# Package 'MAST'

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Type Package

**Title** Model-based Analysis of Single Cell Transcriptomics

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VignetteBuilder knitr

**Imports** Biobase, BiocGenerics, S4Vectors, data.table, ggplot2, plyr, stringr, abind, methods, parallel, reshape2, stats, stats4, graphics, utils, SummarizedExperiment(>= 1.5.3), progress, Matrix

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Suggests knitr, rmarkdown, testthat, lme4(>= 1.0), blme, roxygen2(> 6.0.0), numDeriv, car, gdata, lattice, GGally, GSEABase, NMF, TxDb.Hsapiens.UCSC.hg19.knownGene, rsvd, limma, RColorBrewer, BiocStyle, scater, DelayedArray, HDF5Array, zinbwave, dplyr

**Description** Methods and models for handling zero-inflated single cell assay data.

**License** GPL(>= 2)

Collate 'AllGenerics.R' 'AllClasses.R' 'CovFromBoots.R'
 'Fluidigm-methods.R' 'GSEA-by-boot.R' 'Hypothesis.R'
 'LmWrapper.R' 'MAST-package.R' 'MultidimensionalScaling.R'
 'RNASeqAssay-methods.R' 'Readers.R' 'SingleCellAssay-methods.R'
 'UtilityFunctions.R' 'ZlmFit-bootstrap.R' 'ZlmFit-logFC.R'
 'ZlmFit.R' 'bayesglm.R' 'convertMASTClassic.R'
 'ebayes-helpers.R' 'filterEval.R' 'helper-methods.R'
 'lmWrapper-bayesglm.R' 'lmWrapper-glm.R' 'lmWrapper-glmer.R'
 'lmWrapper-ridge.R' 'Irtest.R' 'predict.R' 'stat\_ell.R'
 'thresholdSCRNA.R' 'zeroinf.R' 'zlmHooks.R'

RoxygenNote 7.2.3

LazyData true

**biocViews** GeneExpression, DifferentialExpression, GeneSetEnrichment, RNASeq, Transcriptomics, SingleCell

BugReports https://github.com/RGLab/MAST/issues

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# $\mathsf{R}$ topics documented:

| MAST-package                               |
|--|
| applyFlat                                  |
| BayesGLMlike-class                         |
| calcZ 5                                    |
| colData<-,SingleCellAssay,DataFrame-method |
| collectResiduals                           |
| computeEtFromCt                            |
| convertMASTClassicToSingleCellAssay        |
| CovFromBoots                               |
| defaultPrior                               |
| dof  |
| Drop                                       |
| ebayes                                     |
| expavg                                     |
| filterLowExpressedGenes                    |
| fit  |
| freq                                       |
| FromFlatDF                                 |
| FromMatrix                                 |
| getConcordance                             |
| getwellKey                                 |
| GLMlike-class                              |
| gseaAfterBoot                              |
| GSEATests-class                            |
| hushWarning                                |
| Hypothesis                                 |
| impute                                     |
| influence.bayesglm                         |
| invlogit                                   |
| LMERlike-class                             |
| LMlike-class                               |

| logFC                                  |    |
|--|----|
| logmean                                | 3  |
| LRT                                    | 32 |
| lrTest                                 | 33 |
| lrTest,ZlmFit,character-method         | 34 |
| magic_assay_names                      | 35 |
| maits                                  | 36 |
| MAST-defunct                           | 36 |
| mast_filter                            | 3  |
| meld_list_left                         | 38 |
| melt.SingleCellAssay                   | 39 |
| model.matrix                           | 39 |
| model.matrix<                          |    |
| myBiplot                               |    |
| new_with_repaired_slots                | 4  |
| pbootVcov1                             | 42 |
| plot.thresholdSCRNACountMatrix         | 43 |
| plotlrt                                |    |
| plotSCAConcordance                     |    |
| predict.ZlmFit                         |    |
| predicted_sig                          |    |
| primerAverage                          |    |
| print.summaryZlmFit                    |    |
| read.fluidigm                          |    |
| removeResponse                         |    |
| rstandard.bayesglm                     |    |
| SceToSingleCellAssay                   |    |
| se.coef                                |    |
| show,LMlike-method                     |    |
| split,SingleCellAssay,character-method |    |
| stat_ell                               |    |
| subset,SingleCellAssay-method          |    |
| summarize                              |    |
| summary,GSEATests-method               |    |
| summary,ZlmFit-method                  |    |
| · · · · · · · · · · · · · · · · · · ·  |    |
| summary.thresholdSCRNACountMatrix      |    |
| thresholdSCRNACountMatrix              | 59 |
| vbeta                                  | 60 |
| vbetaFA                                | 60 |
| waldTest                               | 6  |
| waldTest,ZlmFit,matrix-method          | 6  |
| xform                                  | 62 |
| zlm                                    | 62 |
| ZlmFit-class                           | 64 |

4 applyFlat

MAST-package

MAST: Model-based Analysis of Single- cell Transcriptomics

# Description

Methods for analysing single cell assay data using hurdle models.

### **Details**

This packages provides data structures and functions for statistical analysis of single-cell assay data such as Fluidigm single cell gene expression assays.

### Author(s)

Maintainer: Andrew McDavid <Andrew\_McDavid@urmc.rochester.edu>

Authors:

- Greg Finak <gfinak@fredhutch.org>
- Masanao Yajima <myajima@fredhutch.org>

### References

Finak, et al. MAST: a flexible statistical framework for assessing transcriptional changes and characterizing heterogeneity in single-cell RNA sequencing data. Genome Biology (2015).

#### See Also

Useful links:

- https://github.com/RGLab/MAST/
- Report bugs at https://github.com/RGLab/MAST/issues

applyFlat

Apply a vectorized binary operation recycling over last dimension

### **Description**

When x is an array of order K, and y is an array of order K-1, whose dimensions otherwise agree, apply FUN by recycling y as necessary over dimension K of x.

### Usage

```
applyFlat(x, y, FUN = "-")
```

BayesGLMlike-class 5

#### **Arguments**

x array, order K y array, order K-1

FUN vectorized binary operation

#### Value

```
array, order K equal to FUN(x,y)
```

### **Examples**

```
##Dumb example, could be done with scale(...,scale=FALSE)
x0 = matrix(1:10, nrow=2)
y0 = rowMeans(x0)
dim(y0) = c(1, 2)
x1 = MAST:::applyFlat(x0,y0)
stopifnot(rowMeans(x1)==0)
```

BayesGLMlike-class

Wrapper for bayesian GLM

### **Description**

Wrapper for bayesian GLM

#### Slots

prior numeric optional 3d array used to specify prior for coefficients useContinuousBayes logical should bayesglm be used to fit the continuous component as well?

calcZ

Get Z or T statistics and P values after running gseaAfterBoot

### **Description**

The Z or T statistics may be reported by component (discrete/continuous) when combined='no' or combined by Fisher's or Stouffer's method (combined='fisher' or combined='stouffer'. Fisher's method uses the product of the p-values, while Stouffer's method uses the sum of the Z/T scores. The "Z" score returned by Fisher is the normal quantile that would yield the observed Fisher P-value, whose sign is derived from the sign of the maximum component Z score. The "Z" score returned by Stouffer when testType='normal' is the sum of the Z scores, over sqrt(2). When testType='t' it is a weighted combination of the Z scores, with weights correponding to the degrees of freedom in each of the t statistics. A t-approximation to this sum of t-variables is derived by matching moments. It seems to be fairly accurate in practice.

### Usage

```
calcZ(gseaObj, testType = "t", combined = "none")
```

### **Arguments**

gseaObj output from gseaAfterBoot

testType either 'normal' or 't'. The 't' test adjusts for excess kurtosis due to the finite

number of bootstrap replicates used to estimate the variance of the statistics.

This will result in more conservative inference.

combined character one of 'none', 'fisher' or 'stouffer'

### Value

3D array with dimensions set (modules) comp ('cont'inuous or 'disc'rete) and metric ('Z' stat and two sided 'P' value that P(z>|Z|)) if combined='no', otherwise just a matrix.

#### See Also

gseaAfterBoot

### **Examples**

```
## See the examples in gseaAfterBoot
example(gseaAfterBoot)
```

#### **Description**

Replace colData with a DataFrame. Checks to make sure that row.names(value) match colnames $\{x\}$ , in contrast to the parent method Checks for a wellKey column, as well.

### Usage

```
## S4 replacement method for signature 'SingleCellAssay,DataFrame' colData(x) \leftarrow value
```

# Arguments

x SingleCellAssay

value DataFrame

# Value

modified SingleCellAssay

collectResiduals 7

| collectResiduals | Residual hooks and collection methods |  |
|------------------|---------------------------------------|--|
|------------------|---------------------------------------|--|

### **Description**

After each gene is fit, a hook function can optionally be run and the output saved. This allows extended computations to be done using the fitted model, without keeping it in memory. Here this is used to calculate various residuals, though in some cases they can be done using only the information contained in the Z1mFit-class.

# Usage

```
collectResiduals(x, sca, newLayerName = "Residuals")
discrete_residuals_hook(x)
continuous_residuals_hook(x)
combined_residuals_hook(x)
deviance_residuals_hook(x)
fitted_phat(x)
partialScore(x, effectRegex)
```

### **Arguments**

x ZlmFit-class

sca SingleCellAssay object to which the residuals should be added

newLayerName character name of the assay layer

effectRegex a regular expression naming columns of the design corresponding to  $Z_0$ . Gen-

erally these should be the treatment effects of interest.

### Value

copy of sca with new layer

### **Functions**

- discrete\_residuals\_hook(): Hook to get the discrete residuals, ie, difference between expected probability of expression and observed
- continuous\_residuals\_hook(): Hook to get the continuous residuals, ie, residuals for conditionally positive observations. If an observation is zero, it's residual is defined to be zero as well.
- combined\_residuals\_hook(): Hook to get the combined residuals, ie, Y-E(U)\*E(V)

8 computeEtFromCt

• deviance\_residuals\_hook(): Standardized deviance residuals hook. Computes the sum of the standardized deviance residuals for the discrete and continuous models (scaled to have unit variance). If the observation is zero then only the discrete component is used.

- fitted\_phat(): Hook to return p\_hat, the predicted probability of expression.
- partialScore(): Compute  $Y_i E(V_i|X_i, Z_0)E(U|X_i, Z_0)$ , where  $Z_0$  is a treatment effect (being left in) and  $X_i$  is a nuisance effect (being regressed out).

#### Total residual types

Each component of the model contributes several flavors of residual, which can be combined in various fashions. The discrete residual can be on the response scale (thus subtracting the predicted probability of expression from the 0/1 expression value). Or it can be a deviance residual, revealing something about the log-likelihood.

### Partial residuals

It's also possible to consider partial residuals, in which the contribution of a particular covariate is added back into the model.

