

# Package ‘dreamlet’

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**Type** Package

**Title** Scalable differential expression analysis of single cell transcriptomics datasets with complex study designs

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**Description** Recent advances in single cell/nucleus transcriptomic technology has enabled collection of cohort-scale datasets to study cell type specific gene expression differences associated disease state, stimulus, and genetic regulation. The scale of these data, complex study designs, and low read count per cell mean that characterizing cell type specific molecular mechanisms requires a user-friendly, purpose-build analytical framework. We have developed the dreamlet package that applies a pseudobulk approach and fits a regression model for each gene and cell cluster to test differential expression across individuals associated with a trait of interest. Use of precision-weighted linear mixed models enables accounting for repeated measures study designs, high dimensional batch effects, and varying sequencing depth or observed cells per biosample.

**VignetteBuilder** knitr

**License** Artistic-2.0

**Encoding** UTF-8

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**BugReports** <https://github.com/DiseaseNeurogenomics/dreamlet/issues>

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---

aggregateNonCountSignal

*Aggregation of single-cell signals*

---

## Description

Aggregation of single-cell to pseudobulk data for non-count data.

## Usage

```
aggregateNonCountSignal(
  sce,
  assay = NULL,
  sample_id = NULL,
  cluster_id = NULL,
  min.cells = 10,
  min.signal = 0.01,
  min.samples = 4,
  min.prop = 0.4,
  verbose = TRUE,
```

```
BPPARAM = SerialParam(progressbar = verbose)
)
```

### Arguments

|             |                                                                                                                                          |
|-------------|------------------------------------------------------------------------------------------------------------------------------------------|
| sce         | a <a href="#">SingleCellExperiment</a> .                                                                                                 |
| assay       | character string specifying the assay slot to use as input data. Defaults to the 1st available (assayNames(x)[1]).                       |
| sample_id   | character string specifying which variable to use as sample id                                                                           |
| cluster_id  | character string specifying which variable to use as cluster id                                                                          |
| min.cells   | minimum number of observed cells for a sample to be included in the analysis                                                             |
| min.signal  | minimum signal value for a gene to be considered expressed in a sample. Proper value for this cutoff depends on the type of signal value |
| min.samples | minimum number of samples passing cutoffs for cell cluster to be retained                                                                |
| min.prop    | minimum proportion of retained samples with non-zero counts for a gene to be                                                             |
| verbose     | logical. Should information on progress be reported?                                                                                     |
| BPPARAM     | a <a href="#">BiocParallelParam</a> object specifying how aggregation should be parallelized.                                            |

### Details

The dreamlet workflow can also be applied to non-count data. In this case, a signal is averaged across all cells from a given sample and cell type. Here `aggregateNonCountSignal()` performs the roles of `aggregateToPseudoBulk()` followed by `processAssays()` but using non-count data.

For each cell cluster, samples with at least `min.cells` are retained. Only clusters with at least `min.samples` retained samples are kept. Features are retained if they have at least `min.signal` in at least `min.prop` fraction of the samples.

The precision of a measurement is the inverse of its sampling variance. The precision weights are computed as  $1/\text{sem}^2$ , where  $\text{sem} = \text{sd}(\text{signal}) / \sqrt{n}$ , `signal` stores the values averaged across cells, and `n` is the number of cells.

### Value

a `dreamletProcessedData` object

### Examples

```
library(muscat)
library(SingleCellExperiment)

data(example_sce)

# create pseudobulk for each sample and cell cluster
# using non-count signal
pb.signal <- aggregateNonCountSignal(example_sce,
  assay = "logcounts",
  cluster_id = "cluster_id",
  sample_id = "sample_id",
  verbose = FALSE
)

# Differential expression analysis within each assay,
# evaluated on the voom normalized data
res.dl <- dreamlet(pb.signal, ~group_id)
```

---

aggregateToPseudoBulk *Aggregation of single-cell to pseudobulk data*

---

## Description

Aggregation of single-cell to pseudobulk data. Adapted from `muscat::aggregateData` and has same syntax and results. But can be much faster for `SingleCellExperiment` backed by H5AD files using on-disk storage.

## Usage

```
aggregateToPseudoBulk(
  x,
  assay = NULL,
  sample_id = NULL,
  cluster_id = NULL,
  fun = c("sum", "mean", "median", "prop.detected", "num.detected", "sem", "number"),
  scale = FALSE,
  verbose = TRUE,
  BPPARAM = SerialParam(progressbar = verbose),
  checkValues = TRUE,
  h5adBlockSizes = 1e+09
)
```

## Arguments

|                             |                                                                                                                                                      |
|-----------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------|
| <code>x</code>              | a <a href="#">SingleCellExperiment</a> .                                                                                                             |
| <code>assay</code>          | character string specifying the assay slot to use as input data. Defaults to the 1st available ( <code>assayNames(x)[1]</code> ).                    |
| <code>sample_id</code>      | character string specifying which variable to use as sample id                                                                                       |
| <code>cluster_id</code>     | character string specifying which variable to use as cluster id                                                                                      |
| <code>fun</code>            | a character string. Specifies the function to use as summary statistic. Passed to <code>summarizeAssayByGroup2</code> .                              |
| <code>scale</code>          | logical. Should pseudo-bulks be scaled with the effective library size & multiplied by 1M?                                                           |
| <code>verbose</code>        | logical. Should information on progress be reported?                                                                                                 |
| <code>BPPARAM</code>        | a <a href="#">BiocParallelParam</a> object specifying how aggregation should be parallelized.                                                        |
| <code>checkValues</code>    | logical. Should we check that signal values are positive integers?                                                                                   |
| <code>h5adBlockSizes</code> | set the automatic block size block size (in bytes) for <code>DelayedArray</code> to read an H5AD file. Larger values use more memory but are faster. |

## Details

Adapted from `muscat::aggregateData` and has similar syntax and same results. This is much faster for `SingleCellExperiment` backed by H5AD files using `DelayedMatrix` because this summarizes counts using [DelayedMatrixStats](#). But this function also includes optimizations for `sparseMatrix` used by [Seurat](#) by using `sparseMatrixStats`.

Keeps variables from `colData()` that are constant within `sample_id`. For example, sex will be constant for all cells from the same `sample_id`, so it is retained as a variable in the pseudobulk result. But number of expressed genes varies across cells within each `sample_id`, so it is dropped from `colData()`. Instead the mean value per cell type is stored in `metadata(pb)$aggr_means`, and these can be included in regression formulas downstream. In that case, the value of the covariates used per sample will depend on the cell type analyzed.

## Value

a `SingleCellExperiment`.

Aggregation parameters (`assay`, `by`, `fun`, `scaled`) are stored in `metadata()``$aggr_pars`, where `by = c(cluster_id, sample_id)`. The number of cells that were aggregated are accessible in `int_colData()``$n_cells`.

## Author(s)

Gabriel Hoffman, Helena L Crowell & Mark D Robinson

## References

Crowell, HL, Sonesson, C, Germain, P-L, Calini, D, Collin, L, Raposo, C, Malhotra, D & Robinson, MD: Muscat detects subpopulation-specific state transitions from multi-sample multi-condition single-cell transcriptomics data. *Nature Communications* **11(1):6077** (2020). doi: <https://doi.org/10.1038/s41467-020-19894-4>

## Examples

```
library(muscat)
library(SingleCellExperiment)

data(example_sce)

# create pseudobulk for each sample and cell cluster
pb <- aggregateToPseudoBulk(example_sce,
  assay = "counts",
  cluster_id = "cluster_id",
  sample_id = "sample_id",
  verbose = FALSE
)

# pseudobulk data from each cell type
# is stored as its own assay
pb

# aggregate by cluster only,
# collapsing all samples into the same pseudobulk
pb2 <- aggregateToPseudoBulk(example_sce,
  cluster_id = "cluster_id",
  verbose = FALSE)

pb2
#
```

aggregateVar

*Per-sample variance of single-cell counts***Description**

Aggregation function for single-cell log-normalized counts to calculate per-sample variance for dreamlet.

**Usage**

```
aggregateVar(
  sce,
  assay = NULL,
  cluster_id = NULL,
  sample_id = NULL,
  min.cells = 10,
  min.var = 0.01,
  min.samples = 4,
  min.prop = 0.4,
  verbose = TRUE,
  BPPARAM = SerialParam(progressbar = verbose)
)
```

**Arguments**

|             |                                                                                                                    |
|-------------|--------------------------------------------------------------------------------------------------------------------|
| sce         | a <a href="#">SingleCellExperiment</a> .                                                                           |
| assay       | character string specifying the assay slot to use as input data. Defaults to the 1st available (assayNames(x)[1]). |
| cluster_id  | character string specifying which variable to use as cluster id                                                    |
| sample_id   | character string specifying which variable to use as sample id                                                     |
| min.cells   | minimum number of observed cells for a sample to be included in the analysis                                       |
| min.var     | minimum variance for a gene to be considered expressed in a sample                                                 |
| min.samples | minimum number of samples passing cutoffs for cell cluster to be retained                                          |
| min.prop    | minimum proportion of retained samples with non-zero counts for a gene to be                                       |
| verbose     | logical. Should information on progress be reported?                                                               |
| BPPARAM     | a <a href="#">BiocParallelParam</a> object specifying how aggregation should be parallelized.                      |

**Details**

The dreamlet workflow can also be applied to model gene expression variance. In this case, a per-sample per-gene variance is calculated across all cells from a given sample and cell type. Here `aggregateVar()` performs the roles of `aggregateToPseudoBulk()` followed by `processAssays()` but using log-normalized count data.

For each cell cluster, samples with at least `min.cells` are retained. Only clusters with at least `min.samples` retained samples are kept. Features are retained if they have at least `min.var` in at least `min.prop` fraction of the samples.

The precision of a measurement is the inverse of its sampling variance. The precision weights are computed as  $1/\text{sem}^2$ , where  $\text{sem} = \text{sd} / \sqrt{n}$  and  $n$  is the number of cells.

**Value**

a dreamletProcessedData object

**Examples**

```
library(muscat)
library(SingleCellExperiment)

data(example_sce)

# Compute variance for each sample and cell cluster
pbVar <- aggregateVar(example_sce,
  assay = "counts",
  cluster_id = "cluster_id",
  sample_id = "sample_id",
  verbose = FALSE
)
```

---

|                   |                                                          |
|-------------------|----------------------------------------------------------|
| as.dreamletResult | <i>Convert list of regression fits to dreamletResult</i> |
|-------------------|----------------------------------------------------------|

---

**Description**

Convert list of regression fits to dreamletResult for downstream analysis

**Usage**

```
as.dreamletResult(fitList, df_details = NULL)
```

**Arguments**

|            |                                     |
|------------|-------------------------------------|
| fitList    | list of regression fit with dream() |
| df_details | data.frame storing assay details    |

**Details**

Useful for combining multiple runs of dreamletCompareClusters() into a single dreamletResult for downstream analysis

**Value**

object of class dreamletResult

**Examples**

```
library(muscat)
library(SingleCellExperiment)

data(example_sce)

# create pseudobulk for each sample and cell cluster
pb <- aggregateToPseudoBulk(example_sce,
  assay = "counts",
```

```

    cluster_id = "cluster_id",
    sample_id = "sample_id",
    verbose = FALSE
  )

  # first comparison
  ct.pairs <- c("B cells", "CD14+ Monocytes")
  fit <- dreamletCompareClusters(pb, ct.pairs, method = "fixed")

  # second comparison
  ct.pairs2 <- c("B cells", "CD8 T cells")
  fit2 <- dreamletCompareClusters(pb, ct.pairs2, method = "fixed")

  # Make a list storing each result with a meaningful name
  fitList <- list()

  id <- paste0("[", ct.pairs[1], "]-vs-", ct.pairs[2], "]")
  fitList[[id]] <- fit

  id <- paste0("[", ct.pairs2[1], "]-vs-", ct.pairs2[2], "]")
  fitList[[id]] <- fit2

  # create a dreamletResult form this list
  res.compare <- as.dreamletResult(fitList)
  res.compare

```

---

assay,dreamletResult,ANY-method

*Get assay*


---

## Description

Get assay

Get assay

Get assays by name

## Usage

```

## S4 method for signature 'dreamletResult,ANY'
assay(x, i, withDimnames = TRUE, ...)

## S4 method for signature 'dreamletProcessedData,ANY'
assay(x, i, withDimnames = TRUE, ...)

## S4 method for signature 'vpDF,ANY'
assay(x, i, withDimnames = TRUE, ...)

