

Package: sctransform (via r-universe)

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Type Package

Title Variance Stabilizing Transformations for Single Cell UMI Data

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Description A normalization method for single-cell UMI count data using a variance stabilizing transformation. The transformation is based on a negative binomial regression model with regularized parameters. As part of the same regression framework, this package also provides functions for batch correction, and data correction. See Hafemeister and Satija (2019) <[doi:10.1186/s13059-019-1874-1](https://doi.org/10.1186/s13059-019-1874-1)>, and Choudhary and Satija (2022) <[doi:10.1186/s13059-021-02584-9](https://doi.org/10.1186/s13059-021-02584-9)> for more details.

URL <https://github.com/satijalab/sctransform>

BugReports <https://github.com/satijalab/sctransform/issues>

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compare_expression	<i>Compare gene expression between two groups</i>
--------------------	---

Description

Compare gene expression between two groups

Usage

```
compare_expression(
  x,
  umi,
  group,
  val1,
  val2,
  method = "LRT",
  bin_size = 256,
  cell_attr = x$cell_attr,
  y = x$y,
  min_cells = 5,
  weighted = TRUE,
  randomize = FALSE,
  verbosity = 2,
  verbose = NULL,
  show_progress = NULL
)
```

Arguments

x	A list that provides model parameters and optionally meta data; use output of vst function
umi	A matrix of UMI counts with genes as rows and cells as columns
group	A vector indicating the groups
val1	A vector indicating the values of the group vector to treat as group 1
val2	A vector indicating the values of the group vector to treat as group 2
method	Either 'LRT' for likelihood ratio test, or 't_test' for t-test
bin_size	Number of genes that are processed between updates of progress bar
cell_attr	Data frame of cell meta data
y	Only used if method = 't_test', this is the residual matrix; default is x\$y
min_cells	A gene has to be detected in at least this many cells in at least one of the groups being compared to be tested
weighted	Balance the groups by using the appropriate weights
randomize	Boolean indicating whether to shuffle group labels - only set to TRUE when testing methods
verbosity	An integer specifying whether to show only messages (1), messages and progress bars (2) or nothing (0) while the function is running; default is 2
verbose	Deprecated; use verbosity instead
show_progress	Deprecated; use verbosity instead

Value

Data frame of results

correct	<i>Correct data by setting all latent factors to their median values and reversing the regression model</i>
---------	---

Description

Correct data by setting all latent factors to their median values and reversing the regression model

Usage

```
correct(
  x,
  data = "y",
  cell_attr = x$cell_attr,
  as_is = FALSE,
  do_round = TRUE,
  do_pos = TRUE,
  scale_factor = NA,
  verbosity = 2,
  verbose = NULL,
  show_progress = NULL
)
```

Arguments

x	A list that provides model parameters and optionally meta data; use output of vst function
data	The name of the entry in x that holds the data
cell_attr	Provide cell meta data holding latent data info
as_is	Use cell attributes as is and do not use the median; set to TRUE if you want to manually control the values of the latent factors; default is FALSE
do_round	Round the result to integers
do_pos	Set negative values in the result to zero
scale_factor	Replace all values of UMI in the regression model by this value. Default is NA which uses median of total UMI as the latent factor.
verbosity	An integer specifying whether to show only messages (1), messages and progress bars (2) or nothing (0) while the function is running; default is 2
verbose	Deprecated; use verbosity instead
show_progress	Deprecated; use verbosity instead

Value

Corrected data as UMI counts

Examples

```
vst_out <- vst(pbmc, return_cell_attr = TRUE)
umi_corrected <- correct(vst_out)
```

correct_counts	<i>Correct data by setting all latent factors to their median values and reversing the regression model</i>
----------------	---

Description

This version does not need a matrix of Pearson residuals. It takes the count matrix as input and calculates the residuals on the fly. The corrected UMI counts will be rounded to the nearest integer and negative values clipped to 0.

