thresholds = c(0.05),
  use_parallel = FALSE,
  seed = 123456L,
  job_id = NULL,
  result_dir = tempdir(),
  ...
)

```

## Arguments

catalogue_matrix	a numeric matrix V with row representing components and columns representing samples, typically you can get nmf_matrix from sig_tally() and transpose it by t().
methods	a subset of c("NNLS", "QP", "SA").
n	the number of bootstrap replicates.
min_count	minimal exposure in a sample, default is 1. Any patient has total exposure less than this value will be filtered out.
p_val_thresholds	a vector of relative exposure threshold for calculating p values.
use_parallel	if TRUE, use parallel computation based on <b>furrr</b> package. It can also be an integer for specifying cores.
seed	random seed to reproduce the result.

**job\_id** a job ID, default is NULL, can be a string. When not NULL, all bootstrapped results will be saved to local machine location defined by `result_dir`. This is very useful for running more than 10 times for more than 100 samples.  
**result\_dir** see above, default is temp directory defined by R.  
**...** other common parameters passing to [sig\\_fit\\_bootstrap](#), including `sig`, `sig_index`, `sig_db`, `db_type`, `mode`, `auto_reduce` etc.

**Value**

a list of `data.table`.

**See Also**

[sig\\_fit](#), [sig\\_fit\\_bootstrap](#)

**Examples**

```

W <- matrix(c(1, 2, 3, 4, 5, 6), ncol = 2)
colnames(W) <- c("sig1", "sig2")
W <- apply(W, 2, function(x) x / sum(x))

H <- matrix(c(2, 5, 3, 6, 1, 9, 1, 2), ncol = 4)
colnames(H) <- paste0("samp", 1:4)

V <- W %*% H
V

if (requireNamespace("quadprog")) {
  z10 <- sig_fit_bootstrap_batch(V, sig = W, n = 10)
  z10
}

```

---

sig\_operation

*Obtain or Modify Signature Information*

---

**Description**

Obtain or Modify Signature Information

**Usage**

```

sig_names(sig)

sig_modify_names(sig, new_names)

sig_number(sig)

sig_attrs(sig)

```

```
sig_signature(sig, normalize = c("row", "column", "raw", "feature"))
sig_exposure(sig, type = c("absolute", "relative"))
```

### Arguments

sig	a Signature object obtained either from <a href="#">sig_extract</a> or <a href="#">sig_auto_extract</a> .
new_names	new signature names.
normalize	one of 'row', 'column', 'raw' and "feature", for row normalization (signature), column normalization (component), raw data, row normalization by feature, respectively.
type	one of 'absolute' and 'relative'.

### Value

a Signature object or data.

### Examples

```
## Operate signature names
load(system.file("extdata", "toy_mutational_signature.RData",
  package = "sigminer", mustWork = TRUE
))
sig_names(sig2)
cc <- sig_modify_names(sig2, new_names = c("Sig2", "Sig1", "Sig3"))
sig_names(cc)

# The older names are stored in tags.
print(attr(cc, "tag"))
## Get signature number
sig_number(sig2)
## Get signature attributes
sig_number(sig2)
## Get signature matrix
z <- sig_signature(sig2)
z <- sig_signature(sig2, normalize = "raw")
## Get exposure matrix
## Of note, this is different from get_sig_exposure()
## it returns a matrix instead of data table.
z <- sig_exposure(sig2) # it is same as sig$Exposure
z <- sig_exposure(sig2, type = "relative") # it is same as sig2$Exposure.norm
```

## Description

Tally a variation object like [MAF](#), [CopyNumber](#) and return a matrix for NMF de-composition and more. This is a generic function, so it can be further extended to other mutation cases. **Please read details about how to set sex for identifying copy number signatures.** Please read <https://osf.io/s93d5/> for the generation of SBS, DBS and ID (INDEL) components.

## Usage

```
sig_tally(object, ...)