#### See Also

zlm

### **Examples**

```
data(vbetaFA)
svbeta <- subset(vbetaFA, ncells==1)
svbeta <- svbeta[freq(svbeta)>.4,]
window <- function(x1) lapply(assays(x1), function(x2) x2[1:3, 1:6])
#total residuals of the response
z1 <- zlm(~ Stim.Condition, svbeta, hook=discrete_residuals_hook)
window(collectResiduals(z1, svbeta))
z2 <- zlm(~ Stim.Condition, svbeta, hook=continuous_residuals_hook)
window(collectResiduals(z2, svbeta))
z3 <- zlm(~ Stim.Condition, svbeta, hook=combined_residuals_hook)
window(collectResiduals(z3, svbeta))
#partial residuals
colData(svbeta)$ngeneson <- colMeans(assay(svbeta)>0)
z5 <- zlm(~ Stim.Condition + ngeneson, svbeta)
partialScore(z5, 'Stim.Condition')</pre>
```

computeEtFromCt

Compute the Et from the Ct

#### **Description**

Computes the Et value from the Ct value in an existing data frame and returns a new data frame with the Et column appended

### Usage

```
computeEtFromCt(df, column = "Ct", Cmax = 40)
```

### **Arguments**

df a data.frame

column The name of the Ct column. A character. 'Ct' by default.

Cmax the maximum number of cycles performed. 40 by default.

#### Value

A copy of df with the 'Et' column appended

#### Author(s)

Greg Finak

# **Examples**

```
data(vbeta)
vbeta <- computeEtFromCt(vbeta)</pre>
```

convertMASTClassicToSingleCellAssay

Convert a MASTClassic SingleCellAssay

### **Description**

Convert a SingleCellAssay object created with the MASTClassic package to an object recognized by the new MAST package

### Usage

```
convertMASTClassicToSingleCellAssay(object = NULL)
```

### **Arguments**

object of class SingleCellAssay created by MASTClassic

#### **Details**

The function will extract the relevant information from the attributes of the old object and construct a new SingleCellAssay that is recognized by MAST. This function checks that the object is a MASTClassic SingleCellAssay object. It will stop if it is not a SingleCellAssay, return a converted SingleCellAssay if object was created by MASTClassic, and return the original object if the object is already compatible.

10 CovFromBoots

### Value

A MAST SingleCellAssay object.

#### Note

Type checking for old object is not performed.

### **Examples**

```
data(vbetaFA)
convertMASTClassicToSingleCellAssay(vbetaFA)
```

CovFromBoots

Extract the inter-gene covariance matrices for continuous and discrete components of a MAST model for a given coefficient from bootstrap replicates

### **Description**

Computes the genewise covariance for a model coefficient from bootstrap replicates from 'MAST::bootVcov1()'. If coefficients are unestimable (i.e. NA) for a gene, that row/column in the covariance matrix will be NA. Returns a list with components "C" and "D" containing the covariance matrices for the "C"ontinuous and "D"iscrete components of the MAST model.

### Usage

```
CovFromBoots(boots = NULL, coefficient = NULL)
```

### **Arguments**

boots a multidimensional array returned by 'bootVcov1' or 'pbootVcov1'.

coefficient 'character' the name of the model coefficient for which to return the inter-gene

covariance matrices.

#### Value

list with components "C" and "D" containing covariance matrices for the continuous and discrete components of the model.

defaultPrior 11

defaultPrior

Initialize a prior to be used a prior for BayeGLMlike/BayesGLMlike2

### **Description**

Initialize a prior to be used a prior for BayeGLMlike/BayesGLMlike2

### Usage

```
defaultPrior(names)
```

### **Arguments**

names

character vector of coefficients. The '(Intercept)' will be ignored.

#### Value

3d array, with leading dimension giving the prior 'loc'ation, 'scale' and degrees of freedom (df), second dimension giving the component ('C'ontinuous or 'D'iscrete) and trailing dimension giving the coefficient to which the prior applies. The location is initialized to be 0, the scale to 2, and degrees of freedom of 1, following the default of bayesglm.

### **Examples**

```
dp <- defaultPrior('Stim.ConditionUnstim')
## Not run:
data(vbetaFA)
zlmVbeta <- zlm(~ Stim.Condition, vbetaFA, method='bayesglm', coefPrior=dp)
## End(Not run)</pre>
```

dof

Degrees of freedom of Zero inflated model

# Description

Degrees of freedom of Zero inflated model

### Usage

```
dof(object)
```

### **Arguments**

object

LMlike or subclass

12 ebayes

### Value

vector giving the model degrees of freedom for continuous and discrete

Drop

Drop specified dimension from an array

# Description

Like drop(x) but only dropping specified dimensions. There is no testing that the specified dimensions are actually singletons.

# Usage

```
Drop(x, d)
```

### **Arguments**

x array of at least d dimensions d dimension(s) to drop

### Value

```
array x
```

### **Examples**

```
x = array(1:4, dim=c(1, 2, 1, 2))
dx = MAST:::Drop(x, 1)
stopifnot(all(dim(dx)==c(2,1,2)))
```

ebayes

Estimate hyperparameters for hierarchical variance model for continuous component

# Description

ebayesControl is a named list with (optional) components 'method' (one of 'MOM' or 'MLE') and 'model' (one of 'H0' or 'H1') method MOM uses a method-of-moments estimator, while MLE using the marginal likelihood. H0 model estimates the precisions using the intercept alone in each gene, while H1 fits the full model specified by mm

### Usage

```
ebayes(assay_t, ebayesControl, mm, truncate = Inf)
```

expavg 13

### **Arguments**

assay\_t cells X genes matrix

ebayesControl list with (optional) components 'method', 'model'. See details.

mm a model matrix, used when model='H1'.

truncate Genes with sample precisions exceeding this value are discarded when estimat-

ing the hyper parameters

### Value

numeric of length two, giving the hyperparameters in terms of a variance (v) and prior observations (df), inside a structure, with component hess, giving the Fisher Information of the hyperparameters.

expavg

Exponential average

# Description

Puts log transformed values onto natural scale and takes mean of vector. Calculates mean(2^x - 1)

### Usage

```
expavg(x)
```

### **Arguments**

x numeric

### Value

numeric

# **Examples**

```
x <- 1:10
logmean(expavg(x))</pre>
```

14 fit

```
filterLowExpressedGenes
```

Filter low-expressing genes

### **Description**

Filter out genes that have less than some percent threshold expression across all libraries

### Usage

```
filterLowExpressedGenes(assay, threshold = 0.1)
```

### Arguments

assay a SingleCellAssay object

threshold a numeric between 0, and 1, specifying the threshold frequency below which

genes will be filtered out

### Value

```
SingleCellAssay
```

# **Examples**

```
data(vbetaFA)
filterLowExpressedGenes(vbetaFA)
```

fit

fit a zero-inflated regression

# Description

Given a design and formula, fit the zero inflated regression, storing the fits in slots fitC and fitD

# Usage

```
fit(object, response, ...)
## S4 method for signature 'LMERlike,missing'
fit(object, response, silent = TRUE, ...)
```

# **Arguments**

object inheriting from LMlike

response a vector, same length as the design, or if missing then use the current response

... currently ignored

silent mute some warnings emitted from the underlying modeling functions

freq 15

### Value

LMlike or subclass

freq

Summary statistics for genes in an experiment

# Description

freq returns the frequency of expression, i.e., the proportion of non-zero values in sc. NAs can be optionally removed

# Usage

```
freq(sc, na.rm = TRUE)
condmean(sc)
condSd(sc)
numexp(sc)
```

# Arguments

sc SingleCellAssay
na.rm should NAs be removed, or carried through?

#### Value

vector of proportions

### **Functions**

- condmean(): Report the mean non-zero expression value for each gene. NAs are always removed.
- condSd(): Report standard deviation of expression, for positive et for each gene. NAs are always removed.
- numexp(): Report number of expressing cells (\$>0\$) per gene. NAs are removed.

### **Examples**

```
data(vbetaFA)
freq(vbetaFA)
condmean(vbetaFA)
```

16 FromFlatDF

| FromFlatDF | Construct a SingleCellAssay (or derived subclass) from a 'flat' |
|------------|---|
|            | (melted) data.frame/data.table                                  |

# Description

SingleCellAssay are a generic container for such data and are simple wrappers around SummarizedExperiment objects. Subclasses exist that embue the container with additional attributes, eg FluidigmAssay.

# Usage

```
FromFlatDF(
  dataframe,
  idvars,
  primerid,
  measurement,
  id = numeric(0),
  cellvars = NULL,
  featurevars = NULL,
  phenovars = NULL,
  class = "SingleCellAssay",
  check_sanity = TRUE,
  ...
)
```

# Arguments

| dataframe    | A 'flattened' data.frame or data.table containing columns giving cell and feature identifiers and a measurement column  |
|--------------|---|
| idvars       | character vector naming columns that uniquely identify a cell   |
| primerid     | character vector of length $\boldsymbol{1}$ that names the column that identifies what feature (i.e. gene) was measured   |
| measurement  | character vector of length 1 that names the column containing the measurement   |
| id           | An identifier (eg, experiment name) for the resulting object  |
| cellvars     | Character vector naming columns containing additional cellular metadata   |
| featurevars  | Character vector naming columns containing additional feature metadata  |
| phenovars    | Character vector naming columns containing additional phenotype metadata  |
| class        | desired subclass of object. Default SingleCellAssay.  |
| check_sanity | (default: TRUE) Set FALSE to override sanity checks that try to ensure that the default assay is log-transformed and has at least one exact zero. See defaultAssay for details on the "default assay" which is assumed to contain log transformed data. |
|              | additional arguments are ignored  |

FromMatrix 17

### Value

SingleCellAssay, or derived, object

### **Examples**

```
data(vbeta)
colnames(vbeta)
vbeta <- computeEtFromCt(vbeta)
vbeta.fa <- FromFlatDF(vbeta, idvars=c("Subject.ID", "Chip.Number", "Well"),
primerid='Gene', measurement='Et', ncells='Number.of.Cells',
geneid="Gene",cellvars=c('Number.of.Cells', 'Population'),
phenovars=c('Stim.Condition','Time'), id='vbeta all', class='FluidigmAssay')
show(vbeta.fa)
nrow(vbeta.fa)
nrow(vbeta.fa)
head(mcols(vbeta.fa)$primerid)
table(colData(vbeta.fa)$Subject.ID)
vbeta.sub <- subset(vbeta.fa, Subject.ID=='Sub01')
show(vbeta.sub)</pre>
```

FromMatrix

Construct a SingleCellAssay from a matrix or array of expression

### **Description**

If the gene expression measurements are already in a rectangular form, then this function allows an easy way to construct a SingleCellAssay object while still doing some sanity checking of inputs.