```

## Arguments

|   |                                                     |
|---|-----------------------------------------------------|
| x | vpDF object                                         |
| i | number indicating index, or string indicating assay |

withDimnames    not used  
...              other arguments

**Value**

return ith assay

---

assayNames,dreamletResult-method  
*Get assayNames*

---

**Description**

Get assayNames  
Get assayNames  
Get assayNames

**Usage**

```
## S4 method for signature 'dreamletResult'  
assayNames(x, ...)  
  
## S4 method for signature 'dreamletProcessedData'  
assayNames(x, ...)  
  
## S4 method for signature 'vpDF'  
assayNames(x, ...)
```

**Arguments**

x                      vpDF object  
...                    additional arguments

**Value**

array of assay names

---

buildClusterTreeFromPB  
*Hierarchical clustering on cell types from pseudobulk*

---

**Description**

Perform hierarchical clustering on cell types from pseudobulk by aggregating read counts from each cell type.

**Usage**

```
buildClusterTreeFromPB(
  pb,
  method = c("complete", "ward.D", "single", "average", "mcquitty", "median", "centroid",
    "ward.D2"),
  dist.method = c("euclidean", "maximum", "manhattan", "canberra", "binary", "minkowski"),
  assays = assayNames(pb)
)
```

**Arguments**

|             |                                                                            |
|-------------|----------------------------------------------------------------------------|
| pb          | SingleCellObject storing pseudobulk for each cell type in in assay() field |
| method      | clustering method for hclust()                                             |
| dist.method | distance metric                                                            |
| assays      | which assays to include                                                    |

**Value**

hierarchical clustering object of class hclust

**Examples**

```
library(muscat)
library(SingleCellExperiment)

data(example_sce)

# create pseudobulk for each sample and cell cluster
pb <- aggregateToPseudoBulk(example_sce,
  assay = "counts",
  cluster_id = "cluster_id",
  sample_id = "sample_id",
  verbose = FALSE
)

# Hierarchical clustering of cell types
hcl <- buildClusterTreeFromPB(pb)

plot(hcl)
```

---

cellCounts

*Extract cell counts*


---

**Description**

Extract matrix of cell counts from SingleCellExperiment

**Usage**

```
cellCounts(x)
```

**Arguments**

x                      a SingleCellExperiment

**Value**

matrix of cell counts with samples as rows and cell types as columns

**See Also**

computeCellCounts()

**Examples**

```
library(muscat)
library(SingleCellExperiment)

data(example_sce)

# create pseudobulk for each sample and cell cluster
pb <- aggregateToPseudoBulk(example_sce,
  assay = "counts",
  cluster_id = "cluster_id",
  sample_id = "sample_id",
  verbose = FALSE
)

# get matrix of cell counts for each sample
cellCounts(pb)
```

---

cellSpecificityValues-class

*Class cellSpecificityValues*

---

**Description**

Class cellSpecificityValues cell type specificity values for each gene and cell type

**Value**

none

---

|                     |                                                     |
|---------------------|-----------------------------------------------------|
| cellTypeSpecificity | <i>Get cell type specificity of gene expression</i> |
|---------------------|-----------------------------------------------------|

---

**Description**

For each gene, compute fraction of overall expression attributable to each cell type

**Usage**

```
cellTypeSpecificity(pb, ...)
```

**Arguments**

|     |                                                                          |
|-----|--------------------------------------------------------------------------|
| pb  | SingleCellExperiment of pseudobulk data where easy assay is a cell type. |
| ... | other arguments passed to <code>edgeR::calcNormFactors()</code>          |

**Details**

Sum counts for each cell type, and compute the fraction of counts-per-million attributable to each cell type for each gene

**Value**

matrix of the fraction of expression attributable to each cell type for each gene.

**Examples**

```
library(muscat)
library(SingleCellExperiment)

data(example_sce)

# create pseudobulk for each sample and cell cluster
pb <- aggregateToPseudoBulk(example_sce,
  assay = "counts",
  cluster_id = "cluster_id",
  sample_id = "sample_id",
  verbose = FALSE
)

# Compute cell type specificity of each gene
df <- cellTypeSpecificity(pb)

# Violin plot of specificity scores for each cell type
# Dashed line indicates genes that are equally expressed
# across all cell types. For K cell types, this is 1/K
plotViolin(df)

# Compute the maximum specificity score for each gene
scoreMax <- apply(df, 1, max)
head(scoreMax)

# For each cell type, get most specific gene
```

```

genes <- rownames(df)[apply(df, 2, which.max)]

# Barplot of 5 genes
plotPercentBars(df, genes = genes)

# heatmap of 5 genes that are most cell type specific
dreamlet::plotHeatmap(df, genes = genes)

```

---

|              |                                     |
|--------------|-------------------------------------|
| checkFormula | <i>Check variables in a formula</i> |
|--------------|-------------------------------------|

---

### Description

Check that variables in formula are present in the data

### Usage

```
checkFormula(formula, data)
```

### Arguments

|         |                                             |
|---------|---------------------------------------------|
| formula | formula of variables to check               |
| data    | data.frame storing variables in the formula |

### Value

If formula is valid, return TRUE. Else throw error

### Examples

```

# Valid formula
dreamlet:::checkFormula(~speed, cars)

# Not valid formula
# dreamlet:::checkFormula( ~ speed + a, cars)

```

---

|           |                              |
|-----------|------------------------------|
| coefNames | <i>Get coefficient names</i> |
|-----------|------------------------------|

---

### Description

Get coefficient names

### Usage

```
coefNames(obj)

## S4 method for signature 'dreamletResult'
coefNames(obj)
```

**Arguments**

obj                    A dreamletResult object

**Value**

array storing names of coefficients

**Examples**

```
library(muscat)
library(SingleCellExperiment)

data(example_sce)

# create pseudobulk for each sample and cell cluster
pb <- aggregateToPseudoBulk(example_sce,
  assay = "counts",
  cluster_id = "cluster_id",
  sample_id = "sample_id",
  verbose = FALSE
)

# voom-style normalization
res.proc <- processAssays(pb, ~group_id)

# Differential expression analysis within each assay,
# evaluated on the voom normalized data
res.dl <- dreamlet(res.proc, ~group_id)

# show coefficients estimated for each cell type
coefNames(res.dl)
```

---

colData,dreamletProcessedData-method

*Extract colData from dreamletProcessedData*

---

**Description**

Extract colData from dreamletProcessedData

**Usage**

```
## S4 method for signature 'dreamletProcessedData'
colData(x, ...)
```

**Arguments**

x                    A dreamletProcessedData object  
 ...                  other arguments

**Value**

object from colData field

---

```
colData<- ,dreamletProcessedData,ANY-method
```

*Set colData*

---

### Description

Set colData of dreamletProcessedData, and check for same dimensions and rownames

### Usage

```
## S4 replacement method for signature 'dreamletProcessedData,ANY'
colData(x, ...) <- value
```

### Arguments

|       |                                                |
|-------|------------------------------------------------|
| x     | dreamletProcessedData object                   |
| ...   | other arguments                                |
| value | data.frame or object that can be coerced to it |

### Value

none

---

```
compositePosteriorTest
```

*Perform composite test on results from mashr*

---

### Description

The posterior probabilities for all genes and conditions is obtained as 1-lFSR. Let prob be an array storing results for one gene. The probability that `_no_` conditions in the exclusion set are non-zero is `prod(1 - prob[exclude])`. The probability that `_all_` conditions in the inclusion set are non-zero is `prod(prob[include])`. The probability that `_at least one_` condition in the inclusion set is non-zero is `1 - prod(1 - prob[include])`. The composite test is the product of the probabilities computed from the inclusion and exclusion sets.

### Usage

```
compositePosteriorTest(
  x,
  include,
  exclude = NULL,
  test = c("at least 1", "all")
)
```

**Arguments**

|                      |                                                                                             |
|----------------------|---------------------------------------------------------------------------------------------|
| <code>x</code>       | "dreamlet_mash_result" from <code>run_mash()</code>                                         |
| <code>include</code> | array of conditions in the inclusion set                                                    |
| <code>exclude</code> | array of conditions in the exclusion set. Defaults to NULL for no exclusion                 |
| <code>test</code>    | evaluate the posterior probability of a non-zero effect in "at least 1" or "all" conditions |

**Details**

Perform composite test evaluating the specificity of an effect. Evaluate the posterior probability that an a non-zero effect present in `_all_` or `_at least one_` condition in the inclusion set, but `_no conditions_` in the exclusion set.

**See Also**

`run_mash()`

**Examples**

```
library(muscat)
library(mashr)
library(SingleCellExperiment)

data(example_sce)

# create pseudobulk for each sample and cell cluster
pb <- aggregateToPseudoBulk(example_sce[1:100, ],
  assay = "counts",
  cluster_id = "cluster_id",
  sample_id = "sample_id",
  verbose = FALSE
)

# voom-style normalization
res.proc <- processAssays(pb, ~group_id)

# Differential expression analysis within each assay,
# evaluated on the voom normalized data
res.dl <- dreamlet(res.proc, ~group_id)

# run MASH model
# This can take 10s of minutes on real data
# This small datasets should take ~30s
res_mash <- run_mash(res.dl, "group_idstim")

# Composite test based on posterior probabilities
# to identify effect present in *at least 1* monocyte type
# and *NO* T-cell type.
include <- c("CD14+ Monocytes", "FCGR3A+ Monocytes")
exclude <- c("CD4 T cells", "CD8 T cells")

# Perform composite test
prob <- compositePosteriorTest(res_mash, include, exclude)

# examine the lFSR for top gene
```

```
get_lfsr(res_mash$model)[which.max(prob), , drop = FALSE]

# Test if *all* cell types have non-zero effect
prob <- compositePosteriorTest(res_mash, assayNames(res.dl))
```

---

|                   |                                      |
|-------------------|--------------------------------------|
| computeCellCounts | <i>Get cell counts with metadata</i> |
|-------------------|--------------------------------------|

---

### Description

Get cell counts with metadata for each sample

### Usage

```
computeCellCounts(sce, annotation, sampleIDs)
```

### Arguments

|            |                                                                        |
|------------|------------------------------------------------------------------------|
| sce        | SingleCellExperiment                                                   |
| annotation | string indicating column in colData(sce) storing cell type annotations |
| sampleIDs  | string indicating column in colData(sce) storing sample identifiers    |

### Value

matrix storing cell counts

### Examples

```
library(muscat)
library(SingleCellExperiment)

data(example_sce)

counts <- computeCellCounts(example_sce, "cluster_id", "sample_id")

counts[1:4, 1:4]
```

---

|               |                                      |
|---------------|--------------------------------------|
| computeLogCPM | <i>Compute log normalized counts</i> |
|---------------|--------------------------------------|