Usage

```
correct_counts(
  x,
  umi,
  cell_attr = x$cell_attr,
  scale_factor = NA,
  verbosity = 2,
  verbose = NULL,
  show_progress = NULL
)
```

Arguments

x	A list that provides model parameters and optionally meta data; use output of vst function
umi	The count matrix
cell_attr	Provide cell meta data holding latent data info
scale_factor	Replace all values of UMI in the regression model by this value. Default is NA which uses median of total UMI as the latent factor.
verbosity	An integer specifying whether to show only messages (1), messages and progress bars (2) or nothing (0) while the function is running; default is 2
verbose	Deprecated; use verbosity instead
show_progress	Deprecated; use verbosity instead

Value

Corrected data as UMI counts

Examples

```
vst_out <- vst(pbmc, return_cell_attr = TRUE)
umi_corrected <- correct_counts(vst_out, pbmc)
```

diff_mean_test	<i>Non-parametric differential expression test for sparse non-negative data</i>
----------------	---

Description

Non-parametric differential expression test for sparse non-negative data

Usage

```
diff_mean_test(
  y,
  group_labels,
  compare = "each_vs_rest",
  R = 99,
  log2FC_th = log2(1.2),
  mean_th = 0.05,
  cells_th = 5,
  only_pos = FALSE,
  only_top_n = NULL,
  mean_type = "geometric",
  verbosity = 1
)
```

Arguments

y	A matrix of counts; must be (or inherit from) class dgCMatrix; genes are row, cells are columns
group_labels	The group labels (e.g. cluster identities); will be converted to factor
compare	Specifies which groups to compare, see details; default is 'each_vs_rest'
R	The number of random permutations used to derive the p-values; default is 99
log2FC_th	Threshold to remove genes from testing; absolute log2FC must be at least this large for a gene to be tested; default is log2(1.2)
mean_th	Threshold to remove genes from testing; gene mean must be at least this large for a gene to be tested; default is 0.05
cells_th	Threshold to remove genes from testing; gene must be detected (non-zero count) in at least this many cells in the group with higher mean; default is 5
only_pos	Test only genes with positive fold change (mean in group 1 > mean in group2); default is FALSE

only_top_n	Test only the this number of genes from both ends of the log2FC spectrum after all of the above filters have been applied; useful to get only the top markers; only used if set to a numeric value; default is NULL
mean_type	Which type of mean to use; if 'geometric' (default) the geometric mean is used; to avoid $\log(0)$ we use \log_1p to add 1 to all counts and log-transform, calculate the arithmetic mean, and then back-transform and subtract 1 using \exp_1m ; if this parameter is set to 'arithmetic' the data is used as is
verbosity	Integer controlling how many messages the function prints; 0 is silent, 1 (default) is not

Value

Data frame of results

Details

This model-free test is applied to each gene (row) individually but is optimized to make use of the efficient sparse data representation of the input. A permutation null distribution is used to assess the significance of the observed difference in mean between two groups.

The observed difference in mean is compared against a distribution obtained by random shuffling of the group labels. For each gene every random permutation yields a difference in mean and from the population of these background differences we estimate a mean and standard deviation for the null distribution. This mean and standard deviation are used to turn the observed difference in mean into a z-score and then into a p-value. Finally, all p-values (for the tested genes) are adjusted using the Benjamini & Hochberg method (fdr). The log2FC values in the output are $\log_2(\text{mean}_1 / \text{mean}_2)$. Empirical p-values are also calculated: $\text{emp_pval} = (b + 1) / (R + 1)$ where b is the number of times the absolute difference in mean from a random permutation is at least as large as the absolute value of the observed difference in mean, R is the number of random permutations. This is an upper bound of the real empirical p-value that would be obtained by enumerating all possible group label permutations.

There are multiple ways the group comparisons can be specified based on the compare parameter. The default, 'each_vs_rest', does multiple comparisons, one per group vs all remaining cells. 'all_vs_all', also does multiple comparisons, covering all groups pairs. If compare is set to a length two character vector, e.g. `c('T-cells', 'B-cells')`, one comparison between those two groups is done. To put multiple groups on either side of a single comparison, use a list of length two. E.g. `compare = list(c('cluster1', 'cluster5'), c('cluster3'))`.