### Usage

```
FromMatrix(
  exprsArray,
  cData,
  fData,
  class = "SingleCellAssay",
  check_sanity = TRUE,
  check_logged = check_sanity
)
```

### **Arguments**

| exprsArray | matrix, or a list of matrices, or an array. Columns are cells, rows are genes.  |
|------------|---|
| cData      | cellData an object that can be coerced to a DataFrame, ie, data.frame, AnnotatedDataFrame. Must have as many rows as ncol(exprsArray)     |
| fData      | featureData an object that can be coerced to a DataFrame, ie, data.frame, AnnotatedDataFrame. Must have as many rows as nrow(exprsArray). |
| class      | desired subclass of object. Default SingleCellAssay.  |

18 getConcordance

check\_sanity (default: TRUE) Set FALSE to override sanity checks that try to ensure that the default assay is log-transformed and has at least one exact zero. See defaultAssay for details on the "default assay" which is assumed to contain log transformed data.

check\_logged alias for check\_sanity

#### Value

an object of class class

#### See Also

defaultAssay

### **Examples**

```
ncells <- 10
ngenes <- 5
fData <- data.frame(primerid=LETTERS[1:ngenes])
cData <- data.frame(wellKey=seq_len(ncells))
mat <- matrix(rnorm(ncells*ngenes), nrow=ngenes)
sca <- FromMatrix(mat, cData, fData)
stopifnot(inherits(sca, 'SingleCellAssay'))
stopifnot(inherits(sca, 'SummarizedExperiment'))
##If there are mandatory keywords expected by a class, you'll have to manually set them yourself
cData$ncells <- 1
fd <- FromMatrix(mat, cData, fData)
stopifnot(inherits(fd, 'SingleCellAssay'))</pre>
```

 ${\tt getConcordance}$ 

Get the concordance between two experiments

### **Description**

Return the concordance between two assays (i.e. single cell and hundred cell). The "average" of singleCellRef (after adjusting for the number of cells) and singleCellComp are taken per gene, per groups. A data.frame with one row per gene-groups is returned with some additional columns.

# Usage

```
getConcordance(
    singleCellRef,
    singleCellcomp,
    groups = NULL,
    fun.natural = expavg,
    fun.cycle = logmean
)
```

getConcordance 19

```
getwss(concord, nexp)
getss(concord)
getrc(concord)
```

### **Arguments**

singleCellRef "reference" SingleCellAssay singleCellcomp "comparison" SingleCellAssay

groups character vector giving variable(s) on which the comparison is conditioned fun.natural function to transform the SingleCellAssays to a mRNA proportional level

fun.cycle inverse function of fun.natural

concord data.frame returned by getConcordance

nexp number of expressed cells per row in concord

### Value

concordance between two assays

### **Functions**

- getwss(): getrc the sum of squares, weighted by nexp
- $\bullet$  getss(): return the sum of squares
- getrc(): Return Lin's (1989) concordance correlation coefficient

#### Author(s)

Andrew McDavid

#### See Also

plotSCAConcordance

# Examples

```
data(vbetaFA)
sca1 <- subset(vbetaFA, ncells==1)
sca100 <- subset(vbetaFA, ncells==100)
concord <- getConcordance(sca1, sca100)
getss(concord)
getrc(concord)</pre>
```

20 GLMlike-class

getwellKey

Accessor for wellKey

### **Description**

This returns the wellKey, which is a unique identifier generated by idvars in the mapping

# Usage

```
getwellKey(sc)
```

### **Arguments**

sc

An object with a wellKey

### Value

integer giving the unique id generated

# **Examples**

```
data(vbetaFA)
getwellKey(vbetaFA)
colData(vbetaFA)$wellKey
```

GLMlike-class

Wrapper for regular glm/lm

### **Description**

Wrapper for regular glm/lm

# Usage

```
## S4 method for signature 'GLMlike'
vcov(object, which, ...)
```

# Arguments

object GLMlike

which character, one of 'C', 'D'.

... ignored

### Value

covariance matrix

gseaAfterBoot 21

### Methods (by generic)

• vcov(GLMlike): return the variance/covariance of component which

#### **Slots**

weightFun function to map expression values to probabilities of expression. Currently unused.

gseaAfterBoot

Gene set analysis for hurdle model

### Description

Modules defined in sets are tested for average differences in expression from the "average" gene. By using bootstraps, the between-gene covariance of terms in the hurdle model is found, and is used to adjust for coexpression between genes. We drop genes if the coefficient we are testing was not estimible in original model fit in zFit or in any of the bootstrap replicates (evidenced an NA in the bootstrap array). This might yield overly conservative inference. Since bootstrapping is a randomized procedure, the degrees of freedom of a module (and its variance parameters) might differ from run-to-run. You might try setting var\_estimate='modelbased' to relax this requirement by assuming independence between genes and then using the asymptotic covariance estimates, which are deterministic, but may result in overly-generous inference.

### Usage

```
gseaAfterBoot(
  zFit,
  boots,
  sets,
  hypothesis,
  control = gsea_control(n_randomize = Inf, var_estimate = "bootall")
)
gsea_control(n_randomize = Inf, var_estimate = "bootall")
```

### **Arguments**

zFit object of class ZlmFit boots bootstraps of zFit sets list of indices of genes

hypothesis a Hypothesis to test. Currently only one degree CoefficientHypothesis are

supported.

control parameters as provided by gsea\_control. See details.

n\_randomize the number of genes to sample to approximate the non-module average expres-

sion. Set to Inf to turn off the approximation (the default).

var\_estimate the method used to estimate the variance of the modules, one of bootall, bootdiag,

or modelbased.

22 gseaAfterBoot

#### Value

Object of class GSEATests, containing slots tests, 4D array and bootR, the number of boostrap replicates.

#### **Functions**

• gsea\_control(): set control parameters. See Details.

#### control

control is a list with elements:

- n\_randomize, giving the number of genes to sample to approximate the non-module average expression. Set to Inf to turn off the approximation (the default).
- var\_estimate, giving the method used to estimate the variance of the modules. bootall uses
  the bootstrapped covariance matrices. bootdiag uses only the diagonal of the bootstrapped
  covariance matrix (so assuming independence across genes). modelbased assumes independence across genes and uses the variance estimated from the model.

### **Return Value**

A 4D array is returned, with dimensions "set" (each module), "comp" ('disc'rete or 'cont'inuous), "metric" ('stat' gives the average of the coefficient, 'var' gives the variance of that average, 'dof' gives the number of genes that were actually tested in the set), "group" ('test' for the genes in test-set, "null" for all genes outside the test-set).

### See Also

```
calcZ
```

summary, GSEATests-method

### **Examples**

```
data(vbetaFA)
vb1 = subset(vbetaFA, ncells==1)
vb1 = vb1[,freq(vb1)>.1][1:15,]
zf = zlm(~Stim.Condition, vb1)
boots = bootVcov1(zf, 5)
sets = list(A=1:5, B=3:10, C=15, D=1:5)
gsea = gseaAfterBoot(zf, boots, sets, CoefficientHypothesis('Stim.ConditionUnstim'))
## Use a model-based estimate of the variance/covariance.
gsea_mb = gseaAfterBoot(zf, boots, sets, CoefficientHypothesis('Stim.ConditionUnstim'),
control = gsea_control(var_estimate = 'modelbased'))
calcZ(gsea)
summary(gsea)
```

GSEATests-class 23

GSEATests-class

An S4 class for Gene Set Enrichment output

# Description

This holds output from a call to gseaAfterBoot. It primarily provides a summary method.

### **Slots**

```
tests array: gene sets X discrete,continuous X stat, variance, degrees of freedom, avg correlation X test, null bootR number of bootstrap replicates
```

### See Also

```
gseaAfterBoot
calcZ
summary,GSEATests-method
```

hushWarning

Selectively muffle warnings based on output

### **Description**

Selectively muffle warnings based on output

# Usage

```
hushWarning(expr, regexp)
```

# Arguments

expr an expression

regexp a regexp to be matched (with str\_detect)

#### Value

the result of expr

### **Examples**

```
hushWarning(warning('Beware the rabbit'), 'rabbit')
hushWarning(warning('Beware the rabbit'), 'hedgehog')
```

24 impute

Hypothesis

Describe a linear model hypothesis to be tested

### **Description**

A Hypothesis can be any linear combination of coefficients, compared to zero. Specify it as a character vector that can be parsed to yield the desired equalities ala makeContrasts. A CoefficientHypothesis is a hypothesis for which terms are singly or jointly tested to be zero (generally the case in a t-test or F-test), by dropping coefficients from the model.

### Usage

```
Hypothesis(hypothesis, terms)
```

### **Arguments**

hypothesis a character vector specifying a hypothesis, following makeContrasts, or a char-

acter vector naming coefficients to be dropped.

terms an optional character vector giving the terms (column names from the model.matrix)

out of which the contrasts will be contrasted. If missing then most functions will

attempt to fill this in for you at run time.

### Value

a Hypothesis with a "transformed" component

### See Also

zlm waldTest lrTest

### **Examples**

```
\label{eq:hamilton} $h < - \ Hypothesis('Stim.ConditionUnstim', c('(Intercept)', 'Stim.ConditionUnstim'))$ $h@contrastMatrix$
```

impute

impute missing continuous expression for plotting

# **Description**

If there are no positive observations for a contrast, it is generally not estimible. However, for the purposes of testing we can replace it with the least favorable value with respect to the contrasts that are defined.

### Usage

```
impute(object, groupby)
```

influence.bayesglm 25

# Arguments

object Output of predict

groupby Variables (column names in predict) to group by for imputation (facets of the

plot)

### Value

data.table

# **Examples**

```
##See stat_ell
example(stat_ell)
```

influence.bayesglm

Influence bayesglm object

# Description

The influence function

# Usage

```
## S3 method for class 'bayesglm'
influence(model, do.coef = TRUE, ...)
```

# Arguments

model bayesglm

do.coef see influence.glm

... ignored

### Value

see influence.glm

26 LMERlike-class

invlogit

Inverse of logistic transformation

# Description

Inverse of logistic transformation

### Usage

```
invlogit(x)
```

### **Arguments**

Х

numeric

### Value

numeric

### **Examples**

```
x <- 1:5
invlogit(log(x/(1-x)))</pre>
```

LMERlike-class

Wrapper for lmer/glmer

### **Description**

A horrendous hack is employed in order to do arbitrary likelihood ratio tests: the model matrix is built, the names possibly mangled, then fed in as a symbolic formula to glmer/lmer. This is necessary because there is no (easy) way to specify an arbitrary fixed-effect model matrix in glmer.

# Usage

```
## S4 method for signature 'LMERlike'
update(object, formula., design, keepDefaultCoef = FALSE, ...)
## S4 method for signature 'LMERlike'
vcov(object, which, ...)
## S4 method for signature 'LMERlike'
coef(object, which, singular = TRUE, ...)
## S4 method for signature 'LMERlike'
logLik(object)
```

LMlike-class 27

### **Arguments**

object LMERlike formula. formula

design something coercible to a data. frame

keepDefaultCoef

logical. Should the coefficient names be preserved from object or updated if

the model matrix has changed?

... In the case of vcov, ignored. In the case of update, passed to model.matrix.

which character, one of 'C', 'D'.

singular logical. Should NA coefficients be returned?

#### Value

see the section "Methods (by generic)"

### Methods (by generic)

- update(LMERlike): update the formula or design matrix
- vcov(LMERlike): return the variance/covariance of component which
- coef(LMERlike): return the coefficients. The horrendous hack is attempted to be undone.
- logLik(LMERlike): return the log-likelihood

#### Slots

pseudoMM part of this horrendous hack.

strictConvergence logical (default: TRUE) return results even when the optimizer or \*lmer complains about convergence

optimMsg character record warnings from lme. NA\_character\_ means no warnings.