---

### Description

Compute normalized counts as log2 counts per million

### Usage

```
computeLogCPM(
  sce,
  lib.size = colSums2(counts(sce)),
  prior.count = 2,
  scaledByLib = FALSE
)
```

**Arguments**

|             |                                                                                        |
|-------------|----------------------------------------------------------------------------------------|
| sce         | SingleCellExperiment with counts stored as counts(sce)                                 |
| lib.size    | library size for each cell                                                             |
| prior.count | average count to be added to each observation to avoid taking log of zero              |
| scaledByLib | if TRUE, scale pseudocount by lib.size. Else do standard constant pseudocount addition |

**Details**

This function gives same result as `edgeR::cpm(counts(sce), log=TRUE)`

**Value**

matrix of log CPM values

**See Also**

also `edgeR::cpm()`

**Examples**

```
library(muscat)
library(SingleCellExperiment)

data(example_sce)

logcounts(example_sce) <- computeLogCPM(example_sce)
```

---

|                   |                                  |
|-------------------|----------------------------------|
| computeNormCounts | <i>Compute normalized counts</i> |
|-------------------|----------------------------------|

---

**Description**

Compute normalized counts as counts per million

**Usage**

```
computeNormCounts(sce)
```

**Arguments**

|     |                                                        |
|-----|--------------------------------------------------------|
| sce | SingleCellExperiment with counts stored as counts(sce) |
|-----|--------------------------------------------------------|

**Details**

This function gives same result as `edgeR::cpm(counts(sce), log=FALSE)`

**Value**

matrix of CPM values

**See Also**

also `edgeR::cpm()`

**Examples**

```
library(muscat)
library(SingleCellExperiment)

data(example_sce)

normcounts(example_sce) <- computeNormCounts(example_sce)
```

---

details

*Extract details from dreamletProcessedData*

---

**Description**

Extract details from `dreamletProcessedData`

**Usage**

```
details(object)

## S4 method for signature 'dreamletProcessedData'
details(object)

## S4 method for signature 'dreamletResult'
details(object)

## S4 method for signature 'vpDF'
details(object)
```

**Arguments**

`object`                    A `dreamletProcessedData` object

**Value**

Extract detailed information from some classes

**Examples**

```
library(muscat)
library(SingleCellExperiment)

data(example_sce)

# create pseudobulk for each sample and cell cluster
pb <- aggregateToPseudoBulk(example_sce,
  assay = "counts",
  cluster_id = "cluster_id",
  sample_id = "sample_id",
  verbose = FALSE
```

```

)

# voom-style normalization
res.proc <- processAssays(pb, ~group_id)

# For each cell type, number of samples retained,
# and variables retained
details(res.proc)

```

---

diffVar, dreamletResult-method

*Test differential variance*


---

## Description

Test the association between a covariate of interest and the response's deviation from expectation.

## Usage

```

## S4 method for signature 'dreamletResult'
diffVar(
  fit,
  method = c("AD", "SQ"),
  scale = c("leverage", "none"),
  BPPARAM = SerialParam(),
  ...
)

```

## Arguments

|         |                                                                                      |
|---------|--------------------------------------------------------------------------------------|
| fit     | model fit from dream()                                                               |
| method  | transform the residuals using absolute deviation ("AD") or squared deviation ("SQ"). |
| scale   | scale each observation by "leverage", or no scaling ("none")                         |
| BPPARAM | parameters for parallel evaluation                                                   |
| ...     | other parameters passed to dream()                                                   |

## Details

This method performs a test of differential variance between two subsets of the data, in a way that generalizes to multiple categories, continuous variables and metrics of spread beyond variance. For the two category test, this method is similar to Levene's test. This model was adapted from Phipson, et al (2014), extended to linear mixed models, and adapted to be compatible with `variancePartition::dream()` and `dreamlet::dreamlet()`.

This method is composed of multiple steps where 1) a typical linear (mixed) model is fit with `dreamlet()`, 2) residuals are computed and transformed based on an absolute value or squaring transform, 3) a second regression is performed with `dreamlet()` to test if a variable is associated with increased deviation from expectation. Both regression take advantage of the `dreamlet()` linear (mixed) modelling framework followed by empirical Bayes shrinkage that extends the `limma::voom()` framework.

Note that `diffVar()` takes the results of the first regression as a parameter to use as a starting point.

## References

Phipson B, Oshlack A (2014). “DiffVar: a new method for detecting differential variability with application to methylation in cancer and aging.” *Genome biology*, **15**(9), 1–16.

## See Also

```
variancePartition::diffVar()
variancePartition::diffVar(), missMethyl::diffVar()
```

## Examples

```
library(muscat)
library(SingleCellExperiment)

data(example_sce)

# create pseudobulk for each sample and cell cluster
pb <- aggregateToPseudoBulk(example_sce,
  assay = "counts",
  cluster_id = "cluster_id",
  sample_id = "sample_id",
  verbose = FALSE
)

# voom-style normalization
res.proc <- processAssays(pb, ~group_id)

# Differential expression analysis within each assay,
# evaluated on the voom normalized data
res.dl <- dreamlet(res.proc, ~group_id)

# Differential variance analysis
# result is a dreamlet fit
res.dvar <- diffVar(res.dl)

# Examine results
res.dvar

# Examine details for each assay
details(res.dvar)

# show coefficients estimated for each cell type
coefNames(res.dvar)

# extract results using limma-style syntax
# combines all cell types together
# adj.P.Val gives study-wide FDR
topTable(res.dvar, coef = "group_idstim", number = 3)

# Plot top hit to see differential variance
# Note that this is a toy example with only 4 samples
cellType <- "CD4 T cells"
gene <- "DYNLRB1"

y <- res.proc[[cellType]]$E[gene, ]
x <- colData(res.proc)$group_id
```

```

boxplot(y ~ x,
        xlab = "Stimulation status",
        ylab = "Gene expression",
        main = paste(cellType, gene)
)
#

```

---

dreamlet

*Differential expression for each assay*


---

## Description

Perform differential expression for each assay using linear (mixed) models

## Usage

```

dreamlet(
  x,
  formula,
  data = colData(x),
  assays = assayNames(x),
  contrasts = NULL,
  min.cells = 10,
  robust = FALSE,
  quiet = FALSE,
  BPPARAM = SerialParam(),
  use.eBayes = TRUE,
  ...
)

## S4 method for signature 'dreamletProcessedData'
dreamlet(
  x,
  formula,
  data = colData(x),
  assays = assayNames(x),
  contrasts = NULL,
  min.cells = 10,
  robust = FALSE,
  quiet = FALSE,
  BPPARAM = SerialParam(),
  use.eBayes = TRUE,
  ...
)

```

## Arguments

|         |                                                         |
|---------|---------------------------------------------------------|
| x       | SingleCellExperiment or dreamletProcessedData object    |
| formula | regression formula for differential expression analysis |
| data    | metadata used in regression formula                     |

|                         |                                                                                                                                                                                      |
|-------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| <code>assays</code>     | array of assay names to include in analysis. Defaults to <code>assayNames(x)</code>                                                                                                  |
| <code>contrasts</code>  | character vector specifying contrasts specifying linear combinations of fixed effects to test. This is fed into <code>makeContrastsDream( formula, data, contrasts=contrasts)</code> |
| <code>min.cells</code>  | minimum number of observed cells for a sample to be included in the analysis                                                                                                         |
| <code>robust</code>     | logical, use eBayes method that is robust to outlier genes                                                                                                                           |
| <code>quiet</code>      | show messages                                                                                                                                                                        |
| <code>BPPARAM</code>    | parameters for parallel evaluation                                                                                                                                                   |
| <code>use.eBayes</code> | should eBayes be used on result? (default: TRUE)                                                                                                                                     |
| <code>...</code>        | other arguments passed to <code>dream</code>                                                                                                                                         |

### Details

Fit linear (mixed) model on each cell type separately. For advanced use of contrasts see `variancePartition::makeContrasts` and vignette <https://gabrielhoffman.github.io/variancePartition/articles/dream.html#advanced-hypothesis-testing-1>.

### Value

Object of class `dreamletResult` storing results for each cell type

### See Also

`variancePartition::dream()`, `variancePartition::makeContrastsDream()`

### Examples

```
library(muscat)
library(SingleCellExperiment)

data(example_sce)

# create pseudobulk for each sample and cell cluster
pb <- aggregateToPseudoBulk(example_sce,
  assay = "counts",
  cluster_id = "cluster_id",
  sample_id = "sample_id",
  verbose = FALSE
)

# voom-style normalization
res.proc <- processAssays(pb, ~group_id)

# Differential expression analysis within each assay,
# evaluated on the voom normalized data
res.dl <- dreamlet(res.proc, ~group_id)

# Examine results
res.dl

# Examine details for each assay
details(res.dl)

# show coefficients estimated for each cell type
coefNames(res.dl)
```

```
# extract results using limma-style syntax
# combines all cell types together
# adj.P.Val gives study-wide FDR
topTable(res.dl, coef = "group_idstim", number = 3)
```

---

dreamletCompareClusters

*Differential expression between pair of assays*


---

## Description

Perform differential expression between a pair of assays using linear (mixed) models

## Usage

```
dreamletCompareClusters(
  pb,
  assays,
  method = c("fixed", "random", "none"),
  formula = ~0,
  collapse = TRUE,
  min.cells = 10,
  min.count = 10,
  min.samples = 4,
  isCounts = TRUE,
  normalize.method = "TMM",
  robust = FALSE,
  quiet = FALSE,
  contrasts = c(compare = paste("cellClustertest - cellClusterbaseline")),
  BPPARAM = SerialParam(),
  errorsAsWarnings = FALSE,
  ...
)
```

## Arguments

|           |                                                                                                                                                                                                                                                                                  |
|-----------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| pb        | pseudobulk data as SingleCellExperiment object                                                                                                                                                                                                                                   |
| assays    | array of two entries specifying assays (i.e. cell clusters) to compare, or a list of two sets of assays.                                                                                                                                                                         |
| method    | account for repeated measures from donors using a "random" effect, a "fixed" effect, or "none"                                                                                                                                                                                   |
| formula   | covariates to include in the analysis.                                                                                                                                                                                                                                           |
| collapse  | if TRUE (default), combine all cell clusters within the test set, and separately the baseline set. If FALSE, estimate coefficient for each cell cluster and then identify differential expression using linear contrasts with <code>variancePartition::makeContrastsDream</code> |
| min.cells | minimum number of observed cells for a sample to be included in the analysis                                                                                                                                                                                                     |
| min.count | minimum number of reads for a gene to be consider expressed in a sample. Passed to <code>edgeR::filterByExpr</code>                                                                                                                                                              |

|                               |                                                                                                                                                                                                                                                            |
|-------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| <code>min.samples</code>      | minimum number of samples passing cutoffs for cell cluster to be retained                                                                                                                                                                                  |
| <code>isCounts</code>         | logical, indicating if data is raw counts                                                                                                                                                                                                                  |
| <code>normalize.method</code> | normalization method to be used by <code>calcNormFactors</code>                                                                                                                                                                                            |
| <code>robust</code>           | logical, use eBayes method that is robust to outlier genes                                                                                                                                                                                                 |
| <code>quiet</code>            | show messages                                                                                                                                                                                                                                              |
| <code>contrasts</code>        | cell type is encoded in variable <code>cellCluster</code> with levels <code>test</code> and <code>baseline</code> . <code>contrasts</code> specifies contrasts passed to <code>variancePartition::makeContrastsDream()</code> . Note, advanced users only. |
| <code>BPPARAM</code>          | parameters for parallel evaluation                                                                                                                                                                                                                         |
| <code>errorsAsWarnings</code> | if TRUE, convert error to a warning and return NULL                                                                                                                                                                                                        |
| <code>...</code>              | other arguments passed to <code>dream</code>                                                                                                                                                                                                               |

## Details

Analyze pseudobulk data to identify differential gene expression between two cell clusters or sets of clusters while modeling the cross-donor expression variation and other aspects of the study design.