## S3 method for class 'CopyNumber'
sig_tally(
  object,
  method = "Wang",
  ignore_chrs = NULL,
  indices = NULL,
  add_loh = FALSE,
  feature_setting = sigminer::CN.features,
  cores = 1,
  keep_only_matrix = FALSE,
  ...
)

## S3 method for class 'RS'
sig_tally(object, keep_only_matrix = FALSE, ...)

## S3 method for class 'MAF'
sig_tally(
  object,
  mode = c("SBS", "DBS", "ID", "ALL"),
  ref_genome = "BSgenome.Hsapiens.UCSC.hg19",
  genome_build = NULL,
  add_trans_bias = FALSE,
  ignore_chrs = NULL,
  use_syn = TRUE,
  keep_only_matrix = FALSE,
  ...
)
```

## Arguments

object	a <a href="#">CopyNumber</a> object or <a href="#">MAF</a> object or SV object (from <a href="#">read_sv_as_rs</a> ).
...	custom setting for operating object. Detail see S3 method for corresponding class (e.g. CopyNumber).
method	method for feature classification, can be one of "Wang" ("W"), "S" (for method described in Steele et al. 2019).
ignore_chrs	Chromosomes to ignore from analysis. e.g. chrX and chrY.

indices	integer vector indicating segments to keep.
add_loh	flag to add LOH classifications.
feature_setting	a data.frame used for classification. <b>Only used when method is "Wang" ("W")</b> . Default is <code>CN.features</code> . Users can also set custom input with "feature", "min" and "max" columns available. Valid features can be printed by <code>unique(CN.features\$feature)</code> .
cores	number of computer cores to run this task. You can use <code>future::availableCores()</code> function to check how many cores you can use.
keep_only_matrix	if TRUE, keep only matrix for signature extraction. For a MAF object, this will just return the most useful matrix.
mode	type of mutation matrix to extract, can be one of 'SBS', 'DBS' and 'ID'.
ref_genome	'BSgenome.Hsapiens.UCSC.hg19', 'BSgenome.Hsapiens.UCSC.hg38', 'BSgenome.Mmusculus.UCSC.BSgenome.Mmusculus.UCSC.mm9', etc.
genome_build	genome build 'hg19', 'hg38', 'mm9' or "mm10", if not set, guess it by ref_genome.
add_trans_bias	if TRUE, consider transcriptional bias categories. 'T:' for Transcribed (the variant is on the transcribed strand); 'U:' for Un-transcribed (the variant is on the untranscribed strand); 'B:' for Bi-directional (the variant is on both strand and is transcribed either way); 'N:' for Non-transcribed (the variant is in a non-coding region and is untranslated); 'Q:' for Questionable. <b>NOTE:</b> the result counts of 'B' and 'N' labels are a little different from SigProfilerMatrixGenerator, the reason is unknown (may be caused by annotation file).
use_syn	Logical. If TRUE, include synonymous variants in analysis.

## Details

For identifying copy number signatures, we have to derive copy number features firstly. Due to the difference of copy number values in sex chromosomes between male and female, we have to do an extra step **if we don't want to ignore them**.

I create two options to control this, the default values are shown as the following, you can use the same way to set (per R session).