LMlike-class Linear Model-like Class

# Description

Wrapper around modeling function to make them behave enough alike that Wald tests and Likelihood ratio are easy to do. To implement a new type of zero-inflated model, extend this class. Depending on how different the method is, you will definitely need to override the fit method, and possibly the model.matrix, model.matrix<-, update, coef, vcov, and logLik methods. 28 LMlike-class

#### Usage

```
## S4 method for signature 'LMlike'
summary(object)
## S4 method for signature 'LMlike'
update(object, formula., design, keepDefaultCoef = FALSE, ...)
## S4 method for signature 'LMlike, CoefficientHypothesis'
waldTest(object, hypothesis)
## S4 method for signature 'LMlike,matrix'
waldTest(object, hypothesis)
## S4 method for signature 'LMlike,character'
lrTest(object, hypothesis)
## S4 method for signature 'LMlike, CoefficientHypothesis'
lrTest(object, hypothesis)
## S4 method for signature 'LMlike, Hypothesis'
lrTest(object, hypothesis)
## S4 method for signature 'LMlike, matrix'
lrTest(object, hypothesis)
## S4 method for signature 'GLMlike'
logLik(object)
```

### Arguments

object LMlike formula. formula

design something coercible to a data.frame

keepDefaultCoef

logical. Should the coefficient names be preserved from object or updated if

the model matrix has changed?

... passed to model.matrix

hypothesis one of a CoefficientHypothesis, Hypothesis or contrast matrix.

#### Value

```
see section "Methods (by generic)"
```

# Methods (by generic)

- summary (LMlike): Print a summary of the coefficients in each component.
- update(LMlike): update the formula or design from which the model.matrix is constructed

LMlike-class 29

• waldTest(object = LMlike, hypothesis = CoefficientHypothesis): Wald test dropping single term specified by CoefficientHypothesis hypothesis

- waldTest(object = LMlike, hypothesis = matrix): Wald test of contrast specified by contrast matrix hypothesis
- 1rTest(object = LMlike, hypothesis = character): Likelihood ratio test dropping entire term specified by character hypothesis naming a term in the symbolic formula.
- lrTest(object = LMlike, hypothesis = CoefficientHypothesis): Likelihood ratio test dropping single term specified by CoefficientHypothesis hypothesis
- lrTest(object = LMlike, hypothesis = Hypothesis): Likelihood ratio test dropping single term specified by Hypothesis hypothesis
- lrTest(object = LMlike, hypothesis = matrix): Likelihood ratio test dropping single term specified by contrast matrix hypothesis
- logLik(GLMlike): return the log-likelihood of a fitted model

#### Slots

design a data frame from which variables are taken for the right hand side of the regression

fitC The continuous fit

fitD The discrete fit

**response** The left hand side of the regression

fitted A logical with components "C" and "D", TRUE if the respective component has converged

formula A formula for the regression

fitArgsC

**fitArgsD** Both lists giving arguments that will be passed to the fitter (such as convergence criteria or case weights)

### See Also

coef

**IrTest** 

waldTest

vcov

logLik

 $\log FC$ 

logFC

Calculate log-fold changes from hurdle model components

### **Description**

Using the delta method, estimate the log-fold change from a state given by a vector contrast0 and the state(s) given by contrast1.

# Usage

```
logFC(zlmfit, contrast0, contrast1)
getLogFC(zlmfit, contrast0, contrast1)
```

### **Arguments**

zlmfit ZlmFit output

contrast0 vector of coefficients giving baseline contrast, or a Hypothesis. If missing, then

the '(Intercept)' is used as baseline.

contrast1 matrix of coefficients giving comparison contrasts, or a Hypothesis. If missing,

then all non-(Intercept) coefficients are compared.

#### **Details**

The log-fold change is defined as follows. For each gene, let u(x) be the expected value of the continuous component, given a covariate x and the estimated coefficients coefC, ie, u(x) = crossprod(x, coefC). Likewise, Let  $v(x) = 1/(1+\exp(-\text{crossprod}(\text{coefD}, x)))$  be the expected value of the discrete component. The log fold change from contrast0 to contrast1 is defined as

```
u(contrast1)v(contrast1) - u(contrast0)v(contrast0).
```

Note that for this to be a log-fold change, then the regression for u must have been fit on the log scale. This is returned in the matrix logFC. An approximation of the variance of logFC (applying the delta method to formula defined above) is provided in varLogFC.

#### Value

list of matrices 'logFC' and 'varLogFC', giving the log-fold-changes for each contrast (columns) and genes (rows) and the estimated sampling variance thereof

#### **Functions**

• getLogFC(): Return results as a perhaps friendlier data.table

logmean 31

#### **Caveats**

1. When method='bayesglm' (the default), it's no longer necessarily true that the log fold change from condition A to B will be the inverse of the log fold change from B to A if the models are fit separately. This is due to the shrinkage in bayesglm.

- 2. The log fold change can be small, but the Hurdle p-value small and significant when the sign of the discrete and continuous model components are discordant so that the marginal log fold change cancels out. The large sample sizes present in many single cell experiments also means that there is substantial power to detect even small changes.
- 3. When there is no expression in a gene for a coefficient that is non-zero in either condition0 or condition1 we return NA because there is not any information to estimate the continuous component. Technically we might return plus or minus infinity, but there is not a straightforward way to estimate a confidence interval in any case. See https://support.bioconductor.org/p/99244/for details

#### See Also

Hypothesis summary,ZlmFit-method

### **Examples**

```
data(vbetaFA)
zz <- zlm( ~ Stim.Condition+Population, vbetaFA[1:5,])</pre>
##log-fold changes in terms of intercept (which is Stim(SEB) and CD154+VbetaResponsive)
lfcStim <- logFC(zz)</pre>
##If we want to compare against unstim, we can try the following
coefnames <- colnames(coef(zz, 'D'))</pre>
contrast0 <- setNames(rep(0, length(coefnames)), coefnames)</pre>
contrast0[c('(Intercept)', 'Stim.ConditionUnstim')] <- 1</pre>
contrast1 <- diag(length(coefnames))</pre>
rownames(contrast1)<-colnames(contrast1)<-coefnames</pre>
contrast1['(Intercept)',]<-1</pre>
lfcUnstim <- logFC(zz, contrast0, contrast1)</pre>
##log-fold change with itself is 0
stopifnot(all(lfcUnstim$logFC[,2]==0))
##inverse of log-fold change with Stim as reference
stopifnot(all(lfcStim$logFC[,1]==(-lfcUnstim$logFC[,1])))
##As a data.table:
getLogFC(zz)
```

logmean

Log mean

#### **Description**

Takes mean of natural scaled values and then logrithm Approximately the inverse operation of expavg Calculates log2(mean(x) + 1)

32 LRT

### Usage

```
logmean(x)
```

### **Arguments**

x numeric

#### Value

numeric

### **Examples**

```
x <- 1:10
expavg(logmean(x))</pre>
```

**LRT** 

Likelihood Ratio Tests for SingleCellAssays

### **Description**

Tests for a change in ET binomial proportion or mean of positive ET Likelihood Ratio Test for SingleCellAssay objects

### Usage

```
LRT(sca, comparison, ...)
## S4 method for signature 'SingleCellAssay,character'
LRT(sca, comparison, referent = NULL, groups = NULL, returnall = FALSE)
```

### **Arguments**

sca A SingleCellAssay class object

comparison A character specifying the factor for comparison

... ignored

referent A character specifying the reference level of comparison.

groups A optional character specifying a variable on which to stratify the test. For

each level of groups, there will be a separate likelihood ratio test.

returnall A logical specifying if additional rows should be returned with information

about the different components of the test.

### Details

Combined Likelihood ratio test (binomial and normal) for SingleCellAssay and derived objects. This function is deprecated, please use lrTest instead.

lrTest 33

### Value

data.frame

# See Also

zlm ZlmFit

### **Examples**

```
data(vbetaFA)
LRT(vbetaFA, 'Stim.Condition', 'Unstim')
```

lrTest

Run a likelihood-ratio test

### **Description**

Compares the change in likelihood between the current model and one subject to contrasts tested in hypothesis. hypothesis can be one of a character giving complete factors or terms to be dropped from the model, CoefficientHypothesis giving names of coefficients to be dropped, Hypothesis giving contrasts using the symbolically, or a contrast matrix, with one row for each coefficient in the full model, and one column for each contrast being tested.

### Usage

```
lrTest(object, hypothesis, ...)
```

### **Arguments**

object LMlike or subclass

hypothesis the hypothesis to be tested. See details.

... optional arguments, passed to fitting functions

### Value

array giving test statistics

### See Also

fit

waldTest

Hypothesis

CoefficientHypothesis

### **Examples**

```
#see ZlmFit-class for examples
example('ZlmFit-class')
```

### **Description**

A 3D array with first dimension being the genes, next dimension giving information about the test (the degrees of freedom, Chisq statistic, and P value), and final dimension being the value of these quantities on the discrete, continuous and hurdle (combined) levels.

### Usage

```
## S4 method for signature 'ZlmFit,character'
lrTest(object, hypothesis, ...)
```

### **Arguments**

object ZlmFit hypothesis See Details

... Arguments passed on to zlm

formula a formula with the measurement variable on the LHS and predictors present in colData on the RHS

sca SingleCellAssay object

method character vector, either 'glm', 'glmer' or 'bayesglm'

silent Silence common problems with fitting some genes

ebayes if TRUE, regularize variance using empirical bayes method

ebayesControl list with parameters for empirical bayes procedure. See ebayes.

force Should we continue testing genes even after many errors have occurred? hook a function called on the fit after each gene.

parallel If TRUE and option(mc.cores)>1 then multiple cores will be used in fitting.

LMlike if provided, then the model defined in this object will be used, rather than following the formulas. This is intended for internal use.

onlyCoef If TRUE then only an array of model coefficients will be returned (probably only useful for bootstrapping).

exprs\_values character or integer passed to 'assay' specifying which assay to use for testing

#### Value

3D array

magic\_assay\_names 35

magic\_assay\_names

Default assay returned

### **Description**

Methods in this package operate on log-transformed (multiplicative scale) expression. We attempt to check for this at construction, and then over-ride the assay method to return the "layer" containing such log-transformed data.

#### Usage

```
magic_assay_names()
assay_idx(x)
## S4 method for signature 'SingleCellAssay,missing'
assay(x, i, withDimnames = TRUE, ...)
```

### **Arguments**

x SingleCellAssay

i must be missing for this method to apply

withDimnames

A logical(1), indicating whether the dimnames of the SummarizedExperiment object should be applied (i.e. copied) to the extracted assays. More precisely, setting withDimnames=FALSE in the *getter* returns the assays *as-is* whereas setting withDimnames=FALSE return them with possibly modified dimnames.

Setting withDimnames=FALSE in the *setter* (assays<-) is required when the dimnames on the supplied assays are not identical to the dimnames on the SummarizedExperiment object; it does not influence actual assignment of dimnames to assays (they're always stored as-is).

Note that

assays(x, withDimnames=FALSE) <- assays(x, withDimnames=FALSE)</pre>

is guaranteed to always work and be a no-op. This is not the case if withDimnames=TRUE is used or if withDimnames is not specified.

... passed to parent method

#### **Details**

By default we return the assay whose names, as given by assayNames(x), matches the first element in the vector c('thresh', 'et', 'Et', 'lCount', 'logTPM', 'logCounts', 'logcounts').