`dreamletCompareClusters()` is useful for finding genes that are differentially expressed between cell clusters and estimating their fold change. However, the p-values and number of differentially expressed genes are problematic for two reasons, so users must be careful not to overinterpret them:

1. Cell clusters are typically identified with the same gene expression data used for this differential expression analysis between clusters. The same data is used both for discovery and testing, and this means that the p-values from the differential expression analysis will not be uniform under the null. This will produce a lot of findings with small p-values even in the absence of true biological differences.
2. The `dreamlet` package is designed for large datasets with many subjects. The sample sizes from cohort studies are an order of magnitude larger than typical single cell studies. This means that these analyses have huge power to detect even subtle difference in expression between cell clusters. While cluster-specific marker genes are often discovered from an handful of samples, the `dreamlet` package is applicable to 100s or 1000s of subjects.

`method` indicates the regression method used to test differential expression between sets of cell clusters. Since the same biosample will usually be represented in both sets of cell clusters, `method` determines how the paired design is modeled. For `method = "mixed"`, the sample is modeled as a random effect:  $\sim (1|\text{Sample}) + \dots$ . For `method = "fixed"`, the sample is modeled as a fixed effect:  $\sim \text{Sample} + \dots$ . For `method = "none"`, the pairing is ignored.

When `collapse=TRUE` (default) combine all cell clusters within the test set, and separately the baseline set, and estimate a coefficient indicating the differential expression between sets for a given gene. If `collapse=FALSE`, estimate a coefficient for each cell type and then identify differential expression using linear contrasts with `variancePartition::makeContrastsDream()`.

## Value

Object of class `dreamletResult` storing results for each comparison

**Examples**

```

library(muscat)
library(SingleCellExperiment)

data(example_sce)

# create pseudobulk for each sample and cell cluster
pb <- aggregateToPseudoBulk(example_sce,
  assay = "counts",
  cluster_id = "cluster_id",
  sample_id = "sample_id",
  verbose = FALSE
)

# Evaluate the specificity of each gene for each cluster
df_cts <- cellTypeSpecificity(pb)

# compare first two assays (i.e. cell types)
ct.pairs <- c("B cells", "CD14+ Monocytes")

# run comparison
# use method = 'fixed' here since it is faster
fit <- dreamletCompareClusters(pb, ct.pairs, method = "fixed")

# Extract top 10 differentially expressed genes
# The coefficient 'compare' is the value logFC between test and baseline:
# compare = cellClustertest - cellClusterbaseline
res <- topTable(fit, coef = "compare", number = 10)

# genes with highest logFC are most highly expressed in
# B cells compared to CD14+ Monocytes
head(res)

dreamlet::plotHeatmap(df_cts, genes = rownames(res)[1:5])

# compare B cells versus the rest of the cell types
# 'rest' is a keyword indicating all other assays
fit <- dreamletCompareClusters(pb, c("B cells", "rest"), method = "fixed")

res <- topTable(fit, coef = "compare", number = 10)

# genes with highest logFC are most highly expressed in
# B cells compared to all others
head(res)

# Get genes upregulated in B cells
idx <- with(res, which(logFC > 0))[1:5]
dreamlet::plotHeatmap(df_cts, genes = rownames(res)[idx])

lst <- list(
  test = c("CD14+ Monocytes", "FCGR3A+ Monocytes"),
  baseline = c("CD4 T cells", "CD8 T cells")
)

# compare 2 monocyte clusters to two T cell clusters
fit <- dreamletCompareClusters(pb, lst, method = "fixed")

```

```

res <- topTable(fit, coef = "compare", number = 10)

# genes with highest logFC are most highly expressed in
# monocytes compared to T cells
head(res)

# Get genes upregulated in monocytes
idx <- with(res, which(logFC > 0))[1:5]
dreamlet::plotHeatmap(df_cts, genes = rownames(res)[idx])

```

---

`dreamletProcessedData-class`

*Class dreamletProcessedData*

---

### Description

Class `dreamletProcessedData`

### Value

none

none

---

`dreamletResult-class`    *Class dreamletResult*

---

### Description

Class `dreamletResult` stores results produced by `dreamlet()` to give a standard interface for downstream analysis

Class `dreamletResult` stores results produced by `dreamlet()` to give a standard interface for downstream analysis

### Value

none

none

---

`dreamlet_mash_result-class`

*Class dreamlet\_mash\_result*

---

### Description

Class `dreamlet_mash_result`

### Value

`dreamlet_mash_result` class

---

|                    |                                            |
|--------------------|--------------------------------------------|
| dropRedundantTerms | <i>Drop redundant terms from the model</i> |
|--------------------|--------------------------------------------|

---

**Description**

Detect co-linear fixed effects and drop the last one

**Usage**

```
dropRedundantTerms(formula, data, tol = 0.001)
```

**Arguments**

|         |                                                          |
|---------|----------------------------------------------------------|
| formula | original formula                                         |
| data    | data.frame                                               |
| tol     | tolerance to test difference of correlation from 1 or -1 |

**Value**

a formula, possibly with terms omitted.

**Examples**

```
# Valid formula
dropRedundantTerms(~ group + extra, sleep)
```

---

|               |                                        |
|---------------|----------------------------------------|
| equalFormulas | <i>Check if two formulas are equal</i> |
|---------------|----------------------------------------|

---

**Description**

Check if two formulas are equal by evaluating the formulas and extracting terms

**Usage**

```
equalFormulas(formula1, formula2)
```

**Arguments**

|          |                |
|----------|----------------|
| formula1 | first formula  |
| formula2 | second formula |

**Value**

boolean value indicating if formulas are equivalent

**Examples**

```
# These formulas are equivalent
formula1 <- ~ Size + 1
formula2 <- ~ 1 + Size

dreamlet:::equalFormulas(formula1, formula2)
```

---

|             |                                                  |
|-------------|--------------------------------------------------|
| extractData | <i>Extract normalized expression and colData</i> |
|-------------|--------------------------------------------------|

---

**Description**

Extract normalized expression and colData

Extract normalized (i.e. log2 CPM) expression and colData from dreamletProcessedData

**Usage**

```
extractData(x, assay, cols = colnames(colData(x)), genes = rownames(x))

## S4 method for signature 'dreamletProcessedData,character'
extractData(
  x,
  assay,
  cols = colnames(colData(x)),
  genes = rownames(assay(x, assay))
)
```

**Arguments**

|       |                                                                                   |
|-------|-----------------------------------------------------------------------------------|
| x     | dreamletProcessedData object                                                      |
| assay | assay to extract                                                                  |
| cols  | columns in colData(x) to extract. defaults to all columns as colnames(colData(x)) |
| genes | genes to extract from assay(x, assay)\$E. defaults to all genes as rownames(x)    |

**Value**

data.frame or DataFrame of merged expression and colData

**Examples**

```
library(muscat)
library(SingleCellExperiment)

data(example_sce)

# create pseudobulk for each sample and cell cluster
pb <- aggregateToPseudoBulk(example_sce,
  assay = "counts",
  cluster_id = "cluster_id",
  sample_id = "sample_id",
```

```

    verbose = FALSE
  )

  # voom-style normalization
  res.proc <- processAssays(pb, ~group_id)

  # Extract all:
  # Extract tibble of colData merged with expression.
  # variables and genes are stored as columns, samples as rows
  df_merge <- extractData(res.proc, "B cells")

  # first few columns
  df_merge[, 1:6]

  # Extract subset:
  df_merge <- extractData(res.proc, "B cells", cols = "group_id", genes = c("SSU72", "U2AF1"))

  df_merge

  # Boxplot of expression
  boxplot(SSU72 ~ group_id, df_merge)
  #

```

---

fitVarPart

---

*Variance Partition analysis for each assay*


---

## Description

Perform Variance Partition analysis for each assay

## Usage

```

fitVarPart(
  x,
  formula,
  data = colData(x),
  assays = assayNames(x),
  quiet = FALSE,
  BPPARAM = SerialParam(),
  ...
)

## S4 method for signature 'dreamletProcessedData'
fitVarPart(
  x,
  formula,
  data = colData(x),
  assays = assayNames(x),
  quiet = FALSE,
  BPPARAM = SerialParam(),
  ...
)

```

**Arguments**

|                      |                                                                                     |
|----------------------|-------------------------------------------------------------------------------------|
| <code>x</code>       | SingleCellExperiment or dreamletProcessedData object                                |
| <code>formula</code> | regression formula for differential expression analysis                             |
| <code>data</code>    | metadata used in regression formula                                                 |
| <code>assays</code>  | array of assay names to include in analysis. Defaults to <code>assayNames(x)</code> |
| <code>quiet</code>   | show messages                                                                       |
| <code>BPPARAM</code> | parameters for parallel evaluation                                                  |
| <code>...</code>     | other arguments passed to <code>dream</code>                                        |

**Value**

Object of class `vpDF` inheriting from `DataFrame` storing the variance fractions for each gene and cell type.

**Examples**

```
library(muscat)
library(SingleCellExperiment)

data(example_sce)

# create pseudobulk for each sample and cell cluster
pb <- aggregateToPseudoBulk(example_sce,
  assay = "counts",
  cluster_id = "cluster_id",
  sample_id = "sample_id",
  verbose = FALSE
)

# voom-style normalization
res.proc <- processAssays(pb, ~group_id)

# variance partitioning analysis
vp <- fitVarPart(res.proc, ~group_id)

# Show variance fractions at the gene-level for each cell type
genes <- vp$gene[2:4]
plotPercentBars(vp[vp$gene %in% genes, ])

# Summarize variance fractions genome-wide for each cell type
plotVarPart(vp)
```

---

getTreat,dreamletResult-method

*Test if coefficient is different from a specified value*

---

**Description**

Test if coefficient is different from a specified value

**Usage**

```
## S4 method for signature 'dreamletResult'
getTreat(fit, lfc = log2(1.2), coef = NULL, number = 10, sort.by = "p")
```

**Arguments**

|         |                                                                                         |
|---------|-----------------------------------------------------------------------------------------|
| fit     | dreamletResult object                                                                   |
| lfc     | a minimum log2-fold-change below which changes not considered scientifically meaningful |
| coef    | which coefficient to test                                                               |
| number  | number of genes to return                                                               |
| sort.by | column to sort by                                                                       |

**Value**

DataFrame storing hypothesis test for each gene and cell type

**See Also**

```
limma::topTreat(), variancePartition::getTreat()
```

**Examples**

```
library(muscat)
library(SingleCellExperiment)

data(example_sce)

# create pseudobulk for each sample and cell cluster
pb <- aggregateToPseudoBulk(example_sce,
  assay = "counts",
  cluster_id = "cluster_id",
  sample_id = "sample_id",
  verbose = FALSE
)

# voom-style normalization
res.proc <- processAssays(pb, ~group_id)

# Differential expression analysis within each assay,
# evaluated on the voom normalized data
res.dl <- dreamlet(res.proc, ~group_id)

# show coefficients estimated for each cell type
coefNames(res.dl)

# extract results using limma-style syntax
# combines all cell types together
# adj.P.Val gives study-wide FDR
getTreat(res.dl, coef = "group_idstim", number = 3)
```