Examples

```
clustering <- 1:ncol(pbmc) %% 2
vst_out <- vst(pbmc, return_corrected_umi = TRUE)
de_res <- diff_mean_test(y = vst_out$umi_corrected, group_labels = clustering)
```

diff_mean_test_conserved

Find differentially expressed genes that are conserved across samples

Description

Find differentially expressed genes that are conserved across samples

Usage

```
diff_mean_test_conserved(
  y,
  group_labels,
  sample_labels,
  balanced = TRUE,
  compare = "each_vs_rest",
  pval_th = 1e-04,
  ...
)
```

Arguments

y	A matrix of counts; must be (or inherit from) class dgCMatrix; genes are rows, cells are columns
group_labels	The group labels (i.e. clusters or time points); will be converted to factor
sample_labels	The sample labels; will be converted to factor
balanced	Boolean, see details for explanation; default is TRUE
compare	Specifies which groups to compare, see details; currently only 'each_vs_rest' (the default) is supported
pval_th	P-value threshold used to call a gene differentially expressed when summarizing the tests per gene
...	Parameters passed to diff_mean_test

Value

Data frame of results

Details

This function calls diff_mean_test repeatedly and aggregates the results per group and gene.

If balanced is TRUE (the default), it is assumed that each sample spans multiple groups, as would be the case when merging or integrating samples from the same tissue followed by clustering. Here the group labels would be the clusters and cluster markers would have support in each sample.

If balanced is FALSE, an unbalanced design is assumed where each sample contributes to one group. An example is a time series experiment where some samples are taken from time point 1

while other samples are taken from time point 2. The time point would be the group label and the goal would be to identify differentially expressed genes between time points that are supported by many between-sample comparisons.

Output columns:

group1 Group label of the first group of cells

group2 Group label of the second group of cells; currently fixed to 'rest'

gene Gene name (from rownames of input matrix)

n_tests The number of tests this gene participated in for this group

log2FC_min,median,max Summary statistics for log2FC across the tests

mean1,2_median Median of group mean across the tests

pval_max Maximum of p-values across tests

de_tests Number of tests that showed this gene having a log2FC going in the same direction as log2FC_median and having a p-value \leq pval_th

The output is ordered by group1, -de_tests, -abs(log2FC_median), pval_max

Examples

```
clustering <- 1:ncol(pbmc) %% 2
sample_id <- 1:ncol(pbmc) %% 3
vst_out <- vst(pbmc, return_corrected_umi = TRUE)
de_res <- diff_mean_test_conserved(y = vst_out$umi_corrected,
group_labels = clustering, sample_labels = sample_id)
```

generate

Generate data from regularized models.

Description

Generate data from regularized models. This generates data from the background, i.e. no residuals are added to the simulated data. The cell attributes for the generated cells are sampled from the input with replacement.

Usage

```
generate(
  vst_out,
  genes = rownames(vst_out$model_pars_fit),
  cell_attr = vst_out$cell_attr,
  n_cells = nrow(cell_attr)
)
```

Arguments

vst_out	A list that provides model parameters and optionally meta data; use output of vst function
genes	The gene names for which to generate data; default is rownames(vst_out\$model_pars_fit)
cell_attr	Provide cell meta data holding latent data info; default is vst_out\$cell_attr
n_cells	Number of cells to generate; default is nrow(cell_attr)

Value

Generated data as dgCMatrix

Examples

```
vst_out <- vst(pbmc, return_cell_attr = TRUE)
generated_data <- generate(vst_out)
```

get_model_var	<i>Return average variance under negative binomial model</i>
---------------	--

Description

This is based on the formula $\text{var} = \mu + \mu^2 / \theta$

Usage

```
get_model_var(
  vst_out,
  cell_attr = vst_out$cell_attr,
  use_nonreg = FALSE,
  bin_size = 256,
  verbosity = 2,
  verbose = NULL,
  show_progress = NULL
)
```

Arguments

vst_out	The output of a vst run
cell_attr	Data frame of cell meta data
use_nonreg	Use the non-regularized parameter estimates; boolean; default is FALSE
bin_size	Number of genes to put in each bin (to show progress)
verbosity	An integer specifying whether to show only messages (1), messages and progress bars (2) or nothing (0) while the function is running; default is 2
verbose	Deprecated; use verbosity instead
show_progress	Deprecated; use verbosity instead

Value

A named vector of variances (the average across all cells), one entry per gene.