### **Functions**

- magic\_assay\_names(): list of names assumed to represent log-transformed data, in order of usage preference
- assay\_idx(): what index is returned by default by 'assay'

36 MAST-defunct

### **Examples**

```
data(vbetaFA)
assay(vbetaFA)[1:3,1:3]
assay(vbetaFA, 'thresh', withDimnames = FALSE) = assay(vbetaFA)*0 - 9
assay(vbetaFA)[1:3, 1:3]
```

maits

MAITs data set, RNASeq

### **Description**

MAITs data set, RNASeq

### **Format**

a list containing an expression matrix (expressionmat), cell cdat and feature fdat.

### See Also

FromMatrix

MAST-defunct

Defunct functions in package 'MAST'

# Description

These functions are defunct or have been renamed.

### Functions (and replacements, if available)

```
filter mast_filter

cData colData

fData mcols

exprs assay

zlm.SingleCellAssay zlm

combine cbind or rbind

deviance_residuals_hook No replacement available, underlying API changed
```

mast\_filter 37

| mast_filter | Filter a SingleCellAssay |  |
|-------------|--------------------------|--|
|             |                          |  |

#### **Description**

Remove, or flag wells that are outliers in discrete or continuous space.

# Usage

```
mast_filter(sc, groups = NULL, filt_control = NULL, apply_filter = TRUE)
burdenOfFiltering(sc, groups, byGroup = FALSE, filt_control = NULL)
```

## **Arguments**

| sc           | The SingleCellAssay object  |
|--------------|---|
| groups       | An optional character naming the grouping variable  |
| filt_control | The list with configuration parameters for the filter.  |
| apply_filter | logical should the filter be applied, or should a matrix of booleans giving if a well would be subject to a filtering criteria be returned? |
| byGroup      | in the case of $burdenOfFiltering$ should the filter be stratified by groups, or only the plotting.   |

#### **Details**

The function filters wells that don't pass filtering criteria described in filt\_control. filt\_control is a list with named elements nOutlier (minimum nmber of outlier cells for a cell to be filtered [default = 2] sigmaContinuous (the z-score outlier threshold for the continuous part of the signal) [default = 7] and sigmaProportion (the z-score outlier threshold for the discrete part of the signal) [default = 7].

If groups is provided, the filtering is calculated within each level of the group, then combined again as output.

## Value

A filtered result

#### **Functions**

• burdenOfFiltering(): plot the proportions of wells are filtered due to different criteria

#### Author(s)

Andrew McDavid

38 meld\_list\_left

## See Also

burdenOfFiltering

# **Examples**

```
data(vbetaFA)
## Split by 'ncells', apply to each component, then recombine
vbeta.filtered <- mast_filter(vbetaFA, groups='ncells')
## Returned as boolean matrix
was.filtered <- mast_filter(vbetaFA, apply_filter=FALSE)
## Wells filtered for being discrete outliers
head(subset(was.filtered, pctout))
burdenOfFiltering(vbetaFA, groups='ncells', byGroup=TRUE)
burdenOfFiltering(vbetaFA, groups='ncells')</pre>
```

meld\_list\_left

Combine lists, preferentially taking elements from x if there are duplicate names

# Description

Combine lists, preferentially taking elements from x if there are duplicate names

# Usage

```
meld_list_left(x, y)
```

# Arguments

```
x list
y list
```

```
MAST:::meld\_list\_left(list(A=1, B=2), list(A = 0))
```

melt.SingleCellAssay 39

```
melt.SingleCellAssay "Melt" a SingleCellAssay matrix
```

#### **Description**

Return a molten (flat) representation, taking the cross-product of the expression values, the colData (column meta data), and the feature data (mcols).

## Usage

```
## S3 method for class 'SingleCellAssay'
melt(data, ..., na.rm = FALSE, value.name = "value")
```

## **Arguments**

```
data SingleCellAssay
... ignored
na.rm ignored
value.name name of 'values' column in returned value
```

#### Value

A data.table, with the cartesian product of the row and column attributes and the expression values

# **Examples**

```
data(vbetaFA)
melt.SingleCellAssay(vbetaFA[1:10,])
as(vbetaFA[1:10,], 'data.table')
```

model.matrix

Model matrix accessor

## **Description**

Model matrix accessor

## Usage

```
model.matrix(object, ...)
## S4 method for signature 'LMlike'
model.matrix(object, ...)
```

40 myBiplot

#### **Arguments**

object LMlike or subclass

... ignored

#### Value

model.matrix if present

# Methods (by class)

• model.matrix(LMlike): return the model.matrix

model.matrix<-

Replace model matrix

# Description

Replace model matrix

## Usage

```
model.matrix(object) <- value</pre>
```

# **Arguments**

object

LMlike or subclass

value

matrix

#### Value

modify object

myBiplot

Makes a nice BiPlot

# Description

Creates a custom BiPlot for visualizing the results of PCA

## Usage

```
myBiplot(pc, colorfactor, scaling = 100, nudge = 1.2, N = 10, dims = 1:2, ...)
```

#### **Arguments**

pc output of prcomp

colorfactor a factor the same length as nrow(pc\$x) to color the points

scaling integer to scale the vectors showing loadings

nudge numeric to offset labels for loadings

N number of variables with longest dim[1] or dim[2] projections to display

dims numeric vector of length 2 indicating which PCs to plot

... passed to plot

#### Value

printed plot

new\_with\_repaired\_slots

Instantiate a class, but warn rather than error for badly named slots

## Description

Instantiate a class, but warn rather than error for badly named slots

#### Usage

```
new_with_repaired_slots(classname, ..., extra)
```

## **Arguments**

classname 'character' naming a class

... slots in 'classname'

extra named list giving other slots in 'classname'

#### Value

```
'new(classname)'
```

```
MAST:::new_with_repaired_slots("SimpleList", listData = list(x = LETTERS), extra = list(elementType = 'character', food = "tasty", beer = "cold"))
```

42 pbootVcov1

# Description

Sample cells with replacement to find bootstrapped distribution of coefficients

## Usage

```
pbootVcov1(cl, zlmfit, R = 99)
bootVcov1(zlmfit, R = 99, boot_index = NULL)
```

#### **Arguments**

cl a cluster object created by makeCluster

zlmfit class ZlmFit

R number of bootstrap replicates

boot\_index list of indices to resample. Only one of R or boot\_index can be offered.

#### Value

```
array of bootstrapped coefficients array of bootstrapped coefficients
```

#### **Functions**

• pbootVcov1(): parallel version of bootstrapping

```
data(vbetaFA)
zlmVbeta <- zlm(~ Stim.Condition, subset(vbetaFA, ncells==1)[1:5,])
#Only run 3 boot straps, which you wouldn't ever want to do in practice...
bootVcov1(zlmVbeta, R=3)</pre>
```

```
plot.thresholdSCRNACountMatrix
```

Plot cutpoints and densities for thresholding

#### **Description**

Plot cutpoints and densities for thresholding

## Usage

```
## S3 method for class 'thresholdSCRNACountMatrix'
plot(x, ask = FALSE, wait.time = 0, type = "bin", indices = NULL, ...)
```

#### Arguments

| x output of thresholdSCRNACountMatrix |
|---------------------------------------|
|---------------------------------------|

ask if TRUE then will prompt before displaying each plot

wait.time pause (in seconds) between each plot

type one or more of the following: 'bin' (plot the genes by the binning used for

thresholding), or 'gene' (plot thresholding by gene – see next argument)

indices if type is equal to 'gene', and is a integer of length 1, then a random sample

of indices genes is taken. If it is NULL, then 10 genes are sampled. If it is a integer vector of length > 1, then it is interpreted as giving a list of indices of

genes to be displayed.

... further arguments passed to plot

#### Value

displays plots

#### **Examples**

```
## See thresholdSCRNACountMatrix
example(thresholdSCRNACountMatrix)
```

plotlrt

Plot a likelihood ratio test object

#### **Description**

Constructs a forest-like plot of signed log10 p-values, possibly adjusted for multiple comparisons adjust can be one of "holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none".

44 plotSCAConcordance

#### Usage

```
plotlrt(lr, adjust = "fdr", thres = 0.1, trunc = 1e-06, groups = NULL)
```

#### **Arguments**

1r output from lrtest, with returnall=FALSE

adjust character, passed along to p.adjust, see below

thres numeric genes with adjusted pvalues above this value are not depicted

trunc numeric p values below this value are truncated at this value

groups character grouping value. If provided, must match groups argument passed to

lrtest. Plots done separately for each group.

#### Value

Constructs a dotplot

#### Author(s)

andrew

plotSCAConcordance

Concordance plots of filtered single vs n-cell assays

#### Description

Plot the average expression value of two subsets of the data. Generally these might be 1 cell and multiple-cell replicates, in which case if the mcols column ncells is set then the averages will be adjusted accordingly. But it could be any grouping.

#### Usage

```
plotSCAConcordance(
    SCellAssay,
    NCellAssay,
    filterCriteria = list(nOutlier = 2, sigmaContinuous = 9, sigmaProportion = 9),
    groups = NULL,
    ...
)
```

#### **Arguments**

SCellAssay is a FluidigmAssay for the 1-cell per well assay NCellAssay is a FluidigmAssay for the n-cell per well assay

filterCriteria is a list of filtering criteria to apply to the SCellAssay and NCellAssay

groups is a character vector naming the group within which to perform filtering. NULL

by default.

... passed to getConcordance

predict.ZlmFit 45

## Value

printed plot

## See Also

getConcordance

## **Examples**

```
data(vbetaFA)
sca1 <- subset(vbetaFA, ncells==1)
sca100 <- subset(vbetaFA, ncells==100)
plotSCAConcordance(sca1, sca100)</pre>
```

predict.ZlmFit

Return predictions from a ZlmFit object.

## **Description**

Return predictions from a ZlmFit object.

## Usage

```
## S3 method for class 'ZlmFit'
predict(object, newdata = NULL, modelmatrix = NULL, ...)
```

# **Arguments**

object A ZlmFit

newdata The data to predict from. Currently ignored, will use the data in the object.

modelmatrix The model matrix specifying the linear combination of coefficients.

... ignored

#### Value

Predictions (on the link scale) and standard errors.

```
##See stat_ell
example(stat_ell)
```

primerAverage

| predicted_ | Sig |  |
|------------|-----|--|

Predicted signatures

# Description

Predicted signatures

#### **Format**

A data frame of predicted gene expresion signatures for stimulated and unstimulated cells.

| pri | merA | vera | ige |
|-----|------|------|-----|

Average expression values for duplicated/redundant genes

## Description

Takes an average, potentially on a different scale given by fun.natural of some genes. The average is then transformed with fun.cycle.

## Usage

```
primerAverage(fd, geneGroups, fun.natural = expavg, fun.cycle = logshift)
```

# Arguments

fd SingleCellAssay or subclass

geneGroups character naming a column in the featureData that keys the duplicates fun.natural transformation to be used to collapse the duplicate expression values

fun.cycle transformation to be used after collapsing

#### Value

averaged version of fd.