---

```
metadata, dreamletProcessedData-method
```

*Extract metadata from dreamletProcessedData*

---

### Description

Extract metadata from dreamletProcessedData

### Usage

```
## S4 method for signature 'dreamletProcessedData'
metadata(x)
```

### Arguments

x                      A dreamletProcessedData object

### Value

object from metadata field

---

```
meta_analysis              Meta-analysis across multiple studies
```

---

### Description

Meta-analysis across multiple studies

### Usage

```
meta_analysis(
  x,
  method = "FE",
  group = c("ID", "assay"),
  control = list(maxiter = 2000)
)
```

### Arguments

|         |                                                                                                                                                                                                                                    |
|---------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| x       | data.frame rbind'ing results across genes, cell types and datasets                                                                                                                                                                 |
| method  | meta-analysis method. Values are fed into metafor::rma(), except for 'RE2C' which calls remaCor::RE2C().                                                                                                                           |
| group   | columns in x to group by. For results from dreamlet::topTable(), results are aggregated by gene and cell type (i.e. 'ID' and 'assay'). If x is not from this function, this argument allows the function to group results properly |
| control | passed to rma(..., control)                                                                                                                                                                                                        |

**Details**

'FE': fixed effects meta-analysis  
 'REML': random effects meta-analysis  
 'RE2C': joint testing of fixed and random effects

**Examples**

```
library(dreamlet)
library(muscat)

data(example_sce)

# create pseudobulk for each sample and cell cluster
pb <- aggregateToPseudoBulk(example_sce,
  assay = "counts",
  cluster_id = "cluster_id",
  sample_id = "sample_id",
  verbose = FALSE
)

# voom-style normalization
# just 'CD14+ Monocytes' for speed
res.proc <- processAssays(pb, ~group_id, assays = "CD14+ Monocytes")

# dreamlet
res.dl <- dreamlet(res.proc, ~group_id)

tab1 <- topTable(res.dl, coef = "group_idstim", number = Inf)
tab1$Dataset <- "1"

# Results from a second cohort
# Here, just a copy of the same results for simplicity
tab2 <- tab1
tab2$Dataset <- "2"

# rbind
tab_combined <- rbind(tab1, tab2)

# Perform fixed effects meta-analysis
res <- meta_analysis(tab_combined, method = "FE")

res[1:3, ]
```

---

outlier

*Multivariate outlier detection*


---

**Description**

Detect multivariate outliers using Mahalanobis distance using mean and covariance estimated either with standard or robust methods.

**Usage**

```
outlier(data, robust = FALSE, ...)
```

**Arguments**

|        |                                                 |
|--------|-------------------------------------------------|
| data   | matrix of data                                  |
| robust | use robust covariance method, defaults to FALSE |
| ...    | arguments passed to MASS::cov.rob()             |

**Details**

The distance follow a chisq distrubtion under the null with standard method for mean and covariance. It is approximate if the robust method is used. So use `qchisq(p = 0.999, df = k)` to get cutoff to keep 99.9% of samples under the null for data with  $k=2$  columns.

**Value**

data.frame storing chisq and z-score for each entry indicating deviation from the mean. The z-score is computed by evaluating the p-value of chisq statistic and converting it into a z-score

**Examples**

```
data <- matrix(rnorm(200), 100, 2)

res <- outlier(data)

res[1:4,]
```

---

outlierByAssay

*Outlier analysis for each assay*


---

**Description**

Compute outlier score for each sample in each assay using `outlier()` run on the top principal components. Mahalanobis distance is used for outlier detect and multivariate normal assumption is used to compute p-values

**Usage**

```
outlierByAssay(object, assays = names(object), nPC = 2, robust = FALSE, ...)
```

**Arguments**

|        |                                                        |
|--------|--------------------------------------------------------|
| object | dreamletProcessedData from processAssays()             |
| assays | assays / cell types to analyze                         |
| nPC    | number of PCs to uses for outlier score with outlier() |
| robust | use robust covariance method, defaults to FALSE        |
| ...    | arguments passed to MASS::cov.rob()                    |

**Value**

ID: sample identifier

assay: specify assay

PCs: principal components

chisq: mahalanobis distance that is distributed as  $\text{chisq}(k)$   $k = n\text{PC}$  if the data is multivariate gaussian

z: z-score corresponding to the chisq distance

**See Also**

outlier()

**Examples**

```
library(muscat)
library(SingleCellExperiment)

data(example_sce)

# create pseudobulk for each sample and cell cluster
pb <- aggregateToPseudoBulk(example_sce,
  assay = "counts",
  cluster_id = "cluster_id",
  sample_id = "sample_id",
  verbose = FALSE
)

# voom-style normalization
res.proc <- processAssays(pb, ~group_id)

# Compute PCs and outlier scores
outlierByAssay( res.proc, c("B cells", "CD14+ Monocytes"))
```

---

plotBeeswarm

*Beeswarm plot of effect sizes for each assay*


---

**Description**

Beeswarm plot of effect sizes for each assay, colored by sign and FDR

**Usage**

```
plotBeeswarm(res.dl, coef, fdr.range = 4, assays = assayNames(res.dl))
```

**Arguments**

|           |                                       |
|-----------|---------------------------------------|
| res.dl    | dreamletResult object from dreamlet() |
| coef      | coefficient name fed to topTable()    |
| fdr.range | range for coloring FDR                |
| assays    | which assays to plot                  |

**Value**

ggplot2 of logFC by assay

**Examples**

```
library(muscat)
library(SingleCellExperiment)

data(example_sce)

# create pseudobulk for each sample and cell cluster
pb <- aggregateToPseudoBulk(example_sce,
  assay = "counts",
  cluster_id = "cluster_id",
  sample_id = "sample_id",
  verbose = FALSE
)

# voom-style normalization
res.proc <- processAssays(pb, ~group_id)

# Differential expression analysis within each assay,
# evaluated on the voom normalized data
res.dl <- dreamlet(res.proc, ~group_id)

# Beeswarm plot of effect sizes for each assay,
# colored by sign and FDR
plotBeeswarm(res.dl, "group_idstim")
```

---

plotCellComposition     *Bar plot of cell compositions*

---

**Description**

Bar plot of cell compositions

**Usage**

```
plotCellComposition(obj, col, width = NULL)

## S4 method for signature 'SingleCellExperiment'
plotCellComposition(obj, col, width = NULL)

## S4 method for signature 'matrix'
plotCellComposition(obj, col, width = NULL)

## S4 method for signature 'data.frame'
plotCellComposition(obj, col, width = NULL)
```

**Arguments**

obj                    matrix of [cells] x [samples] or SingleCellExperiment from aggregateToPseudoBulk

col                    array of colors. If missing, use default colors. If names(col) is the same as arrayNames(obj), then colors will be assigned by assay name#

width                  specify width of bars

**Value**

Barplot showing cell fractions

**Examples**

```
library(muscat)
library(SingleCellExperiment)

data(example_sce)

# create pseudobulk for each sample and cell cluster
pb <- aggregateToPseudoBulk(example_sce,
  assay = "counts",
  cluster_id = "cluster_id",
  sample_id = "sample_id",
  verbose = FALSE
)

# show cell composition bar plots
plotCellComposition(pb)

# extract cell counts
df_cellCounts <- cellCounts(pb)

# show cell composition bar plots
plotCellComposition(df_cellCounts)
```

---

plotForest

*Forest plot*


---

**Description**

Forest plot

**Usage**

```
plotForest(x, gene, coef, ...)
```

## S4 method for signature 'dreamletResult'

```
plotForest(x, gene, coef, assays = names(x), ylim = NULL)
```

## S4 method for signature 'dreamlet\_mash\_result'

```
plotForest(x, gene, coef, assays = colnames(x$logFC.original), ylim = NULL)
```

**Arguments**

|        |                                   |
|--------|-----------------------------------|
| x      | result from dreamlet              |
| gene   | gene to show results for          |
| coef   | coefficient to test with topTable |
| ...    | other arguments                   |
| assays | array of assays to plot           |
| ylim   | limits for the y axis             |

**Value**

Plot showing effect sizes

**Examples**

```
library(muscat)
library(SingleCellExperiment)

data(example_sce)

# create pseudobulk for each sample and cell cluster
pb <- aggregateToPseudoBulk(example_sce,
  assay = "counts",
  cluster_id = "cluster_id",
  sample_id = "sample_id",
  verbose = FALSE
)

# voom-style normalization
res.proc <- processAssays(pb, ~group_id)

# Differential expression analysis within each assay,
# evaluated on the voom normalized data
res.dl <- dreamlet(res.proc, ~group_id)

# show coefficients estimated for each cell type
coefNames(res.dl)

# Show estimated log fold change with in each cell type
plotForest(res.dl, gene = "ISG20", coef = "group_idstim")
```

---

|                 |                                    |
|-----------------|------------------------------------|
| plotGeneHeatmap | <i>Heatmap of genes and assays</i> |
|-----------------|------------------------------------|

---

**Description**

Heatmap of genes and assays

**Usage**

```
plotGeneHeatmap(
  x,
  coef,
  genes,
  assays = assayNames(x),
  zmax = NULL,
  transpose = FALSE
)

## S4 method for signature 'dreamletResult'
plotGeneHeatmap(
  x,
  coef,
  genes,
  assays = assayNames(x),
  zmax = NULL,
  transpose = FALSE
)
```

**Arguments**

|           |                                                                                                           |
|-----------|-----------------------------------------------------------------------------------------------------------|
| x         | A dreamletResult object                                                                                   |
| coef      | column number or column name specifying which coefficient or contrast of the linear model is of interest. |
| genes     | array of genes to include in plot                                                                         |
| assays    | array of assay names to include in analysis. Defaults to assayNames(x)                                    |
| zmax      | maximum z.std value                                                                                       |
| transpose | (default: FALSE) Use 'coord_flip()' to flip axes                                                          |

**Value**

Heatmap plot for specified genes and assays

Heatmap plot for specified genes and assays

**Examples**

```
library(muscat)
library(SingleCellExperiment)

data(example_sce)

# create pseudobulk for each sample and cell cluster
pb <- aggregateToPseudoBulk(example_sce,
  assay = "counts",
  cluster_id = "cluster_id",
  sample_id = "sample_id",
  verbose = FALSE
)

# voom-style normalization
res.proc <- processAssays(pb, ~group_id)
```

```
# Differential expression analysis within each assay,
# evaluated on the voom normalized data
res.dl <- dreamlet(res.proc, ~group_id)

# Heatmap for specified subset of genes
plotGeneHeatmap(res.dl, coef = "group_idstim", genes = rownames(pb)[1:15])
```

---

plotHeatmap

*Plot heatmap*


---

## Description

Plot heatmap

## Usage

```
plotHeatmap(
  x,
  genes = rownames(x),
  color = "darkblue",
  assays = colnames(x),
  useFillScale = TRUE
)

## S4 method for signature 'cellSpecificityValues'
plotHeatmap(
  x,
  genes = rownames(x),
  color = "darkblue",
  assays = colnames(x),
  useFillScale = TRUE
)

## S4 method for signature 'data.frame'
plotHeatmap(
  x,
  genes = rownames(x),
  color = "darkblue",
  assays = colnames(x),
  useFillScale = TRUE
)

## S4 method for signature 'matrix'
plotHeatmap(
  x,
  genes = rownames(x),
  color = "darkblue",
  assays = colnames(x),
  useFillScale = TRUE
)
```

**Arguments**

|              |                                                 |
|--------------|-------------------------------------------------|
| x            | fractions for each gene                         |
| genes        | name of genes to plot                           |
| color        | color of heatmap                                |
| assays       | array of assays to plot                         |
| useFillScale | default TRUE. add scale_fill_gradient() to plot |