Examples

```
vst_out <- vst(pbmc, return_cell_attr = TRUE)
res_var <- get_model_var(vst_out)
```

get_nz_median2	<i>Get median of non zero UMIs from a count matrix</i>
----------------	--

Description

Get median of non zero UMIs from a count matrix

Usage

```
get_nz_median2(umi, genes = NULL)
```

Arguments

umi	Count matrix
genes	A vector of genes to consider for calculating the median. Default is NULL which uses all genes.

Value

A numeric value representing the median of non-zero entries from the UMI matrix

get_residuals	<i>Return Pearson or deviance residuals of regularized models</i>
---------------	---

Description

Return Pearson or deviance residuals of regularized models

Usage

```

get_residuals(
  vst_out,
  umi,
  residual_type = "pearson",
  res_clip_range = c(-sqrt(ncol(umi)), sqrt(ncol(umi))),
  min_variance = vst_out$arguments$min_variance,
  cell_attr = vst_out$cell_attr,
  bin_size = 256,
  verbosity = vst_out$arguments$verbosity,
  verbose = NULL,
  show_progress = NULL
)

```

Arguments

vst_out	The output of a vst run
umi	The UMI count matrix that will be used
residual_type	What type of residuals to return; can be 'pearson' or 'deviance'; default is 'pearson'
res_clip_range	Numeric of length two specifying the min and max values the results will be clipped to; default is c(-sqrt(ncol(umi)), sqrt(ncol(umi)))
min_variance	Lower bound for the estimated variance for any gene in any cell when calculating pearson residual; default is vst_out\$arguments\$min_variance
cell_attr	Data frame of cell meta data
bin_size	Number of genes to put in each bin (to show progress)
verbosity	An integer specifying whether to show only messages (1), messages and progress bars (2) or nothing (0) while the function is running; default is 2
verbose	Deprecated; use verbosity instead
show_progress	Deprecated; use verbosity instead

Value

A matrix of residuals

Examples

```

vst_out <- vst(pbmc, return_cell_attr = TRUE)
pearson_res <- get_residuals(vst_out, pbmc)
deviance_res <- get_residuals(vst_out, pbmc, residual_type = 'deviance')

```

get_residual_var	<i>Return variance of residuals of regularized models</i>
------------------	---

Description

This never creates the full residual matrix and can be used to determine highly variable genes.

Usage

```
get_residual_var(
  vst_out,
  umi,
  residual_type = "pearson",
  res_clip_range = c(-sqrt(ncol(umi)), sqrt(ncol(umi))),
  min_variance = vst_out$arguments$min_variance,
  cell_attr = vst_out$cell_attr,
  bin_size = 256,
  verbosity = vst_out$arguments$verbosity,
  verbose = NULL,
  show_progress = NULL
)
```

Arguments

vst_out	The output of a vst run
umi	The UMI count matrix that will be used
residual_type	What type of residuals to return; can be 'pearson' or 'deviance'; default is 'pearson'
res_clip_range	Numeric of length two specifying the min and max values the residuals will be clipped to; default is c(-sqrt(ncol(umi)), sqrt(ncol(umi)))
min_variance	Lower bound for the estimated variance for any gene in any cell when calculating pearson residual; default is vst_out\$arguments\$min_variance
cell_attr	Data frame of cell meta data
bin_size	Number of genes to put in each bin (to show progress)
verbosity	An integer specifying whether to show only messages (1), messages and progress bars (2) or nothing (0) while the function is running; default is 2
verbose	Deprecated; use verbosity instead
show_progress	Deprecated; use verbosity instead

Value

A vector of residual variances (after clipping)