#### Note

This code needs to be tested more extensively after a refactoring. Caveat calculator.

print.summaryZlmFit 47

## **Description**

Shows the top 'n' genes by z score on 'by'

# Usage

```
## S3 method for class 'summaryZlmFit'
print(x, n = 2, by = "logFC", ...)
```

## **Arguments**

| X  | output from summary(ZlmFit)   |
|----|---|
| n  | number of genes to show   |
| by | one of 'C' , 'D' or 'logFC' for continuous, discrete and log fold change z-scores for each contrast |
|    | ignored   |

#### Value

prints a pretty table and invisibly returns a data.table representing the table.

## See Also

summary,ZlmFit-method

read.fluidigm Reads a Fluidigm Biomark (c. 2011) raw data file (or set of files)

# Description

This function reads a raw Fluidigm Biomark data file or set of files and constructs a SingleCellAssay (or FluigidmAssay) object. This was written c. 2011 and has not been tested lately. The Biomark format may have changed.

48 read.fluidigm

#### Usage

```
read.fluidigm(
  files = NULL,
  metadata = NULL,
 header.size = 2,
  skip = 8,
  cycle.threshold = 40,
 metadataColClasses = NULL,
 meta.key = NULL,
  idvars = NULL,
  splitby = NULL,
  unique.well.id = "Chamber.ID",
  raw = TRUE,
  assay = NULL,
  geneid = "Assay.Name",
  sample = NULL,
 well = "Well",
 measurement = "X40.Ct",
 measurement.processed = "Ct",
  ncells = "SampleRConc"
)
```

#### **Arguments**

files A character vector of files to read.

A character path and filename of a CSV file containing additional metadata metadata

about the samples

header.size A numeric indicating the number of lines in the header (default 2)

numeric how many lines to skip before reading (default 8) skip

cycle.threshold

The maximum number of PCR cycles performed (default 40) numeric

metadataColClasses

Optional character vector giving the column classes of the metadata file. See

read.table.

Optional character vector that identifies the key column between the metadata meta.key

and the fluidigm data

idvars Optional character vector that defines the set of columns uniquely identifying

a well (unique cell, gene, and condition).

splitby Optional character that defines the column / variable used to split the resulting

data into a list of SingleCellAssay, such that unique levels of splitby each fall into their own SingleCellAssay. Ususally the experimental unit subjected to

different treatments.

unique.well.id The column that uniquely identifies a sample well in the data. Default is "Cham-

ber.ID".

logical flag indicating this is raw data coming off the instrument. Thus we

make some assumptions about the column names that are present.

raw

removeResponse 49

assay character name of a column that uniquely identifies an Assay (i.e. gene). De-

fault is NULL

geneid character names of the column that identifies a gene. Default is "Assay.Name"

sample character name of a column that uniquely identifies a sample

well character name of a column that uniquely identifies a well. Default "Well".

measurement character name of the column that holds the measurement. Default "X40.Ct".

measurement.processed

character one of "Ct", "40-Ct", or "et". If not "Ct", the measurement will be

transformed.

ncells The column with the number of cells in this well.

#### Value

list of SingleCellAssay holding the data.

#### Author(s)

Greg Finak

removeResponse Remove the left hand side (response) from a formula

#### Description

The order of terms will be rearrange to suit R's liking for hierarchy but otherwise the function should be idempotent for

## Usage

```
removeResponse(Formula, warn = TRUE)
```

# **Arguments**

Formula formula

warn Issue a warning if a response variable is found?

## Value

formula

#### Author(s)

Andrew

rstandard.bayesglm

rstandard for bayesglm objects.

## **Description**

rstandard bayesglm object S3 method

## Usage

```
## S3 method for class 'bayesglm'
rstandard(
  model,
  infl = influence(model, do.coef = FALSE),
  type = c("deviance", "pearson"),
  ...
)
```

# Arguments

model bayesglm
infl see rstandard
type see rstandard
... ignored

#### Value

numeric residuals

SceToSingleCellAssay Coerce a SingleCellExperiment to some class defined in MAST

## **Description**

Coerce a SingleCellExperiment to some class defined in MAST

# Usage

```
SceToSingleCellAssay(sce, class = "SingleCellAssay", check_sanity = TRUE)
```

## **Arguments**

sce object inheriting from SingleCellExperiment class character naming the class to be coerced to

check\_sanity (default: TRUE) Set FALSE to override sanity checks that try to ensure that the de-

fault assay is log-transformed and has at least one exact zero. See defaultAssay for details on the "default assay" which is assumed to contain log transformed

data.

se.coef 51

# Value

object of the indicated class.

se.coef

Return coefficient standard errors

## **Description**

Given a fitted model, return the standard errors of the coefficient

# Usage

```
se.coef(object, ...)
```

# **Arguments**

object a model implementing vcov ... passed to methods

## Value

vector or matrix

## See Also

ZlmFit-class

# **Examples**

```
#see ZlmFit-class for examples
example('ZlmFit-class')
```

show, LMlike-method

show

## **Description**

Display info

# Usage

```
## S4 method for signature 'LMlike'
show(object)
## S4 method for signature 'ZlmFit'
show(object)
```

## **Arguments**

object an object of some type

#### **Details**

Prints information on a LMlike object

#### Value

side effect of printing to console

## Methods (by class)

• show(ZlmFit): print info on ZlmFit

## Description

Splits a SingleCellAssay into a list by a factor (or something coercible into a factor) or a character giving a column of colData(x)

## Usage

```
## S4 method for signature 'SingleCellAssay,character'
split(x, f, drop = FALSE, ...)
```

## **Arguments**

x SingleCellAssay

f length-1 character, or atomic of length ncol(x)

drop unused factor levels

... ignored

## Value

List

```
data(vbetaFA)
split(vbetaFA, 'ncells')
fa <- as.factor(colData(vbetaFA)$ncells)
split(vbetaFA, fa)</pre>
```

stat\_ell 53

stat\_ell

Plot confidence ellipse in 2D

# Description

The focus of the ellipse will be the point (x, y) and semi-major axes aligned with the coordinate axes and scaled by xse, yse and the level.

## Usage

```
stat_ell(
  mapping = NULL,
  data = NULL,
  geom = "polygon",
  position = "identity",
  na.rm = FALSE,
  show.legend = NA,
  inherit.aes = TRUE,
  fill = NA,
  level = 0.95,
  lty = 2,
  invert = FALSE,
  alpha = 1,
  ...
)
```

## **Arguments**

| mapping  | Set of aesthetic mappings created by aes or aes If specified and inherit.aes = TRUE (the default), it is combined with the default mapping at the top level of the plot. You must supply mapping if there is no plot mapping.  |
|----------|--|
| data     | The data to be displayed in this layer. There are three options: If NULL, the default, the data is inherited from the plot data as specified in the call to ggplot. A data.frame, or other object, will override the plot data. All objects will be fortified to produce a data frame. See fortify for which variables will be created. A function will be called with a single argument, the plot data. The return value must be a data.frame., and will be used as the layer data. |
| geom     | The geometric object to use display the data   |
| position | Position adjustment, either as a string, or the result of a call to a position adjust-   |

ment function.

na.rm If FALSE (the default), removes missing values with a warning. If TRUE

silently removes missing values.

logical. Should this layer be included in the legends? NA, the default, includes if any aesthetics are mapped. FALSE never includes, and TRUE always includes.

 $\verb|show.legend|$ 

| inherit.aes | If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn't inherit behaviour from the default plot specification, e.g. borders. |
|-------------|---|
| fill        | A color or aesthetic mapping to fill color. Defaults to NA for empty ellipses.  |
| level       | The confidence level at which to draw an ellipse (default is level=0.95).   |
| lty         | The linetype to use. Can map to a variable. Defaults to 2 (dashed line)   |
| invert      | vector of length 1 that should either be "x", "y", or TRUE. Specifies whether to plot the estimates from the discrete component on the inverse logit scale. invert specifies which axis to invert.  |
| alpha       | transparency  |
|             | other arguments passed on to layer. These are often aesthetics, used to set an aesthetic to a fixed value, like color = "red" or size = 3. They may also be parameters to the paired geom/stat.   |

## Value

ggplot layer

## **Examples**

```
data(vbetaFA)
library(ggplot2)
zlmCond <- zlm(~Stim.Condition, vbetaFA[1:10,])
MM <- model.matrix(~Stim.Condition,unique(colData(vbetaFA)[,c("Stim.Condition"),drop=FALSE]))
predicted <- predict(zlmCond,modelmatrix=MM)
plt <- ggplot(predicted)+aes(x=invlogit(etaD),y=muC,xse=seD,yse=seC,col=sample)+
    facet_wrap(~primerid,scales="free_y")+theme_linedraw()+
    geom_point(size=0.5)+scale_x_continuous("Proportion expression")+
    scale_y_continuous("Estimated Mean")+
    stat_ell(aes(x=etaD,y=muC),level=0.95, invert='x')
## plot with inverse logit transformed x-axis
print(plt)
# doesn't do anything in this case because there are no inestimable coefficients
predictI <- impute(predicted, groupby='primerid')</pre>
```

```
subset, SingleCellAssay-method
```

Subset a SingleCellAssay by cells (columns)

#### **Description**

Evaluates the expression in  $\dots$  in the context of colData(x) and returns a subsetted version of x

#### Usage

```
## S4 method for signature 'SingleCellAssay'
subset(x, ...)
```

summarize 55

# Arguments

```
x SingleCellAssay ... expression
```

#### Value

```
{\tt SingleCellAssay}
```

# Examples

```
data(vbetaFA)
subset(vbetaFA, ncells==1)
```

summarize

Return programmatically useful summary of a fit

# Description

Return programmatically useful summary of a fit

# Usage

```
summarize(object, ...)
```

# Arguments

```
object LMlike or subclass
... other arguments
```

## Value

list of parameters characterizing fit

```
summary, GSEATests-method
```

Summarize gene set enrichment tests

## **Description**

```
Returns a data.table with one row per gene set. This data.table contains columns:
```

set name of gene set

cond\_Z Z statistic for continuous component

cont\_P wald P value

cont\_effect difference in continuous regression coefficients between null and test sets (ie, the numerator of the Z-statistic.)

disc Z Z statistic for discrete

disc\_P wald P value

disc\_effect difference in discrete regression coefficients between null and test sets.

combined\_Z combined discrete and continuous Z statistic using Stouffer's method

combined\_P combined P value

combined\_adj FDR adjusted combined P value

#### Usage

```
## S4 method for signature 'GSEATests'
summary(object, ...)
```

## Arguments

object A GSEATests object
... passed to calcZ

#### Value

data.table

#### See Also

gseaAfterBoot

```
## See the examples in gseaAfterBoot
example(gseaAfterBoot)
```

summary, ZlmFit-method Summarize model features from a ZlmFit object

# Description

Returns a data. table with a special print method that shows the top 2 most significant genes by contrast. This data. table contains columns:

primerid the gene

component C=continuous, D=discrete, logFC=log fold change, S=combined using Stouffer's method, H=combined using hurdle method

contrast the coefficient/contrast of interest

ci.hi upper bound of confidence interval

ci.lo lower bound of confidence interval

coef point estimate

**z** z score (coefficient divided by standard error of coefficient)

**Pr(>Chisq)** likelihood ratio test p-value (only if doLRT=TRUE)

Some of these columns will contain NAs if they are not applicable for a particular component or contrast.