**Value**

heatmap

**Examples**

```
library(muscat)
library(SingleCellExperiment)

data(example_sce)

# create pseudobulk for each sample and cell cluster
pb <- aggregateToPseudoBulk(example_sce,
  assay = "counts",
  cluster_id = "cluster_id",
  sample_id = "sample_id",
  verbose = FALSE
)

# Compute cell type specificity of each gene
df <- cellTypeSpecificity(pb)

# For each cell type, get most specific gene
genes <- rownames(df)[apply(df, 2, which.max)]

# heatmap of 5 genes that are most cell type specific
dreamlet::plotHeatmap(df, genes = genes)
```

---

plotPCA

---

*Plot PCA of gene expression for an assay*


---

**Description**

Compute PCA of gene expression for an assay, and plot samples coloring by outlier score

**Usage**

```
## S4 method for signature 'list'
plotPCA(
  object,
  assays = names(object),
  nPC = 2,
  robust = FALSE,
  ...,
```

```

    maxOutlierZ = 20,
    nrow = 2,
    size = 2,
    fdr.cutoff = 0.05
  )

```

### Arguments

|                          |                                                                                    |
|--------------------------|------------------------------------------------------------------------------------|
| <code>object</code>      | dreamletProcessedData from processAssays() or a list from residuals()              |
| <code>assays</code>      | assays / cell types to analyze                                                     |
| <code>nPC</code>         | number of PCs to uses for outlier score with outlier()                             |
| <code>robust</code>      | use robust covariance method, defaults to FALSE                                    |
| <code>...</code>         | arguments passed to MASS::cov.rob()                                                |
| <code>maxOutlierZ</code> | cap outlier z-scores at this value for plotting to maintain consistent color scale |
| <code>nrow</code>        | number of rows in plot                                                             |
| <code>size</code>        | size passed to geom_point()                                                        |
| <code>fdr.cutoff</code>  | FDR cutoff to determine outlier                                                    |

### See Also

outlierByAssay()

### Examples

```

library(muscat)
library(SingleCellExperiment)

data(example_sce)

# create pseudobulk for each sample and cell cluster
pb <- aggregateToPseudoBulk(example_sce,
  assay = "counts",
  cluster_id = "cluster_id",
  sample_id = "sample_id",
  verbose = FALSE
)

# voom-style normalization
res.proc <- processAssays(pb, ~group_id)

# PCA to identify outliers
# from normalized expression
plotPCA( res.proc, c("B cells", "CD14+ Monocytes"))

# Run on regression residuals
#-----

# Regression analysis
fit = dreamlet(res.proc, ~ group_id)

# Extract regression residuals
residsObj = residuals(fit)

```

```
# PCA on residuals
plotPCA( residsObj, c("B cells", "CD14+ Monocytes"))
```

---

```
plotPercentBars, vpDF-method
```

*Bar plot of variance fractions*

---

## Description

Bar plot of variance fractions for a subset of genes

## Usage

```
## S4 method for signature 'vpDF'
plotPercentBars(
  x,
  col = c(ggColorHue(ncol(x) - 3), "grey85"),
  genes = unique(x$gene),
  width = NULL,
  ncol = 3,
  ...
)

## S4 method for signature 'cellSpecificityValues'
plotPercentBars(
  x,
  col = ggColorHue(ncol(x)),
  genes = rownames(x),
  width = NULL,
  ...
)
```

## Arguments

|                    |                                                   |
|--------------------|---------------------------------------------------|
| <code>x</code>     | vpDF object returned by <code>fitVarPart()</code> |
| <code>col</code>   | color of bars for each variable                   |
| <code>genes</code> | name of genes to plot                             |
| <code>width</code> | specify width of bars                             |
| <code>ncol</code>  | number of columns in the plot                     |
| <code>...</code>   | other arguments                                   |

## Value

Bar plot showing variance fractions for each gene

**Examples**

```

library(muscat)
library(SingleCellExperiment)

data(example_sce)

# create pseudobulk for each sample and cell cluster
pb <- aggregateToPseudoBulk(example_sce,
  assay = "counts",
  cluster_id = "cluster_id",
  sample_id = "sample_id",
  verbose = FALSE
)

# voom-style normalization
res.proc <- processAssays(pb, ~group_id)

# variance partitioning analysis
vp <- fitVarPart(res.proc, ~group_id)

# Show variance fractions at the gene-level for each cell type
plotPercentBars(vp, genes = vp$gene[2:4], ncol = 2)

```

---

plotProjection

*Plot 2D projection*


---

**Description**

Plot 2D projection (i.e. UMAP, tSNE) for millions of cells efficiently

**Usage**

```

plotProjection(
  sce,
  type,
  annotation,
  pointsize = 0,
  pixels = c(512, 512),
  legend.position = "none",
  text = TRUE,
  order
)

```

**Arguments**

|            |                                                                   |
|------------|-------------------------------------------------------------------|
| sce        | SingleCellExperiment                                              |
| type       | field in reducedDims(sce) to plot                                 |
| annotation | column in colData(sce) to annotate each cell                      |
| pointsize  | Radius of rasterized point. Use 0 for single pixels(fastest).     |
| pixels     | Vector with X and Y resolution of the raster, default c(512, 512) |

|                 |                                                                                                                    |
|-----------------|--------------------------------------------------------------------------------------------------------------------|
| legend.position | legend.position: the position of legends ("none", "left", "right", "bottom", "top", or two-element numeric vector) |
| text            | show annotation as text. Default TRUE                                                                              |
| order           | specify order of levels for annotation                                                                             |

### Details

Uses `scattermore::geom_scattermore()` to plot millions of points efficiently

### Value

ggplot2 plot of the projection

### Examples

```
library(muscat)
library(SingleCellExperiment)

data(example_sce)

plotProjection(example_sce, "TSNE", "cluster_id", 1)
```

---

plotVarPart,DataFrame-method

*Violin plot of variance fractions*

---

### Description

Violin plot of variance fraction for each gene and each variable

### Usage

```
## S4 method for signature 'DataFrame'
plotVarPart(
  obj,
  col = c(ggColorHue(base::ncol(obj) - 3), "grey85"),
  label.angle = 20,
  main = "",
  ylab = "",
  convertToPercent = TRUE,
  ncol = 3,
  ...
)
```

### Arguments

|             |                                                                                             |
|-------------|---------------------------------------------------------------------------------------------|
| obj         | varParFrac object returned by <code>fitExtractVarPart</code> or <code>extractVarPart</code> |
| col         | vector of colors                                                                            |
| label.angle | angle of labels on x-axis                                                                   |
| main        | title of plot                                                                               |

ylab                    text on y-axis  
convertToPercent       multiply fractions by 100 to convert to percent values  
ncol                   number of columns in the plot  
...                    additional arguments

**Value**

Violin plot showing variance fractions

**Examples**

```
library(muscat)
library(SingleCellExperiment)

data(example_sce)

# create pseudobulk for each sample and cell cluster
pb <- aggregateToPseudoBulk(example_sce,
  assay = "counts",
  cluster_id = "cluster_id",
  sample_id = "sample_id",
  verbose = FALSE
)

# voom-style normalization
res.proc <- processAssays(pb, ~group_id)

# variance partitioning analysis
vp <- fitVarPart(res.proc, ~group_id)

# Summarize variance fractions genome-wide for each cell type
plotVarPart(vp)
```

---

|            |                     |
|------------|---------------------|
| plotViolin | <i>Plot Violins</i> |
|------------|---------------------|

---

**Description**

Plot Violins

**Usage**

```
plotViolin(x, ...)
```

## S4 method for signature 'cellSpecificityValues'

```
plotViolin(x, assays = colnames(x))
```

**Arguments**

x                    fractions for each gene  
...                   other arguments  
assays               array of assays to plot

**Value**

Violin plot

**Examples**

```
library(muscat)
library(SingleCellExperiment)

data(example_sce)

# create pseudobulk for each sample and cell cluster
pb <- aggregateToPseudoBulk(example_sce,
  assay = "counts",
  cluster_id = "cluster_id",
  sample_id = "sample_id",
  verbose = FALSE
)

# Compute cell type specificity of each gene
df <- cellTypeSpecificity(pb)

# Violin plot of specificity scores for each cell type
# Dashed line indicates genes that are equally expressed
# across all cell types. For K cell types, this is 1/K
plotViolin(df)
```

plotVolcano

*Volcano plot for each cell type***Description**

Volcano plot for each cell type

**Usage**

```
plotVolcano(
  x,
  coef,
  nGenes = 5,
  size = 12,
  minp = 9.999999999999997e-311,
  cutoff = 0.05,
  ncol = 3,
  ...
)

## S4 method for signature 'list'
plotVolcano(
  x,
  coef,
  nGenes = 5,
  size = 12,
```

```

    minp = 9.9999999999997e-311,
    cutoff = 0.05,
    ncol = 3,
    assays = names(x),
    ...
)

## S4 method for signature 'MArrayLM'
plotVolcano(
  x,
  coef,
  nGenes = 5,
  size = 12,
  minp = 9.9999999999997e-311,
  cutoff = 0.05,
  ncol = 3,
  ...
)

## S4 method for signature 'dreamlet_mash_result'
plotVolcano(
  x,
  coef,
  nGenes = 5,
  size = 12,
  minp = 1e-16,
  cutoff = 0.05,
  ncol = 3,
  assays = colnames(x$logFC.original),
  ...
)

```

### Arguments

|        |                                                                           |
|--------|---------------------------------------------------------------------------|
| x      | result from dreamlet                                                      |
| coef   | coefficient to test with topTable                                         |
| nGenes | number of genes to highlight in each volcano plot                         |
| size   | text size                                                                 |
| minp   | minimum p-value to show on the y-axis                                     |
| cutoff | adj.P.Val cutoff to distinguish significant from non-significant genes    |
| ncol   | number of columns in the plot                                             |
| ...    | arguments passed to facet_wrap(). Useful for specifying scales = "free_y" |
| assays | which assays to plot                                                      |

### Value

Volcano plot for each cell type

**Examples**

```

library(muscat)
library(SingleCellExperiment)

data(example_sce)

# create pseudobulk for each sample and cell cluster
pb <- aggregateToPseudoBulk(example_sce,
  assay = "counts",
  cluster_id = "cluster_id",
  sample_id = "sample_id",
  verbose = FALSE
)

# voom-style normalization
res.proc <- processAssays(pb, ~group_id)

# Differential expression analysis within each assay,
# evaluated on the voom normalized data
res.dl <- dreamlet(res.proc, ~group_id)

# show coefficients estimated for each cell type
coefNames(res.dl)

# volcano plot for each cell type
plotVolcano(res.dl, coef = "group_idstim")

# volcano plot for first two cell types
plotVolcano(res.dl[1:2], coef = "group_idstim")

```

plotVoom

*Plot voom curves from each cell type***Description**

Plot voom curves from each cell type

**Usage**

```

plotVoom(x, ncol = 3, alpha = 0.5, ...)

## S4 method for signature 'dreamletProcessedData'
plotVoom(x, ncol = 3, alpha = 0.5, assays = names(x))

## S4 method for signature 'EList'
plotVoom(x, ncol = 3, alpha = 0.5)

```

**Arguments**

|      |                               |
|------|-------------------------------|
| x    | dreamletProcessedData         |
| ncol | number of columns in the plot |

|        |                        |
|--------|------------------------|
| alpha  | transparency of points |
| ...    | other arguments        |
| assays | which assays to plot   |

**Value**

Plot of mean-variance trend

**Examples**

```
library(muscat)
library(SingleCellExperiment)

data(example_sce)

# create pseudobulk for each sample and cell cluster
pb <- aggregateToPseudoBulk(example_sce,
  assay = "counts",
  cluster_id = "cluster_id",
  sample_id = "sample_id",
  verbose = FALSE
)

# voom-style normalization
res.proc <- processAssays(pb, ~group_id)

# Show mean-variance trend from voom
plotVoom(res.proc)

# plot for first two cell types
plotVoom(res.proc[1:2])
```

---

*print,dreamletResult-method*

*Print object*

---

**Description**

Print object

Print object

**Usage**

```
## S4 method for signature 'dreamletResult'
print(x, ...)

## S4 method for signature 'dreamletProcessedData'
print(x, ...)
```

**Arguments**

|     |                              |
|-----|------------------------------|
| x   | dreamletProcessedData object |
| ... | other arguments              |

**Value**

print data stored in object

---

processAssays

*Processing SingleCellExperiment to dreamletProcessedData*


---

**Description**

For raw counts, estimate precision weights using linear mixed model weighting by number of cells observed for each sample. For normalized data, only weight by number of cells.