Examples

```
vst_out <- vst(pbmc, return_cell_attr = TRUE)
res_var <- get_residual_var(vst_out, pbmc)
```

is_outlier	<i>Identify outliers</i>
------------	--------------------------

Description

Identify outliers

Usage

```
is_outlier(y, x, th = 10)
```

Arguments

y	Dependent variable
x	Independent variable
th	Outlier score threshold

Value

Boolean vector

make.sparse	<i>Convert a given matrix to dgCMatrix</i>
-------------	--

Description

Convert a given matrix to dgCMatrix

Usage

```
make.sparse(mat)
```

Arguments

mat	Input matrix
-----	--------------

Value

A dgCMatrix

pbmc	<i>Peripheral Blood Mononuclear Cells (PBMCs)</i>
------	---

Description

UMI counts for a subset of cells freely available from 10X Genomics

Usage

```
pbmc
```

Format

A sparse matrix (dgCMatrix, see Matrix package) of molecule counts. There are 914 rows (genes) and 283 columns (cells). This is a downsampled version of a 3K PBMC dataset available from 10x Genomics.

Source

<https://support.10xgenomics.com/single-cell-gene-expression/datasets/1.1.0/pbmc3k>

plot_model	<i>Plot observed UMI counts and model</i>
------------	---

Description

Plot observed UMI counts and model

Usage

```
plot_model(  
  x,  
  umi,  
  goi,  
  x_var = x$arguments$latent_var[1],  
  cell_attr = x$cell_attr,  
  do_log = TRUE,  
  show_fit = TRUE,  
  show_nr = FALSE,  
  plot_residual = FALSE,  
  batches = NULL,  
  as_poisson = FALSE,  
  arrange_vertical = TRUE,  
  show_density = FALSE,  
  gg_cmds = NULL  
)
```

Arguments

x	The output of a vst run
umi	UMI count matrix
goi	Vector of genes to plot
x_var	Cell attribute to use on x axis; will be taken from x\$arguments\$latent_var[1] by default
cell_attr	Cell attributes data frame; will be taken from x\$cell_attr by default
do_log	Log10 transform the UMI counts in plot
show_fit	Show the model fit
show_nr	Show the non-regularized model (if available)
plot_residual	Add panels for the Pearson residuals
batches	Manually specify a batch variable to break up the model plot in segments
as_poisson	Fix model parameter theta to Inf, effectively showing a Poisson model
arrange_vertical	Stack individual ggplot objects or place side by side
show_density	Draw 2D density lines over points
gg_cmds	Additional ggplot layer commands

Value

A ggplot object

Examples

```
vst_out <- vst(pbmc, return_cell_attr = TRUE)
plot_model(vst_out, pbmc, 'EMC4')
```

plot_model_pars	<i>Plot estimated and fitted model parameters</i>
-----------------	---

Description

Plot estimated and fitted model parameters

Usage

```
plot_model_pars(  
  vst_out,  
  xaxis = "gmean",  
  show_theta = FALSE,  
  show_var = FALSE,  
  verbosity = 2,  
  verbose = NULL,  
  show_progress = NULL  
)
```

Arguments

vst_out	The output of a vst run
xaxis	Variable to plot on X axis; default is "gmean"
show_theta	Whether to show the theta parameter; default is FALSE (only the overdispersion factor is shown)
show_var	Whether to show the average model variance; default is FALSE
verbosity	An integer specifying whether to show only messages (1), messages and progress bars (2) or nothing (0) while the function is running; default is 2
verbose	Deprecated; use verbosity instead
show_progress	Deprecated; use verbosity instead

Value

A ggplot object

Examples

```
vst_out <- vst(pbmc, return_gene_attr = TRUE)  
plot_model_pars(vst_out)
```

robust_scale	<i>Robust scale using median and mad</i>
--------------	--

Description

Robust scale using median and mad

Usage

```
robust_scale(x)
```

Arguments

x Numeric

Value

Numeric

robust_scale_binned *Robust scale using median and mad per bin*

Description

Robust scale using median and mad per bin

Usage

```
robust_scale_binned(y, x, breaks)
```

Arguments

y Numeric vector
x Numeric vector
breaks Numeric vector of breaks

Value

Numeric vector of scaled score

row_gmean *Geometric mean per row*

Description

Geometric mean per row

Usage

```
row_gmean(x, eps = 1)
```

Arguments

x matrix of class matrix or dgMatrix
eps small value to add to x to avoid log(0); default is 1