#### Usage

```
## S4 method for signature 'ZlmFit'
summary(
   object,
   logFC = TRUE,
   doLRT = FALSE,
   level = 0.95,
   parallel = FALSE,
   ...
)
```

## **Arguments**

| object   | A ZlmFit object   |
|----------|---|
| logFC    | If TRUE, calculate log-fold changes, or output from a call to getLogFC.                                 |
| doLRT    | if TRUE, calculate lrTests on each coefficient, or a character vector of such coefficients to consider. |
| level    | what level of confidence coefficient to return. Defaults to 95 percent.                                 |
| parallel | If TRUE and option(mc.cores)>1 then multiple cores will be used in fitting.                             |
|          | ignored   |

#### Value

```
data.table
```

#### See Also

print.summaryZlmFit

# **Examples**

```
data(vbetaFA)
z <- zlm(~Stim.Condition, vbetaFA[1:5,])
zs <- summary(z)
names(zs)
print(zs)
##Select `datatable` copmonent to get normal print method
zs$datatable
## Can use parallel processing for LRT now
summary(z, doLRT = TRUE, parallel = TRUE)</pre>
```

```
\verb|summary.thresholdSCRNACountMatrix| \\
```

Summarize the effect of thresholding

#### Description

Returns the proportion of (putative) expression, the variance of expressed cells, and -log10 shapiro-wilk tests for normality on the expressed cells

#### **Usage**

```
## $3 method for class 'thresholdSCRNACountMatrix'
summary(object, ...)
## $3 method for class 'summaryThresholdSCRNA'
print(x, ...)
```

#### **Arguments**

```
    object a thresholdSCRNACountMatrix
    ... currently ignored
    x a summaryThresholdSCRNA object, ie output from summary.thresholdSCRNACountMatrix
```

#### Value

a list of statistics on the original data, and thresholded data

## **Functions**

• print(summaryThresholdSCRNA): prints five-number distillation of the statistics and invisibly returns the table used to generate the summary

thresholdSCRNACountMatrix

Threshold a count matrix using an adaptive threshold.

## **Description**

An adaptive threshold is calculated from the conditional mean of expression, based on 10 bins of the genes with similar expression levels. Thresholds are chosen by estimating cutpoints in the bimodal density estimates of the binned data. These density estimates currently exclude the zeros due to complications with how the bandwidth is selected. (If the bandwith is too small, then extra peaks/modes are found and everything goes haywire). If the diagnostic plots don't reveal any bimodal bins, this is probably the reason, and you may not need to threshold since background in the data are exact zeros.

#### Usage

```
thresholdSCRNACountMatrix(
  data_all,
  conditions = NULL,
  cutbins = NULL,
  nbins = 10,
  bin_by = "median",
  qt = 0.975,
  min_per_bin = 50,
  absolute_min = 0,
  data_log = TRUE,
  adj = 1
)
```

# Arguments

| data_all    | matrix of (possibly log-transformed) counts or TPM. Rows are genes and columns are cells.               |
|-------------|---|
| conditions  | Bins are be determined per gene and per condition. Typically contrasts of interest should be specified. |
| cutbins     | vector of cut points.   |
| nbins       | integer number of bins when cutbins is not specified.   |
| bin_by      | character "median", "proportion", "mean"  |
| qt          | when bin_by is "quantile", what quantile should be used to form the bins                                |
| min_per_bin | minimum number of genes within a bin  |

60 vbetaFA

absolute\_min numeric giving a hard threshold below which everything is assumed to be noise data\_log is data\_all log+1 transformed? If so, it will be returned on the (log+1)-scale as well.

adj bandwith adjustment, passed to density

#### Value

list of thresholded counts (on natural scale), thresholds, bins, densities estimated on each bin, and the original data

#### **Examples**

```
data(maits,package='MAST', envir = environment())
sca <- FromMatrix(t(maits$expressionmat[,1:1000]), maits$cdat, maits$fdat[1:1000,])
tt <- thresholdSCRNACountMatrix(assay(sca))
tt <- thresholdSCRNACountMatrix(2^assay(sca)-1, data_log=FALSE)
opar <- par(no.readonly = TRUE)
on.exit(par(opar))
par(mfrow=c(4,2))
plot(tt)</pre>
```

vbeta

Vbeta Data Set

# Description

Vbeta Data Set

#### **Format**

a data frame with 11 columns. Column Ct contains the cycle threshold, with NA denoting that the threshold was never crossed. So it is inversely proportional to the log2 mRNA, and should be negated (and NAs set to zero) if it is used as a expression measurement for a FluidigmAssay.

vbetaFA

Vbeta Data Set, FluidigmAssay

## Description

Vbeta Data Set, FluidigmAssay

## Format

a FluidigmAssay of the vbeta data set.

#### See Also

```
vbeta, FromFlatDF
```

waldTest 61

waldTest Run a Wald test

#### **Description**

Run a Wald tests on discrete and continuous components hypothesis can be one of a character giving complete factors or terms to be dropped from the model, CoefficientHypothesis giving names of coefficients to be dropped, Hypothesis giving contrasts using the symbolically, or a contrast matrix, with one row for each coefficient in the full model, and one column for each contrast being tested.

#### Usage

```
waldTest(object, hypothesis)
```

## **Arguments**

object LMlike or subclass

hypothesis the hypothesis to be tested. See details.

#### Value

array giving test statistics

#### See Also

fit

lrTest

lht

#### **Examples**

```
#see ZlmFit-class for examples
example('ZlmFit-class')
```

## **Description**

A 3D array with first dimension being the genes, next dimension giving information about the test (the degrees of freedom, Chisq statistic, and P value), and final dimension being the value of these quantities on the discrete, continuous and hurdle (combined) levels.

62 zlm

#### Usage

```
## S4 method for signature 'ZlmFit,matrix'
waldTest(object, hypothesis)
```

## **Arguments**

object ZlmFit hypothesis See Details

## Value

3D array

xform

Make matrix of continuous expression values, orthogonal to discrete

## **Description**

This centers each column of mat around the mean of its non-zero values.

## Usage

```
xform(mat, scale = FALSE)
```

#### **Arguments**

matrix (such as produced by exprs)

scale should the columns also be scaled to have unit variance

## Value

matrix

zlm

Zero-inflated regression for SingleCellAssay

## **Description**

For each gene in sca, fits the hurdle model in formula (linear for et>0), logistic for et==0 vs et>0. Return an object of class ZlmFit containing slots giving the coefficients, variance-covariance matrices, etc. After each gene, optionally run the function on the fit named by 'hook'

zlm 63

## Usage

```
zlm(
  formula,
  sca,
  method = "bayesglm",
  silent = TRUE,
  ebayes = TRUE,
  ebayesControl = NULL,
  force = FALSE,
  hook = NULL,
  parallel = TRUE,
  LMlike,
  onlyCoef = FALSE,
  exprs_values = assay_idx(sca)$aidx,
  ...
)
```

# Arguments

| formula       | a formula with the measurement variable on the LHS and predictors present in colData on the RHS   |
|---------------|---|
| sca           | SingleCellAssay object  |
| method        | character vector, either 'glm', 'glmer' or 'bayesglm'   |
| silent        | Silence common problems with fitting some genes   |
| ebayes        | if TRUE, regularize variance using empirical bayes method   |
| ebayesControl | list with parameters for empirical bayes procedure. See ebayes.   |
| force         | Should we continue testing genes even after many errors have occurred?  |
| hook          | a function called on the fit after each gene.   |
| parallel      | If TRUE and option(mc.cores)>1 then multiple cores will be used in fitting.   |
| LMlike        | if provided, then the model defined in this object will be used, rather than following the formulas. This is intended for internal use. |
| onlyCoef      | If TRUE then only an array of model coefficients will be returned (probably only useful for bootstrapping).                             |
| exprs_values  | character or integer passed to 'assay' specifying which assay to use for testing  |
| • • •         | arguments passed to the S4 model object upon construction. For example, fitArgsC and fitArgsD, or coefPrior.                            |

## Value

a object of class ZlmFit with methods to extract coefficients, etc. OR, if data is a data.frame just a list of the discrete and continuous fits.

64 ZImFit-class

#### **Empirical Bayes variance regularization**

The empirical bayes regularization of the gene variance assumes that the precision (1/variance) is drawn from a gamma distribution with unknown parameters. These parameters are estimated by considering the distribution of sample variances over all genes. The procedure used for this is determined from ebayesControl, a named list with components 'method' (one of 'MOM' or 'MLE') and 'model' (one of 'H0' or 'H1') method MOM uses a method-of-moments estimator, while MLE using the marginal likelihood. H0 model estimates the precisions using the intercept alone in each gene, while H1 fits the full model specified by formula

#### See Also

ZlmFit-class, ebayes, GLMlike-class, BayesGLMlike-class

## **Examples**

```
data(vbetaFA)
zlmVbeta <- zlm(~ Stim.Condition, subset(vbetaFA, ncells==1)[1:10,])</pre>
slotNames(zlmVbeta)
#A matrix of coefficients
coef(zlmVbeta, 'D')['CCL2',]
#An array of covariance matrices
vcov(zlmVbeta, 'D')[,,'CCL2']
waldTest(zlmVbeta, CoefficientHypothesis('Stim.ConditionUnstim'))
## Can also provide just a \code{data.frame} instead
data<- data.frame(x=rnorm(500), z=rbinom(500, 1, .3))</pre>
logit.y \leftarrow with(data, x*2 + z*2); mu.y \leftarrow with(data, 10+10*x+10*z + rnorm(500))
y \leftarrow (runif(500) < exp(logit.y)/(1 + exp(logit.y)))*1
y[y>0] \leftarrow mu.y[y>0]
data$y <- y
fit <- zlm(y \sim x+z, data)
summary.glm(fit$disc)
```

ZlmFit-class

An S4 class to hold the output of a call to zlm

## Description

This holds output from a call to zlm. Many methods are defined to operate on it. See below.

# Usage

```
## S4 method for signature 'ZlmFit,CoefficientHypothesis'
lrTest(object, hypothesis, ...)
## S4 method for signature 'ZlmFit,Hypothesis'
lrTest(object, hypothesis, ...)
```

ZImFit-class 65

```
## S4 method for signature 'ZlmFit,matrix'
lrTest(object, hypothesis, ...)

## S4 method for signature 'ZlmFit,CoefficientHypothesis'
waldTest(object, hypothesis)

## S4 method for signature 'ZlmFit,Hypothesis'
waldTest(object, hypothesis)

## S4 method for signature 'ZlmFit'
coef(object, which, ...)

## S4 method for signature 'ZlmFit'
vcov(object, which, ...)

## S4 method for signature 'ZlmFit'
se.coef(object, which, ...)
```

#### **Arguments**

object ZlmFit

hypothesis call to Hypothesis or CoefficientHypothesis or a matrix giving such contrasts.