**Usage**

```
processAssays(
  sceObj,
  formula,
  assays = assayNames(sceObj),
  min.cells = 5,
  min.count = 5,
  min.samples = 4,
  min.prop = 0.4,
  isCounts = TRUE,
  normalize.method = "TMM",
  span = "auto",
  quiet = FALSE,
  weightsList = NULL,
  BPPARAM = SerialParam(),
  ...
)
```

**Arguments**

|                  |                                                                                                                        |
|------------------|------------------------------------------------------------------------------------------------------------------------|
| sceObj           | SingleCellExperiment object                                                                                            |
| formula          | regression formula for differential expression analysis                                                                |
| assays           | array of assay names to include in analysis. Defaults to assayNames(sceObj)                                            |
| min.cells        | minimum number of observed cells for a sample to be included in the analysis                                           |
| min.count        | used to compute a CPM threshold of $CPM.cutoff = min.count / median(lib.size) * 1e6$ . Passed to edgeR::filterByExpr() |
| min.samples      | minimum number of samples passing cutoffs for cell cluster to be retained                                              |
| min.prop         | minimum proportion of retained samples with $CPM > CPM.cutoff$                                                         |
| isCounts         | logical, indicating if data is raw counts                                                                              |
| normalize.method | normalization method to be used by calcNormFactors                                                                     |

|             |                                                                                                     |
|-------------|-----------------------------------------------------------------------------------------------------|
| span        | Lowess smoothing parameter using by <code>variancePartition::voomWithDreamWeights()</code>          |
| quiet       | show messages                                                                                       |
| weightsList | list storing matrix of precision weights for each cell type. If NULL precision weights are set to 1 |
| BPPARAM     | parameters for parallel evaluation                                                                  |
| ...         | other arguments passed to dream                                                                     |

### Details

For each cell cluster, samples with at least `min.cells` are retained. Only clusters with at least `min.samples` retained samples are kept. Genes are retained if they have at least `min.count` reads in at least `min.prop` fraction of the samples. Current values are reasonable defaults, since genes that don't pass these cutoffs are very underpowered for differential expression analysis and only increase the multiple testing burden. But values of `min.cells = 2` and `min.count = 2` are also reasonable to include more genes in the analysis.

The precision weights are estimated using the residuals fit from the specified formula. These weights are robust to changes in the formula as long as the major variables explaining the highest fraction of the variance are included.

If `weightsList` is NULL, precision weights are set to 1 internally.

### Value

Object of class `dreamletProcessedData` storing voom-style normalized expression data

### See Also

`voomWithDreamWeights()`

### Examples

```
library(muscat)
library(SingleCellExperiment)

data(example_sce)

# create pseudobulk for each sample and cell cluster
pb <- aggregateToPseudoBulk(example_sce,
  assay = "counts",
  cluster_id = "cluster_id",
  sample_id = "sample_id",
  verbose = FALSE
)

# voom-style normalization
res.proc <- processAssays(pb, ~group_id)

# Differential expression analysis within each assay,
# evaluated on the voom normalized data
res.dl <- dreamlet(res.proc, ~group_id)
#
```

---

|                 |                                              |
|-----------------|----------------------------------------------|
| processOneAssay | <i>Processing expression data from assay</i> |
|-----------------|----------------------------------------------|

---

## Description

For raw counts, filter genes and samples, then estimate precision weights using linear mixed model weighting by number of cells observed for each sample. For normalized data, only weight by number of cells

## Usage

```
processOneAssay(
  y,
  formula,
  data,
  n.cells,
  min.cells = 5,
  min.count = 2,
  min.samples = 4,
  min.prop = 0.4,
  min.total.count = 15,
  isCounts = TRUE,
  normalize.method = "TMM",
  span = "auto",
  quiet = TRUE,
  weights = NULL,
  rescaleWeightsAfter = FALSE,
  BPPARAM = SerialParam(),
  ...
)
```

## Arguments

|                  |                                                                                                                                                                 |
|------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------|
| y                | matrix of counts or log2 CPM                                                                                                                                    |
| formula          | regression formula for differential expression analysis                                                                                                         |
| data             | metadata used in regression formula                                                                                                                             |
| n.cells          | array of cell count for each sample                                                                                                                             |
| min.cells        | minimum number of observed cells for a sample to be included in the analysis                                                                                    |
| min.count        | used to compute a CPM threshold of $\text{CPM.cutoff} = \text{min.count} / \text{median}(\text{lib.size}) * 1e6$ . Passed to <code>edgeR::filterByExpr()</code> |
| min.samples      | minimum number of samples passing cutoffs for cell cluster to be retained                                                                                       |
| min.prop         | minimum proportion of retained samples with $\text{CPM} > \text{CPM.cutoff}$                                                                                    |
| min.total.count  | minimum total count required per gene for inclusion                                                                                                             |
| isCounts         | logical, indicating if data is raw counts                                                                                                                       |
| normalize.method | normalization method to be used by <code>calcNormFactors</code>                                                                                                 |
| span             | Lowess smoothing parameter using by <code>variancePartition::voomWithDreamWeights()</code>                                                                      |

|                     |                                                                           |
|---------------------|---------------------------------------------------------------------------|
| quiet               | show messages                                                             |
| weights             | matrix of precision weights                                               |
| rescaleWeightsAfter | default = FALSE, should the output weights be scaled by the input weights |
| BPPARAM             | parameters for parallel evaluation                                        |
| ...                 | other arguments passed to dream                                           |

**Value**

EList object storing log2 CPM and precision weights

**See Also**

processAssays()

---

|                     |                                           |
|---------------------|-------------------------------------------|
| removeConstantTerms | <i>Remove constant terms from formula</i> |
|---------------------|-------------------------------------------|

---

**Description**

Remove constant terms from formula. Also remove categorical variables with a max of one example per category

**Usage**

```
removeConstantTerms(formula, data)
```

**Arguments**

|         |                  |
|---------|------------------|
| formula | original formula |
| data    | data.frame       |

**Details**

Adapted from `MoEClust::drop_constants`

**Value**

a formula, possibly with terms omitted.

**Examples**

```
# Valid formula
removeConstantTerms(~ group + extra, sleep)

# there is no variation in 'group' in this dataset
removeConstantTerms(~ group + extra, sleep[1:3, ])
```

---

`residuals,dreamletResult-method`*Extract residuals from dreamletResult*

---

## Description

Extract residuals from dreamletResult

## Usage

```
## S4 method for signature 'dreamletResult'
residuals(object, y, ..., type = c("response", "pearson"))
```

## Arguments

|                     |                                                             |
|---------------------|-------------------------------------------------------------|
| <code>object</code> | dreamletResult object                                       |
| <code>y</code>      | dreamletProcessedData object                                |
| <code>...</code>    | other arguments                                             |
| <code>type</code>   | compute either "response" residuals or "pearson" residuals. |

## Details

"response" residuals are the typical residuals returned from `lm()`. "pearson" residuals divides each residual value by its estimated standard error. This requires specifying `y`

## Value

residuals from model fit

## Examples

```
library(muscat)
library(SingleCellExperiment)

data(example_sce)

# create pseudobulk for each sample and cell cluster
pb <- aggregateToPseudoBulk(example_sce,
  assay = "counts",
  cluster_id = "cluster_id",
  sample_id = "sample_id",
  verbose = FALSE
)

# voom-style normalization
res.proc <- processAssays(pb, ~group_id)

# Differential expression analysis within each assay,
# evaluated on the voom normalized data
res.dl <- dreamlet(res.proc, ~group_id)

# extract typical residuals for each assay (i.e. cell type)
```

```
# Return list with entry for each assay with for retained samples and genes
resid.lst <- residuals(res.dl)

# Get Pearson residuals:
# typical residuals scaled by the standard deviation
residPearson.lst <- residuals(res.dl, res.proc, type = "pearson")
```

run\_mash

*Run mash analysis on dreamlet results***Description**

Run mash analysis on dreamlet results

**Usage**

```
run_mash(fit, coefList)
```

**Arguments**

|          |                                                                                                                                                                                                                                                                              |
|----------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| fit      | result from dreamlet()                                                                                                                                                                                                                                                       |
| coefList | coefficient to be analyzed. Assumes 1) the null distribution of the two coefficients is similar, 2) the effects sizes are on the same scale, and 3) the effect estimates should be shrunk towards each other. If these are not satisfied, run separately on each coefficient |

**Details**

Apply **mashr** analysis (Urbut et al. 2019) on the joint set of coefficients for each gene and cell type. mashr is a Bayesian statistical method that borrows strength across tests (i.e. genes and cell types) by learning the distribution of non-zero effects based the observed logFC and standard errors. The method then estimates the posterior distributions of each coefficient based on the observed value and the genome-wide empirical distribution.

mashr has been previously applied to differential expression in **GTEx** data using multiple tissues from the same set of donors (Oliva et al. 2020).

In single cell data, a given gene is often not sufficiently expressed in all cell types. So it is not evaluated in a subsets of cell types, and its coefficient value is NA. Since mashr assumes coefficients and standard errors for every gene and cell type pair, entries with these missing values are set to have coef = 0, and se = 1e6. The output of mashr is then modified to set the corresponding values to NA, to avoid nonsensical results downstream.

**Value**

a list storing the mashr model as model and the original coefficients as logFC.original

## References

Oliva M, Munoz-Aguirre M, Kim-Hellmuth S, Wucher V, Gewirtz AD, Cotter DJ, Parsana P, Kasela S, Balliu B, Vinuela A, others (2020). “The impact of sex on gene expression across human tissues.” *Science*, **369**(6509), eaba3066. <https://doi.org/10.1126/science.aba3066>.

Urbat SM, Wang G, Carbonetto P, Stephens M (2019). “Flexible statistical methods for estimating and testing effects in genomic studies with multiple conditions.” *Nature genetics*, **51**(1), 187–195. <https://doi.org/10.1038/s41588-018-0268-8>.