Value

geometric means

row_var	<i>Variance per row</i>
---------	-------------------------

Description

Variance per row

Usage

```
row_var(x)
```

Arguments

x matrix of class matrix or dgCMatix

Value

variances

smooth_via_pca	<i>Smooth data by PCA</i>
----------------	---------------------------

Description

Perform PCA, identify significant dimensions, and reverse the rotation using only significant dimensions.

Usage

```
smooth_via_pca(
  x,
  elbow_th = 0.025,
  dims_use = NULL,
  max_pc = 100,
  do_plot = FALSE,
  scale. = FALSE
)
```

Arguments

x A data matrix with genes as rows and cells as columns

elbow_th The fraction of PC sdev drop that is considered significant; low values will lead to more PCs being used

dims_use Directly specify PCs to use, e.g. 1:10

max_pc Maximum number of PCs computed

do_plot	Plot PC sdev and sdev drop
scale.	Boolean indicating whether genes should be divided by standard deviation after centering and prior to PCA

Value

Smoothed data

Examples

```
vst_out <- vst(pbmc)
y_smooth <- smooth_via_pca(vst_out$y, do_plot = TRUE)
```

umify	<i>Quantile normalization of cell-level data to match typical UMI count data</i>
-------	--

Description

Quantile normalization of cell-level data to match typical UMI count data

Usage

```
umify(counts)
```

Arguments

counts	A matrix of class dgCMatrix with genes as rows and columns as cells
--------	---

Value

A UMI-fied count matrix

Details

sctransform::vst operates under the assumption that gene counts approximately follow a Negative Binomial distribution. For UMI-based data that seems to be the case, however, non-UMI data does not behave in the same way. In some cases it might be better to to apply a transformation to such data to make it look like UMI data. This function applies such a transformation function.

Cells in the input matrix are processed independently. For each cell the non-zero data is transformed to quantile values. Based on the number of genes detected a smooth function is used to predict the UMI-like counts.

The functions have be trained on various public data sets and come as part of the package (see umify_data data set in this package).

Examples

```
silly_example <- umify(pbmc)
```

umify_data

Transformation functions for umify

Description

The functions have been trained on various public data sets and relate quantile values to log-counts. Here the expected values at various points are given.

Usage

```
umify_data
```

Format

A list of length two. The first element is a data frame with group, quantile and log-counts values. The second element is a vector of breaks to be used with cut to group observations.

vst

Variance stabilizing transformation for UMI count data

Description

Apply variance stabilizing transformation to UMI count data using a regularized Negative Binomial regression model. This will remove unwanted effects from UMI data and return Pearson residuals. Uses `future_lapply`; you can set the number of cores it will use to `n` with `plan(strategy = "multicore", workers = n)`. If `n_genes` is set, only a (somewhat-random) subset of genes is used for estimating the initial model parameters. For details see doi: [10.1186/s1305901918741](https://doi.org/10.1186/s1305901918741).

Usage

```
vst(
  umi,
  cell_attr = NULL,
  latent_var = c("log_umi"),
  batch_var = NULL,
  latent_var_nonreg = NULL,
  n_genes = 2000,
  n_cells = NULL,
  method = "poisson",
  do_regularize = TRUE,
  theta_regularization = "od_factor",
```

```

res_clip_range = c(-sqrt(ncol(umi)), sqrt(ncol(umi))),
bin_size = 500,
min_cells = 5,
residual_type = "pearson",
return_cell_attr = FALSE,
return_gene_attr = TRUE,
return_corrected_umi = FALSE,
min_variance = -Inf,
bw_adjust = 3,
gmean_eps = 1,
theta_estimation_fun = "theta.ml",
theta_given = NULL,
exclude_poisson = FALSE,
use_geometric_mean = TRUE,
use_geometric_mean_offset = FALSE,
fix_intercept = FALSE,
fix_slope = FALSE,
scale_factor = NA,
vst.flavor = NULL,
verbosity = 2,
verbose = NULL,
show_progress = NULL
)