... ignored

which character vector, one of "C" (continuous) or "D" (discrete) specifying which component should be returned

#### Value

```
see "Methods (by generic)"
```

#### Methods (by generic)

- lrTest(object = ZlmFit, hypothesis = CoefficientHypothesis): Returns an array with likelihood-ratio tests on contrasts defined using CoefficientHypothesis().
- lrTest(object = ZlmFit, hypothesis = Hypothesis): Returns an array with likelihood-ratio tests specified by Hypothesis, which is a Hypothesis.
- lrTest(object = ZlmFit, hypothesis = matrix): Returns an array with likelihood-ratio tests specified by Hypothesis, which is a contrast matrix.
- waldTest(object = ZlmFit, hypothesis = CoefficientHypothesis): Returns an array with Wald Tests on contrasts defined using CoefficientHypothesis().
- waldTest(object = ZlmFit, hypothesis = Hypothesis): Returns an array with Wald Tests on contrasts defined in Hypothesis()
- coef(ZlmFit): Returns the matrix of coefficients for component which.
- vcov(ZlmFit): Returns an array of variance/covariance matrices for component which.
- se.coef(ZlmFit): Returns a matrix of standard error estimates for coefficients on component which.

66 ZlmFit-class

#### Slots

```
coefC matrix of continuous coefficients
coefD matrix of discrete coefficients
vcovC array of variance/covariance matrices for coefficients
vcovD array of variance/covariance matrices for coefficients
LMlike the LmWrapper object used
sca the SingleCellAssay object used
deviance matrix of deviances
loglik matrix of loglikelihoods
df.null matrix of null (intercept only) degrees of freedom
df.resid matrix of residual DOF
dispersion matrix of dispersions (after shrinkage)
dispersionNoShrink matrix of dispersion (before shrinkage)
priorDOF shrinkage weight in terms of number of psuedo-obs
priorVar shrinkage target
converged output that may optionally be set by the underlying modeling function
hookOut a list of length ngenes containing output from a hook function, if zlm was called with one
exprs_values 'character' or 'integer' with the 'assay' used.
```

#### See Also

zlm summary,ZlmFit-method

```
data(vbetaFA)
zlmVbeta <- zlm(~ Stim.Condition+Population, subset(vbetaFA, ncells==1)[1:10,])</pre>
#Coefficients and standard errors
coef(zlmVbeta, 'D')
coef(zlmVbeta, 'C')
se.coef(zlmVbeta, 'C')
#Test for a Population effect by dropping the whole term (a 5 degree of freedom test)
lrTest(zlmVbeta, 'Population')
#Test only if the VbetaResponsive cells differ from the baseline group
lrTest(zlmVbeta, CoefficientHypothesis('PopulationVbetaResponsive'))
# Test if there is a difference between CD154+/Unresponsive and CD154-/Unresponsive.
# Note that because we parse the expression
# the columns must be enclosed in backquotes
# to protect the \quote{+} and \quote{-} characters.
lrTest(zlmVbeta,\ Hypothesis('`PopulationCD154+VbetaUnresponsive`-
        `PopulationCD154-VbetaUnresponsive`'))
waldTest(zlmVbeta, Hypothesis('`PopulationCD154+VbetaUnresponsive` -
        `PopulationCD154-VbetaUnresponsive`'))
```

# **Index**

| assay, SingleCellAssay, missing-method   | applyFlat, 4 assay, 36                                    | <pre>dof,GLMlike-method(dof), 11 dof,LMERlike-method(dof), 11</pre> |
|--|---|---|
| assay_idx (magic_assay_names), 35  BayesGLMlike-class, 5  bootVcov1 (pbootVcov1), 42  burdenOfFiltering (mast_filter), 37  calcZ, 5, 22  ebayes, 12, 34, 63  expavg, 13, 31  expres (MAST-defunct), 36  filter (MAST-defunct), 36  filterLowExpressedGenes, 14 | assay,SingleCellAssay,missing-method                      |   |
| bootVcov1 (pbootVcov1), 42<br>burdenOfFiltering (mast_filter), 37  calcZ, 5, 22  fData (MAST-defunct), 36 filter (MAST-defunct), 36 filterLowExpressedGenes, 14  |   | -   |
| burdenOfFiltering (mast_filter), 37  calcZ, 5, 22  fData (MAST-defunct), 36  filter (MAST-defunct), 36  filterLowExpressedGenes, 14  |   | exprs (MAST-defunct), 36  |
| Carcz, 5, 22   | **  |   |
|  |   | · · · · · · · · · · · · · · · · · · ·                               |
| cData (MAST-defunct), 36 fit, BayesGLMlike, missing-method (fit),  | cData (MAST-defunct), 36                                  | <pre>fit,BayesGLMlike,missing-method(fit),</pre>                    |
| fit, GLMlike, missing-method (fit), 14   | 26  | fit, GLMlike, missing-method (fit), 14                              |
| coef,ZlmFit-method(ZlmFit-class),64 CoefficientHypothesis,65 fit,LMERlike,missing-method(fit),14 fitted_phat(collectResiduals),7   |   | <pre>fitted_phat (collectResiduals), 7</pre>                        |
| CoefficientHypothesis (Hypothesis), 24 colData, 36 FluidigmAssay, 16 FluidigmAssay (FromFlatDF), 16  |   | - · · · · · · · · · · · · · · · · · · ·                             |
| colData<-,SingleCellAssay,DataFrame-method, freq, 15   | <pre>colData&lt;-,SingleCellAssay,DataFrame-method,</pre> | **  |
| collectResiduals, 7 FromMatrix, 17, 36   | collectResiduals, 7                                       |   |
| combined_residuals_hook getConcordance, 18 (collectResiduals), 7 getLogFC (logFC), 30  |   |   |
| computeEffromCt, 8 getrc (getConcordance), 18  | · · · · · · · · · · · · · · · · · · ·                     | getrc (getConcordance), 18  |
| condSd (freq), 15 getwellKey, 20   | condSd (freq), 15   |   |
| continuous_residuals_hook  | (collectResiduals), 7                                     |   |
| convertMASTClassicToSingleCellAssay, 9 CovFromBoots, 10  CovFromBoots, 10  (getWellNey), 20 getwss (getConcordance), 18 GLMlike-class, 20  |   | getwss (getConcordance), 18   |
| defaultAssay 16 18 50 gsea_control (gseaAfterBoot), 21   | defaultAssav. 16. 18. 50                                  | gsea_control (gseaAfterBoot), 21                                    |
| defaultAssay (magic_assay_names), 35 gseaArterBoot, 21 GSEATests=class 23  | defaultAssay (magic_assay_names), 35                      | - ·   |
| defaultPrior, 11 deviance_residuals_hook hushWarning, 23   | deviance_residuals_hook                                   | hushWarning, 23   |
| (collectResiduals), 7 Hypothesis, 24, 30, 31, 65 discrete_residuals_hook   |   | Hypothesis, 24, 30, 31, 65  |
| (collectResiduals), 7 impute, 24 dof, 11 influence.bayesglm, 25  | (collectResiduals), 7                                     | •   |

68 INDEX

| influence.glm, 25                                     | pbootVcov1, 42                                       |
|---|--|
| invlogit, 26  | plot.thresholdSCRNACountMatrix,43                    |
|   | plotlrt, 43  |
| LMERlike-class, 26                                    | plotSCAConcordance, 19, 44                           |
| LMlike-class, 27                                      | predict.ZlmFit, 45                                   |
| logFC, 30   | predicted_sig, 46                                    |
| logLik,GLMlike-method(LMlike-class),27                | primerAverage, 46                                    |
| logLik,LMERlike-method                                | print.summaryThresholdSCRNA                          |
| (LMERlike-class), 26                                  | (summary.thresholdSCRNACountMatrix),                 |
| logmean, 31   | 58   |
| LRT, 32   | print.summaryZlmFit,47                               |
| LRT,SingleCellAssay,character-method                  | ,  |
| (LRT), 32   | rbind, <i>36</i>                                     |
| lrTest, 32, 33  | read.fluidigm, 47                                    |
| lrTest,LMlike,character-method                        | read.table,48  |
| (LMlike-class), 27                                    | removeResponse, 49                                   |
| lrTest,LMlike,CoefficientHypothesis-method            | rstandard, 50  |
| (LMlike-class), 27                                    | rstandard.bayesglm, 50                               |
| lrTest,LMlike,Hypothesis-method                       | <b>3 3</b> 7   |
| (LMlike-class), 27                                    | SceToSingleCellAssay, 50                             |
| lrTest,LMlike,matrix-method                           | se.coef, 51  |
| (LMlike-class), 27                                    | se.coef,ZlmFit-method(ZlmFit-class),64               |
| lrTest,ZlmFit,character-method,34                     | show, LMlike-method, 51                              |
| <pre>lrTest,ZlmFit,CoefficientHypothesis-method</pre> | show,ZlmFit-method                                   |
| (ZlmFit-class), 64                                    | (show,LMlike-method),51                              |
| lrTest,ZlmFit,Hypothesis-method                       | SingleCellAssay (FromFlatDF), 16                     |
| (ZlmFit-class), 64                                    | split, SingleCellAssay, character-method,            |
| lrTest,ZlmFit,matrix-method                           | 52   |
| (ZlmFit-class), 64                                    | split,SingleCellAssay,factor-method                  |
| (12 10 01.000), 0.                                    | <pre>(split,SingleCellAssay,character-method),</pre> |
| magic_assay_names,35                                  | 52   |
| maits, 36   | split, SingleCellAssay, list-method                  |
| MAST (MAST-package), 4                                | <pre>(split,SingleCellAssay,character-method),</pre> |
| MAST-defunct, 36                                      | 52   |
| MAST-package, 4                                       | stat_ell,53  |
| mast_filter, 37                                       | subset, SingleCellAssay-method, 54                   |
| mcols, 36   | summarize, 55  |
| meld_list_left,38                                     | summary.GSEATests-method.56                          |
| melt.SingleCellAssay, 39                              | summary, LMlike-method (LMlike-class), 27            |
| model.matrix,39                                       | summary, ZlmFit-method, 31, 57                       |
| model.matrix,LMlike-method                            | summary-ZlmFit,                                      |
| (model.matrix), 39                                    | (summary,ZlmFit-method), 57                          |
| model.matrix<-,40                                     | summary.thresholdSCRNACountMatrix, 58                |
| myBiplot, 40  | Summary: em constacon mocuraria en 1x, 50            |
|   | thresholdSCRNACountMatrix, 59                        |
| new_with_repaired_slots,41                            | ,  |
| numexp(freq), 15                                      | update,LMERlike-method                               |
| •               | (LMERlike-class), 26                                 |
| partialScore(collectResiduals),7                      | update,LMlike-method(LMlike-class), 27               |

INDEX 69

```
vbeta, 60, 60
vbetaFA, 60
vcov, GLMlike-method (GLMlike-class), 20
vcov, LMERlike-method (LMERlike-class),
vcov,ZlmFit-method(ZlmFit-class),64
waldTest, 61
waldTest, LMlike, CoefficientHypothesis-method
        (LMlike-class), 27
waldTest,LMlike,matrix-method
        (LMlike-class), 27
{\tt waldTest,ZlmFit,CoefficientHypothesis-method}
        (ZlmFit-class), 64
waldTest,ZlmFit,Hypothesis-method
        (ZlmFit-class), 64
waldTest,ZlmFit,matrix-method,61
xform, 62
zlm, 34, 36, 62
ZlmFit (ZlmFit-class), 64
ZlmFit-class, 64
ZlmFit-summary,
        (summary,ZlmFit-method), 57
```