```
options(sigminer.sex = "female", sigminer.copynumber.max = NA_integer_)
```

- If your cohort are all females, you can totally ignore this.
- If your cohort are all males, set `sigminer.sex` to 'male' and `sigminer.copynumber.max` to a proper value (the best is consistent with [read\\_copynumber](#)).
- If your cohort contains both males and females, set `sigminer.sex` as a data.frame with two columns "sample" and "sex". And set `sigminer.copynumber.max` to a proper value (the best is consistent with [read\\_copynumber](#)).

## Value

a list contains a matrix used for NMF de-composition.

**Methods (by class)**

- CopyNumber: Returns copy number features, components and component-by-sample matrix
- RS: Returns genome rearrangement sample-by-component matrix
- MAF: Returns SBS mutation sample-by-component matrix and APOBEC enrichment

**Author(s)**

Shixiang Wang

**References**

- Wang, Shixiang, et al. "Copy number signature analyses in prostate cancer reveal distinct etiologies and clinical outcomes." medRxiv (2020).
- Steele, Christopher D., et al. "Undifferentiated sarcomas develop through distinct evolutionary pathways." Cancer Cell 35.3 (2019): 441-456.
- Mayakonda, Anand, et al. "Maftools: efficient and comprehensive analysis of somatic variants in cancer." Genome research 28.11 (2018): 1747-1756.
- Roberts SA, Lawrence MS, Klimczak LJ, et al. An APOBEC Cytidine Deaminase Mutagenesis Pattern is Widespread in Human Cancers. Nature genetics. 2013;45(9):970-976. doi:10.1038/ng.2702.
- Bergstrom EN, Huang MN, Mahto U, Barnes M, Stratton MR, Rozen SG, Alexandrov LB: Sig-ProfilerMatrixGenerator: a tool for visualizing and exploring patterns of small mutational events. BMC Genomics 2019, 20:685 <https://bmcgenomics.biomedcentral.com/articles/10.1186/s12864-019-6041-2>

**See Also**

[sig\\_estimate](#) for estimating signature number for [sig\\_extract](#), [sig\\_auto\\_extract](#) for extracting signatures using automatic relevance determination technique.

**Examples**

```
# Load copy number object
load(system.file("extdata", "toy_copynumber.RData",
  package = "sigminer", mustWork = TRUE
))

# Use method designed by Wang, Shixiang et al.
cn_tally_W <- sig_tally(cn, method = "W")

# Use method designed by Steele et al.
# See example in read_copynumber

# Prepare SBS signature analysis
laml.maf <- system.file("extdata", "tcga_laml.maf.gz", package = "maftools")
laml <- read_maf(maf = laml.maf)
if (require("BSgenome.Hsapiens.UCSC.hg19")) {
  mt_tally <- sig_tally(
    laml,
```



```

    ref_genome = "BSgenome.Hsapiens.UCSC.hg19",
    use_syn = TRUE
  )
  mt_tally$nmf_matrix[1:5, 1:5]

  ## Use strand bias categories
  mt_tally <- sig_tally(
    laml,
    ref_genome = "BSgenome.Hsapiens.UCSC.hg19",
    use_syn = TRUE, add_trans_bias = TRUE
  )
  ## Test it by enrichment analysis
  enrich_component_strand_bias(mt_tally$nmf_matrix)
  enrich_component_strand_bias(mt_tally$all_matrices$SBS_24)
} else {
  message("Please install package 'BSgenome.Hsapiens.UCSC.hg19' firstly!")
}

```

---

sig\_unify\_extract      *An Unified Interface to Extract Signatures*

---

### Description

This function provides an unified interface to signature extractor implemented in **sigminer**. If you determine a specific approach, please also read the documentation of corresponding extractor. See "Arguments" part.

### Usage

```

sig_unify_extract(
  nmf_matrix,
  range = 2:5,
  nrun = 10,
  approach = c("bayes_nmf", "repeated_nmf", "bootstrap_nmf", "sigprofiler"),
  cores = 1L,
  ...
)

```

### Arguments

nmf_matrix	a matrix used for NMF decomposition with rows indicate samples and columns indicate components.
range	signature number range, i.e. 2:5.
nrun	the number of iteration to be performed to extract each signature number.
approach	approach name. <ul style="list-style-type: none"> <li>• "repeated_nmf" - <a href="#">sig_extract</a></li> </ul>

- "bayes\_nmf" - [sig\\_auto\\_extract](#)
- "bootstrap\_nmf" - [bp\\_extract\\_signatures](#)
- "sigprofiler" - [sigprofiler](#)

cores            number of cores used for computation.

...             other parameters passing to signature extractor based on the approach setting.

**Value**

Result dependent on the approach setting.

**See Also**

[sig\\_extract](#), [sig\\_auto\\_extract](#), [bp\\_extract\\_signatures](#), [sigprofiler](#)

**Examples**