## See Also

`mashr::mash_estimate_corr_em()`, `mashr::cov_canonical`, `mashr::mash_set_data`

## Examples

```
library(muscat)
library(mashr)
library(SingleCellExperiment)

data(example_sce)

# create pseudobulk for each sample and cell cluster
pb <- aggregateToPseudoBulk(example_sce[1:100, ],
  assay = "counts",
  cluster_id = "cluster_id",
  sample_id = "sample_id",
  verbose = FALSE
)

# voom-style normalization
res.proc <- processAssays(pb, ~group_id)

# Differential expression analysis within each assay,
# evaluated on the voom normalized data
res.dl <- dreamlet(res.proc, ~group_id)

# run MASH model
# This can take 10s of minutes on real data
# This small datasets should take ~30s
res_mash <- run_mash(res.dl, "group_idstim")

# extract statistics from mashr model
# NA values indicate genes not sufficiently expressed
# in a given cell type

# original logFC
head(res_mash$logFC.original)

# posterior mean for logFC
head(get_pm(res_mash$model))

# how many gene-by-celltype tests are significant
# i.e. if a gene is significant in 2 celltypes, it is counted twice
table(get_lfsr(res_mash$model) < 0.05, useNA = "ifany")

# how many genes are significant in at least one cell type
```

```

table(apply(get_lfsr(res_mash$model), 1, min, na.rm = TRUE) < 0.05)

# how many genes are significant in each cell type
apply(get_lfsr(res_mash$model), 2, function(x) sum(x < 0.05, na.rm = TRUE))

# examine top set of genes
# which genes are significant in at least 1 cell type
sort(names(get_significant_results(res_mash$model)))[1:10]

# Lets examine EN01
# There is a lot of variation in the raw logFC
res_mash$logFC.original["EN01", ]

# posterior mean after borrowing across cell type and genes
get_pm(res_mash$model)["EN01", ]

# forest plot based on mashr results
plotForest(res_mash, "EN01")

# volcano plot based on mashr results
# yaxis uses local false sign rate (lfsr)
plotVolcano(res_mash)

# Comment out to reduce package runtime
# gene set analysis using mashr results
# library(zenith)
# go.gs = get_GeneOntology("CC", to="SYMBOL")
# df_gs = zenith_gsa(res_mash, go.gs)

# Heatmap of results
# plotZenithResults(df_gs, 2, 1)

```

---

seeErrors

*Get error text*


---

## Description

Get error text

## Usage

```

seeErrors(obj)

## S4 method for signature 'dreamletResult'
seeErrors(obj)

## S4 method for signature 'dreamletProcessedData'
seeErrors(obj)

## S4 method for signature 'vpDF'
seeErrors(obj)

```

**Arguments**

obj                    A dreamletResult object

**Value**

tibble storing error text

**Examples**

```
library(muscat)
library(SingleCellExperiment)

data(example_sce)

# create pseudobulk for each sample and cell cluster
pb <- aggregateToPseudoBulk(example_sce,
  assay = "counts",
  cluster_id = "cluster_id",
  sample_id = "sample_id",
  verbose = FALSE
)

# voom-style normalization
res.proc <- processAssays(pb, ~group_id)

# Differential expression analysis within each assay,
# evaluated on the voom normalized data
res.dl <- dreamlet(res.proc, ~group_id)

# show errors
# but none are reported
res.err = seeErrors(res.dl)
```

---

show,dreamletResult-method

*Show object*

---

**Description**

Show object

Show object

**Usage**

```
## S4 method for signature 'dreamletResult'
show(object)

## S4 method for signature 'dreamletProcessedData'
show(object)
```

**Arguments**

object                  dreamletProcessedData object

**Value**

show data stored in object

---

sortCols, vpDF-method    *Sort variance partition statistics*

---

**Description**

Sort variance partition statistics

**Usage**

```
## S4 method for signature 'vpDF'
sortCols(
  x,
  FUN = sum,
  decreasing = TRUE,
  last = c("Residuals", "Measurement.error"),
  ...
)
```

**Arguments**

x                      object returned by fitVarPart()  
 FUN                    function giving summary statistic to sort by. Defaults to sum  
 decreasing            logical. Should the sorting be increasing or decreasing?  
 last                   columns to be placed on the right, regardless of values in these columns  
 ...                    other arguments to sort

**Value**

data.frame with columns sorted by mean value, with Residuals in last column

**Examples**

```
library(muscat)
library(SingleCellExperiment)

data(example_sce)

# create pseudobulk for each sample and cell cluster
pb <- aggregateToPseudoBulk(example_sce,
  assay = "counts",
  cluster_id = "cluster_id",
  sample_id = "sample_id",
  verbose = FALSE
)
```

```
# voom-style normalization
res.proc <- processAssays(pb, ~group_id)

# variance partitioning analysis
vp <- fitVarPart(res.proc, ~group_id)

# Summarize variance fractions genome-wide for each cell type
plotVarPart(sortCols(vp))
```

---

|             |                                     |
|-------------|-------------------------------------|
| stackAssays | <i>Stack assays from pseudobulk</i> |
|-------------|-------------------------------------|

---

## Description

Stack assays from pseudobulk to perform analysis across cell types

## Usage

```
stackAssays(pb, assays = assayNames(pb))
```

## Arguments

|        |                                                                         |
|--------|-------------------------------------------------------------------------|
| pb     | pseudobulk SingleCellExperiment from aggregateToPseudoBulk()            |
| assays | array of assay names to include in analysis. Defaults to assayNames(pb) |

## Value

pseudobulk SingleCellExperiment cbind'ing expression values and rbind'ing colData. The column stackedAssay in colData() stores the assay information of the stacked data.

## Examples

```
library(muscat)
library(SingleCellExperiment)

data(example_sce)

# create pseudobulk for each sample and cell cluster
pb <- aggregateToPseudoBulk(example_sce,
  assay = "counts",
  cluster_id = "cluster_id",
  sample_id = "sample_id",
  verbose = FALSE
)

# Stack assays for joint analysis
pb.stack <- stackAssays(pb)

# voom-style normalization
# assay (i.e. cell type) can now be included as a covariate
res.proc <- processAssays(pb.stack, ~ group_id + stackedAssay)
```

```
# variance partitioning analysis
vp <- fitVarPart(res.proc, ~ group_id + stackedAssay)

# Summarize variance fractions across cell types
plotVarPart(sortCols(vp))

# Interaction analysis allows group_id
# to have a different effect within each stacedAssay
vp2 <- fitVarPart(res.proc, ~ group_id * stackedAssay)

plotVarPart(sortCols(vp2))

# Interaction model using random effects
form <- ~ (1 | group_id) + (1 | stackedAssay) + (1 | group_id:stackedAssay)
#
```

---

|             |                                        |
|-------------|----------------------------------------|
| tabToMatrix | <i>Convert results table to matrix</i> |
|-------------|----------------------------------------|

---

### Description

Convert results table to matrix

### Usage

```
tabToMatrix(tab, col, rn = "ID", cn = "assay")
```

### Arguments

|     |                               |
|-----|-------------------------------|
| tab | results table from topTable() |
| col | which column to extract       |
| rn  | column id storing rownames    |
| cn  | column id storing colnames    |

### Value

matrix storing values of column col in rows defined by rn and columns defined by cn

---

|                                |                                             |
|--------------------------------|---------------------------------------------|
| topTable,dreamletResult-method | <i>Table of Top Genes from dreamlet fit</i> |
|--------------------------------|---------------------------------------------|

---

### Description

Extract a table of the top-ranked genes from a dreamlet fit.

**Usage**

```
## S4 method for signature 'dreamletResult'
topTable(
  fit,
  coef = NULL,
  number = 10,
  genelist = NULL,
  adjust.method = "BH",
  sort.by = "P",
  resort.by = NULL,
  p.value = 1,
  lfc = 0,
  confint = FALSE
)
```

**Arguments**

|               |                       |
|---------------|-----------------------|
| fit           | dreamletResult object |
| coef          | coef                  |
| number        | number                |
| genelist      | genelist              |
| adjust.method | adjust.method         |
| sort.by       | sort.by               |
| resort.by     | resort.by             |
| p.value       | p.value               |
| lfc           | lfc                   |
| confint       | confint               |

**Value**

data.frame storing hypothesis test for each gene and cell type

**See Also**

limma::topTable(), variancePartition::topTable()

**Examples**

```
library(muscat)
library(SingleCellExperiment)

data(example_sce)

# create pseudobulk for each sample and cell cluster
pb <- aggregateToPseudoBulk(example_sce,
  assay = "counts",
  cluster_id = "cluster_id",
  sample_id = "sample_id",
  verbose = FALSE
)
```

```
# voom-style normalization
res.proc <- processAssays(pb, ~group_id)

# Differential expression analysis within each assay,
# evaluated on the voom normalized data
res.dl <- dreamlet(res.proc, ~group_id)

# show coefficients estimated for each cell type
coefNames(res.dl)

# extract results using limma-style syntax
# combines all cell types together
# adj.P.Val gives study-wide FDR
topTable(res.dl, coef = "group_idstim", number = 3)
```

---

vpDF-class

*Class vpDF*


---

### Description

Class vpDF stores results for each gene for each assay

### Value

none

none

---

zenith\_gsa,dreamletResult,GeneSetCollection-method

*Perform gene set analysis using zenith*


---

### Description

Perform a competitive gene set analysis accounting for correlation between genes.

### Usage

```
## S4 method for signature 'dreamletResult,GeneSetCollection'
zenith_gsa(
  fit,
  geneSets,
  coefs,
  use.ranks = FALSE,
  n_genes_min = 10,
  inter.gene.cor = 0.01,
  progressbar = TRUE,
  ...
)
```

```
## S4 method for signature 'dreamlet_mash_result, GeneSetCollection'
zenith_gsa(
  fit,
  geneSets,
  coefs,
  use.ranks = FALSE,
  n_genes_min = 10,
  inter.gene.cor = 0.01,
  progressbar = TRUE,
  ...
)
```

### Arguments

|                             |                                                                       |
|-----------------------------|-----------------------------------------------------------------------|
| <code>fit</code>            | results from <code>dreamlet()</code>                                  |
| <code>geneSets</code>       | <code>GeneSetCollection</code>                                        |
| <code>coefs</code>          | coefficients to test using <code>topTable(fit, coef=coefs[i])</code>  |
| <code>use.ranks</code>      | do a rank-based test TRUE or a parametric test FALSE? default: FALSE  |
| <code>n_genes_min</code>    | minimum number of genes in a geneset                                  |
| <code>inter.gene.cor</code> | if NA, estimate correlation from data. Otherwise, use specified value |
| <code>progressbar</code>    | if TRUE, show progress bar                                            |
| <code>...</code>            | other arguments                                                       |

### Details

This code adapts the widely used `camera()` analysis (Wu and Smyth 2012) in the `limma` package (Ritchie et al. 2015) to the case of linear (mixed) models used by `variancePartition::dream()`.

### Value

data.frame of results for each gene set and cell type  
 data.frame of results for each gene set and cell type

### Examples

```
library(muscat)
library(SingleCellExperiment)

data(example_sce)

# create pseudobulk for each sample and cell cluster
pb <- aggregateToPseudoBulk(example_sce,
  assay = "counts",
  cluster_id = "cluster_id",
  sample_id = "sample_id",
  verbose = FALSE
)

# voom-style normalization
res.proc <- processAssays(pb, ~group_id)

# Differential expression analysis within each assay,
```

```

# evaluated on the voom normalized data
res.dl <- dreamlet(res.proc, ~group_id)

# Load Gene Ontology database
# use gene 'SYMBOL', or 'ENSEMBL' id
# use get_MSigDB() to load MSigDB
library(zenith)
go.gs <- get_GeneOntology("CC", to = "SYMBOL")

# Run zenith gene set analysis on result of dreamlet
res_zenith <- zenith_gsa(res.dl, go.gs, "group_idstim", progressbar = FALSE)

# for each cell type select 3 genesets with largest t-statistic
# and 1 geneset with the lowest
# Grey boxes indicate the gene set could not be evaluated because
#   to few genes were represented
plotZenithResults(res_zenith, 3, 1)

```

---

```

[,dreamletResult,ANY,ANY,ANY-method
  Subset with brackets

```

---

## Description

Subset with brackets

Subset with brackets

## Usage

```

## S4 method for signature 'dreamletResult,ANY,ANY,ANY'
x[i]

## S4 method for signature 'dreamletProcessedData,ANY,ANY,ANY'
x[i]

```

## Arguments

|   |                              |
|---|------------------------------|
| x | dreamletProcessedData object |
| i | indeces to extract           |

## Value

entries stored at specified index

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