```

Arguments

<code>umi</code>	A matrix of UMI counts with genes as rows and cells as columns
<code>cell_attr</code>	A data frame containing the dependent variables; if omitted a data frame with <code>umi</code> and <code>gene</code> will be generated
<code>latent_var</code>	The independent variables to regress out as a character vector; must match column names in <code>cell_attr</code> ; default is <code>c("log_umi")</code>
<code>batch_var</code>	The dependent variables indicating which batch a cell belongs to; no batch interaction terms used if omitted
<code>latent_var_nonreg</code>	The non-regularized dependent variables to regress out as a character vector; must match column names in <code>cell_attr</code> ; default is <code>NULL</code>
<code>n_genes</code>	Number of genes to use when estimating parameters (default uses 2000 genes, set to <code>NULL</code> to use all genes)
<code>n_cells</code>	Number of cells to use when estimating parameters (default uses all cells)
<code>method</code>	Method to use for initial parameter estimation; one of <code>'poisson'</code> , <code>'qpoisson'</code> , <code>'nb_fast'</code> , <code>'nb'</code> , <code>'nb_theta_given'</code> , <code>'glmGamPoi'</code> , <code>'offset'</code> , <code>'offset_shared_theta_estimate'</code> , <code>'glmGamPoi_offset'</code> ; default is <code>'poisson'</code>
<code>do_regularize</code>	Boolean that, if set to <code>FALSE</code> , will bypass parameter regularization and use all genes in first step (ignoring <code>n_genes</code>); default is <code>FALSE</code>
<code>theta_regularization</code>	Method to use to regularize theta; use <code>'log_theta'</code> for the behavior prior to version 0.3; default is <code>'od_factor'</code>

res_clip_range	Numeric of length two specifying the min and max values the results will be clipped to; default is <code>c(-sqrt(ncol(umi)), sqrt(ncol(umi)))</code>
bin_size	Number of genes to process simultaneously; this will determine how often the progress bars are updated and how much memory is being used; default is 500
min_cells	Only use genes that have been detected in at least this many cells; default is 5
residual_type	What type of residuals to return; can be 'pearson', 'deviance', or 'none'; default is 'pearson'
return_cell_attr	Make cell attributes part of the output; default is FALSE
return_gene_attr	Calculate gene attributes and make part of output; default is TRUE
return_corrected_umi	If set to TRUE output will contain corrected UMI matrix; see <code>correct</code> function
min_variance	Lower bound for the estimated variance for any gene in any cell when calculating pearson residual; one of 'umi_median', 'model_median', 'model_mean' or a numeric. default is -Inf. When set to 'umi_median' uses $(\text{median of non-zero UMIs} / 5)^2$ as the minimum variance so that a median UMI (often 1) results in a maximum pearson residual of 5. When set to 'model_median' or 'model_mean' uses the mean/median of the model estimated μ per gene as the minimum_variance.#'
bw_adjust	Kernel bandwidth adjustment factor used during regularization; factor will be applied to output of <code>bw.SJ</code> ; default is 3
gmean_eps	Small value added when calculating geometric mean of a gene to avoid $\log(0)$; default is 1
theta_estimation_fun	Character string indicating which method to use to estimate theta (when method = poisson); default is 'theta.ml', but 'theta.mm' seems to be a good and fast alternative
theta_given	If method is set to <code>nb_theta_given</code> , this should be a named numeric vector of fixed theta values for the genes; if method is <code>offset</code> , this should be a single value; default is NULL
exclude_poisson	Exclude poisson genes (i.e. $\mu < 0.001$ or $\mu > \text{variance}$) from regularization; default is FALSE
use_geometric_mean	Use geometric mean instead of arithmetic mean for all calculations ; default is TRUE
use_geometric_mean_offset	Use geometric mean instead of arithmetic mean in the offset model; default is FALSE
fix_intercept	Fix intercept as defined in the offset model; default is FALSE
fix_slope	Fix slope to $\log(10)$ (equivalent to using library size as an offset); default is FALSE
scale_factor	Replace all values of UMI in the regression model by this value instead of the median UMI; default is NA