```
load(system.file("extdata", "toy_copynumber_tally_W.RData",
  package = "sigminer", mustWork = TRUE
))
# Extract signatures
# It is same as sig_extract(cn_tally_W$nmf_matrix, 2, nrun = 1)
res <- sig_unify_extract(cn_tally_W$nmf_matrix, 2,
  nrun = 1,
  approach = "repeated_nmf"
)
# Auto-extract signatures based on bayesian NMF
res2 <- sig_unify_extract(cn_tally_W$nmf_matrix,
  nrun = 1,
  approach = "bayes_nmf"
)
```

---

simulated\_catalogs      *A List of Simulated SBS-96 Catalog Matrix*

---

**Description**

Data from [doi:10.1038/s4301802000275](https://doi.org/10.1038/s4301802000275). 5 simulated mutation catalogs are used by the paper but only 4 are available. The data are simulated from COSMIC mutational signatures 1, 2, 3, 5, 6, 8, 12, 13, 17 and 18. Each sample is a linear combination of 5 randomly selected signatures with the addition of Poisson noise. The number of mutation in each sample is randomly selected between 1,000 and 50,000 mutations, in log scale so that a lower number of mutations is more likely to be selected. The proportion of each signature in each sample is also random.

**Format**

A list of matrix

**Source**

Generate from code under data\_raw/

**Examples**

```
data(simulated_catalogs)
```

---

simulation

*Simulation Analysis*

---

**Description**

- `simulate_signature()` - Simulate signatures from signature pool.
- `simulate_catalogue()` - Simulate catalogs from signature/catalog pool.
- `simulate_catalogue_matrix()` - Simulate a bootstrapped catalog matrix.

**Usage**

```
simulate_signature(x, weights = NULL)
```

```
simulate_catalogue(x, n, weights = NULL)
```

```
simulate_catalogue_matrix(x)
```

**Arguments**

`x` a numeric vector representing a signature/catalog or matrix with rows representing signatures/samples and columns representing components.

`weights` a numeric vector for weights.

`n` an integer indicating mutation number to be generated in a catalog.

**Value**

a matrix.

**Examples**

```
# Generate a catalog
set.seed(1234)
catalog <- as.integer(table(sample(1:96, 1000, replace = TRUE)))
names(catalog) <- paste0("comp", 1:96)
# Generate a signature
sig <- catalog / sum(catalog)

# Simulate catalogs
x1 <- simulate_catalogue(catalog, 10) # 10 mutations
x1
```

```
x2 <- simulate_catalogue(catalog, 100) # 100 mutations
x2
x3 <- simulate_catalogue(catalog, 1000) # 1000 mutations
x3
# Similar with a signature
x4 <- simulate_catalogue(sig, 10) # 10 mutations
x4

# Load SBS signature
load(system.file("extdata", "toy_mutational_signature.RData",
  package = "sigminer", mustWork = TRUE
))
s <- t(sig2$Signature.norm)
# Generate a signature from multiple signatures/catalogs
s1 <- simulate_signature(s)
s1
s2 <- simulate_signature(s, weights = 1:3)
s2
# Generate a catalog from multiple signatures/catalogs
c1 <- simulate_catalogue(s, 100, weights = 1:3)
c1
```

---

subset.CopyNumber      *Subsetting CopyNumber object*

---

## Description

Subset data slot of [CopyNumber](#) object, un-selected rows will move to dropoff.segs slot, annotation slot will update in the same way.

## Usage

```
## S3 method for class 'CopyNumber'
subset(x, subset = TRUE, ...)
```

## Arguments

x	a <a href="#">CopyNumber</a> object to be subsetted.
subset	logical expression indicating rows to keep.
...	further arguments to be passed to or from other methods. Useless here.

## Value

a [CopyNumber](#) object

## Author(s)

Shixiang Wang

---

transcript.hg19

*Merged Transcript Location at Genome Build hg19*

---

**Description**

Merged Transcript Location at Genome Build hg19

**Format**

A data.table

**Source**

from GENCODE release v33.

**Examples**

```
data(transcript.hg19)
```

---

transcript.hg38

*Merged Transcript Location at Genome Build hg38*

---

**Description**

Merged Transcript Location at Genome Build hg38

**Format**

A data.table

**Source**

from GENCODE release v33.

**Examples**

```
data(transcript.hg38)
```

---

`transcript.mm10`*Merged Transcript Location at Genome Build mm10*

---

**Description**

Merged Transcript Location at Genome Build mm10

**Format**

A `data.table`

**Source**

from GENCODE release M25.

**Examples**

```
data(transcript.mm10)
```

---

`transcript.mm9`*Merged Transcript Location at Genome Build mm9*

---

**Description**

Merged Transcript Location at Genome Build mm9

**Format**

A `data.table`

**Source**

from UCSC <http://hgdownload.cse.ucsc.edu/goldenPath/mm9/database/transcriptome.txt.gz>

**Examples**

```
data(transcript.mm9)
```

---

transform\_seg\_table *Transform Copy Number Table*


---

## Description

Transform Copy Number Table

## Usage

```
transform_seg_table(
  data,
  genome_build = c("hg19", "hg38", "mm10", "mm9"),
  ref_type = c("cytoband", "gene"),
  values_fill = NA,
  values_fn = function(x, ...) {
    round(mean(x, ...))
  },
  resolution_factor = 1L
)
```

## Arguments

data	a CopyNumber object or a data.frame containing at least 'chromosome', 'start', 'end', 'segVal', 'sample' these columns.
genome_build	genome build version, used when data is a data.frame, should be 'hg19' or 'hg38'.
ref_type	annotation data type used for constructing matrix.
values_fill	Optionally, a (scalar) value that specifies what each value should be filled in with when missing. This can be a named list if you want to apply different fill values to different value columns.
values_fn	Optionally, a function applied to the value in each cell in the output. You will typically use this when the combination of id_cols and names_from columns does not uniquely identify an observation. This can be a named list if you want to apply different aggregations to different values_from columns.
resolution_factor	an integer to control the resolution. When it is 1 (default), compute frequency in each cytoband. When it is 2, use compute frequency in each half cytoband.

## Value

a data.table.

**Examples**

```
load(system.file("extdata", "toy_copynumber.RData",
  package = "sigminer", mustWork = TRUE
))
# Compute the mean segVal in each cytoband
x <- transform_seg_table(cn, resolution_factor = 1)
x
# Compute the mean segVal in each half-cytoband
x2 <- transform_seg_table(cn, resolution_factor = 2)
x2
```

---

use_color_style	<i>Set Color Style for Plotting</i>
-----------------	-------------------------------------

---

**Description**

Set Color Style for Plotting

**Usage**

```
use_color_style(
  style,
  mode = c("SBS", "copynumber", "DBS", "ID", "RS"),
  method = "Wang"
)
```

**Arguments**

style	one of 'default' and 'cosmic'.
mode	only used when the style is 'cosmic', can be one of "SBS", "copynumber", "DBS", "ID".
method	used to set a more custom palette for different methods.

**Value**

color values.

**Examples**