<code>vst.flavor</code>	When set to 'v2' sets <code>method = glmGamPoi_offset</code> , <code>n_cells=2000</code> , and <code>exclude_poisson = TRUE</code> which causes the model to learn <code>theta</code> and <code>intercept</code> only besides excluding poisson genes from learning and regularization; default is <code>NULL</code> which uses the original <code>sctransform</code> model
<code>verbosity</code>	An integer specifying whether to show only messages (1), messages and progress bars (2) or nothing (0) while the function is running; default is 2
<code>verbose</code>	Deprecated; use <code>verbosity</code> instead
<code>show_progress</code>	Deprecated; use <code>verbosity</code> instead

Value

A list with components

<code>y</code>	Matrix of transformed data, i.e. Pearson residuals, or deviance residuals; empty if <code>residual_type = 'none'</code>
<code>umi_corrected</code>	Matrix of corrected UMI counts (optional)
<code>model_str</code>	Character representation of the model formula
<code>model_pars</code>	Matrix of estimated model parameters per gene (<code>theta</code> and regression coefficients)
<code>model_pars_outliers</code>	Vector indicating whether a gene was considered to be an outlier
<code>model_pars_fit</code>	Matrix of fitted / regularized model parameters
<code>model_str_nonreg</code>	Character representation of model for non-regularized variables
<code>model_pars_nonreg</code>	Model parameters for non-regularized variables
<code>genes_log_gmean_step1</code>	log-geometric mean of genes used in initial step of parameter estimation
<code>cells_step1</code>	Cells used in initial step of parameter estimation
<code>arguments</code>	List of function call arguments
<code>cell_attr</code>	Data frame of cell meta data (optional)
<code>gene_attr</code>	Data frame with gene attributes such as mean, detection rate, etc. (optional)
<code>times</code>	Time stamps at various points in the function

Details

In the first step of the algorithm, per-gene glm model parameters are learned. This step can be done on a subset of genes and/or cells to speed things up. If `method` is set to 'poisson', a poisson regression is done and the negative binomial `theta` parameter is estimated using the response residuals in `theta_estimation_fun`. If `method` is set to 'qpoisson', coefficients and overdispersion (`phi`) are estimated by quasi poisson regression and `theta` is estimated based on `phi` and the mean fitted value - this is currently the fastest method with results very similar to 'glmGamPoi' If `method` is set to 'nb_fast', coefficients and `theta` are estimated as in the 'poisson' method, but coefficients are then re-estimated using a proper negative binomial model in a second call to `glm` with `family = MASS::negative.binomial(theta = theta)`. If `method` is set to 'nb', coefficients and `theta` are

estimated by a single call to `MASS::glm.nb`. If method is set to `'glmGamPoi'`, coefficients and theta are estimated by a single call to `glmGamPoi::glm_gp`.

A special case is `method = 'offset'`. Here no regression parameters are learned, but instead an offset model is assumed. The latent variable is set to `log_umi` and a fixed slope of $\log(10)$ is used (offset). The intercept is given by $\log(\text{gene_mean}) - \log(\text{avg_cell_umi})$. See Lause et al. doi: [10.1186/s13059021024517](https://doi.org/10.1186/s13059021024517) for details. Theta is set to 100 by default, but can be changed using the `theta_given` parameter (single numeric value). If the offset method is used, the following parameters are overwritten: `cell_attr <- NULL`, `latent_var <- c('log_umi')`, `batch_var <- NULL`, `latent_var_nonreg <- NULL`, `n_genes <- NULL`, `n_cells <- NULL`, `do_regularize <- FALSE`. Further, `method = 'offset_shared_theta_estimate'` exists where the 250 most highly expressed genes with detection rate of at least 0.5 are used to estimate a theta that is then shared across all genes. Thetas are estimated per individual gene using 5000 randomly selected cells. The final theta used for all genes is then the average.

Examples

```
vst_out <- vst(pbmc)
```

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