```
use_color_style("default")
use_color_style("cosmic")
```



# Index

- \* **bootstrap**
  - sig\_fit\_bootstrap, 112
- add\_h\_arrow, 4
- add\_labels, 5
  
- bp, 7
- bp\_attribute\_activity (bp), 7
- bp\_cluster\_iter\_list (bp), 7
- bp\_cluster\_iter\_list(), 10
- bp\_extract\_signatures, 39, 122
- bp\_extract\_signatures (bp), 7
- bp\_extract\_signatures(), 10, 11
- bp\_extract\_signatures\_iter (bp), 7
- bp\_extract\_signatures\_iter(), 10
- bp\_get\_clustered\_sigs (bp), 7
- bp\_get\_rank\_score (bp), 7
- bp\_get\_sig\_obj (bp), 7
- bp\_get\_stats (bp), 7
- bp\_show\_survey (bp), 7
- bp\_show\_survey2 (bp), 7
  
- centromeres.hg19, 14
- centromeres.hg38, 14
- centromeres.mm10, 15
- centromeres.mm9, 15
- chromsize.hg19, 16
- chromsize.hg38, 16
- chromsize.mm10, 17
- chromsize.mm9, 17
- circlize::circos.genomicHeatmap, 62
- circlize::circos.genomiclines, 66
- CN.features, 18, 41, 92, 119
- CopyNumber, 27, 52–54, 58, 61, 63, 64, 69, 118, 124
- CopyNumber (CopyNumber-class), 18
- CopyNumber-class, 18
- cosine, 19
- cowplot::save\_plot(), 74
- cytobands.hg19, 19
- cytobands.hg38, 20
- cytobands.mm10, 20
- cytobands.mm9, 21
  
- data.table::fread(), 51
  
- enrich\_component\_strand\_bias, 21
  
- facet, 90
- future::availableCores(), 119
  
- geom\_boxplot, 90
- get\_adj\_p, 22
- get\_Aneuploidy\_score, 23
- get\_bayesian\_result, 25, 100
- get\_cn\_freq\_table, 26
- get\_cn\_ploidy, 24, 26
- get\_genome\_annotation, 27
- get\_group\_comparison, 30, 74, 75
- get\_groups, 28, 74
- get\_intersect\_size, 31
- get\_pLOH\_score, 32
- get\_shannon\_diversity\_index, 33
- get\_sig\_cancer\_type\_index, 34
- get\_sig\_db, 35
- get\_sig\_exposure, 37
- get\_sig\_exposure(), 33
- get\_sig\_feature\_association, 38, 43, 89
- get\_sig\_feature\_association(), 43
- get\_sig\_rec\_similarity, 39
- get\_sig\_similarity, 6, 36, 40, 100
- get\_tidy\_association, 38, 39, 42, 88, 89
- ggpar, 90
- ggplot2::annotate, 6
- ggplot2::facet\_wrap, 79
- ggpubr::compare\_means(), 22, 75
- ggpubr::ggboxplot, 83
- ggpubr::ggviolin, 83
- ggpubr::stat\_compare\_means(), 22, 75
- ggpubr::stat\_pvalue\_manual(), 22

- group\_enrichment, [43, 78, 79](#)
- handle\_hyper\_mutation, [45](#)
- hello, [46](#)
- legend(), [74](#)
- list.files(), [51](#)
- MAF, [52, 55, 97, 118](#)
- MAF (MAF-class), [46](#)
- MAF-class, [46](#)
- maftools::read.maf, [53](#)
- mean, [77](#)
- NMF::nmf(), [108](#)
- NMF::nmfEstimateRank, [105](#)
- NMF::predict(), [29](#)
- output\_bootstrap, [47](#)
- output\_fit, [48](#)
- output\_sig, [49](#)
- output\_tally, [50](#)
- plot\_grid, [63, 65](#)
- read\_copynumber, [50, 54, 55, 119](#)
- read\_copynumber(), [53](#)
- read\_copynumber\_ascat, [52](#)
- read\_copynumber\_seqz, [53](#)
- read\_maf, [52, 53, 55](#)
- read\_sv\_as\_rs, [54, 118](#)
- read\_vcf, [55](#)
- read\_xena\_variants, [56](#)
- report\_bootstrap\_p\_value, [57, 114](#)
- same\_size\_clustering, [57](#)
- scoring, [58](#)
- show\_catalogue, [60](#)
- show\_cn\_circos, [61](#)
- show\_cn\_components, [62](#)
- show\_cn\_distribution, [63](#)
- show\_cn\_features, [65](#)
- show\_cn\_freq\_circos, [66](#)
- show\_cn\_group\_profile, [67](#)
- show\_cn\_profile, [61, 69](#)
- show\_cor, [70](#)
- show\_cosmic, [72](#)
- show\_cosmic\_sig\_profile, [36, 72](#)
- show\_group\_comparison, [74](#)
- show\_group\_comparison(), [30](#)
- show\_group\_distribution, [76](#)
- show\_group\_enrichment, [44, 78](#)
- show\_group\_mapping, [79](#)
- show\_groups, [29, 74](#)
- show\_sig\_bootstrap, [81](#)
- show\_sig\_bootstrap\_error, [83](#)
- show\_sig\_bootstrap\_error  
(show\_sig\_bootstrap), [81](#)
- show\_sig\_bootstrap\_exposure, [83, 90](#)
- show\_sig\_bootstrap\_exposure  
(show\_sig\_bootstrap), [81](#)
- show\_sig\_bootstrap\_stability, [83](#)
- show\_sig\_bootstrap\_stability  
(show\_sig\_bootstrap), [81](#)
- show\_sig\_consensusmap, [85](#)
- show\_sig\_exposure, [86](#)
- show\_sig\_feature\_corrplot, [71, 87](#)
- show\_sig\_fit, [89](#)
- show\_sig\_number\_survey (sig\_estimate),  
[103](#)
- show\_sig\_number\_survey(), [7](#)
- show\_sig\_number\_survey2, [106](#)
- show\_sig\_number\_survey2 (sig\_estimate),  
[103](#)
- show\_sig\_profile, [5, 61, 73, 91, 97](#)
- show\_sig\_profile(), [94](#)
- show\_sig\_profile\_heatmap, [93, 94](#)
- show\_sig\_profile\_loop, [93, 96](#)
- sig\_attrs (sig\_operation), [116](#)
- sig\_auto\_extract, [12, 25, 28, 37, 86, 92, 95, 96, 98, 99, 102, 106, 108, 109, 111, 113, 117, 120, 122](#)
- sig\_convert, [102](#)
- sig\_estimate, [12, 101, 103, 105–108, 120](#)
- sig\_estimate(), [105](#)
- sig\_exposure (sig\_operation), [116](#)
- sig\_extract, [7, 12, 28, 37, 85, 86, 92, 95, 96, 98, 101–103, 106, 107, 108, 109, 111, 113, 117, 120–122](#)
- sig\_extract(), [28, 29, 101](#)
- sig\_fit, [7, 28, 36, 37, 48, 74, 83, 89, 90, 101, 107, 109, 114, 116](#)
- sig\_fit(), [74](#)
- sig\_fit\_bootstrap, [57, 83, 90, 110, 111, 112, 115, 116](#)
- sig\_fit\_bootstrap\_batch, [47, 82, 83, 90, 111, 114, 115](#)
- sig\_modify\_names (sig\_operation), [116](#)

sig\_names (sig\_operation), 116  
sig\_number (sig\_operation), 116  
sig\_operation, 116  
sig\_signature (sig\_operation), 116  
sig\_tally, 21, 60, 63, 65, 92, 93, 101, 108,  
117  
sig\_tally(), 50  
sig\_unify\_extract, 121  
sigminer, 97  
sigprofiler, 98, 122  
sigprofiler\_extract, 12  
sigprofiler\_extract (sigprofiler), 98  
sigprofiler\_import (sigprofiler), 98  
simulate\_catalogue (simulation), 123  
simulate\_catalogue\_matrix (simulation),  
123  
simulate\_signature (simulation), 123  
simulated\_catalogs, 122  
simulation, 123  
stats::aov, 30  
stats::fisher.test, 30  
stats::median, 77  
stats::p.adjust, 43, 71  
stats::p.adjust(), 22  
stats::TukeyHSD, 30  
subset.CopyNumber, 124  
  
transcript.hg19, 125  
transcript.hg38, 125  
transcript.mm10, 126  
transcript.mm9, 126  
transform\_seg\_table, 127  
  
use\_color\_style, 128