Package: seqgendiff (via r-universe)

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Type Package

Title RNA-Seq Generation/Modification for Simulation

Version 1.2.4

Description Generates/modifies RNA-seq data for use in simulations. We provide a suite of functions that will add a known amount of signal to a real RNA-seq dataset. The advantage of using this approach over simulating under a theoretical distribution is that common/annoying aspects of the data are more preserved, giving a more realistic evaluation of your method. The main functions are select_counts(), thin_diff(), thin_lib(), thin_gene(), thin_2group(), thin_all(), and effective_cor(). See Gerard (2020) <doi:10.1186/s12859-020-3450-9> for details on the implemented methods.

License GPL-3

Encoding UTF-8

URL https://github.com/dcgerard/seqgendiff

BugReports https://github.com/dcgerard/seqgendiff/issues

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Imports assertthat, irlba, sva, pdist, matchingR, clue

VignetteBuilder knitr

Repository https://dcgerard.r-universe.dev

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seqgendiff-package seqgendiff: RNA-Seq Generation/Modification for Simulation

Description

This package is designed to take real RNA-seq data and alter it by adding a known amount of signal. You can then use this modified dataset in simulation studies for differential expression analysis, factor analysis, confounder adjustment, or library size adjustment. The advantage of this way of simulating data is that you can see how your method behaves when the simulated data exhibit common (and annoying) features of real data. For example, in the real world data are not normally (or negative binomially) distributed and unobserved confounding is a major issue. This package will simulate data that exhibit these characteristics. The methods used in this package are described in detail in Gerard (2020).

seqgendiff Functions

- select_counts() Subsample the columns and rows of a real RNA-seq count matrix. You would then feed this sub-matrix into one of the thinning functions below.
- thin_diff() The function most users should be using for general-purpose binomial thinning. For the special applications of the two-group model or library/gene thinning, see the functions listed below.
- thin_2group() The specific application of thinning in the two-group model.
- thin_lib() The specific application of library size thinning.
- thin_gene() The specific application of total gene expression thinning.

corassign

- thin_all() The specific application of thinning all counts.
- effective_cor() Returns an estimate of the actual correlation between surrogate variables and a user-specified design matrix.
- ThinDataToSummarizedExperiment() Converts a ThinData object to a SummarizedExperiment object.

ThinDataToDESeqDataSet() Converts a ThinData object to a DESeqDataSet object.

Author(s)

David Gerard

References

Gerard, D (2020). "Data-based RNA-seq simulations by binomial thinning." BMC Bioinformatics. 21(1), 206. doi:10.1186/s1285902034509.

See Also

Useful links:

- https://github.com/dcgerard/seqgendiff
- Report bugs at https://github.com/dcgerard/seqgendiff/issues

corassign

Group assignment that is correlated with latent factors.

Description

We extract latent factors from the log of mat using an SVD, then generate an underlying groupassignment variable from a conditional normal distribution (conditional on the latent factors). This underlying group-assignment variable is used to assign groups.

Usage

```
corassign(mat, nfac = NULL, corvec = NULL, return = c("group", "full"))
```

Arguments

mat	A matrix of count data. The rows index the individuals and the columns index the genes.
nfac	The number of latent factors. If NULL, then we will use the method of Onatski (2010) to choose the number of latent factors.
corvec	The vector of correlations. corvec[i] is the correlation between latent factor i and the underlying group-assignment variable. You can think of the correla- tions in corvec as a kind of "tetrachoric correlation." If NULL, then it assumes independence between factors and group assignment. Note that the correlations of the latent factors with the observed group-assignment vector (instead of the latent group-assignment vector) will be corvec * sqrt(2 / pi).

return What should we return? Just the group assignment ("group") or a list of a bunch of things ("full").

Details

If nfac is provided, then corvec must be the same length as nfac. If nfac is not provided, then it is assumed that the first nfac elements of corvec are the underlying correlations, if nfac turns out to be smaller than the length of corvec. If nfac turns out to be larger than the length of corvec, then the factors without defined correlations are assumed to have correlation 0.

Value

A list with some or all of the following elements:

- x The vector of group assignments. 0L indicates membership to one group and 1L indicates membership to the other group.
- nfac The number of assumed latent factors.
- facmat A matrix, whose columns contain the latent factors.
- groupfac The underlying group-assignment factor.
- corvec The correlation vector. Note that this is the correlation between random variables observed in groupfac and facmat,

If return = "group", then the list only contains x.

Author(s)

David Gerard

References

• A. Onatski (2010), Determining the number of factors from empirical distribution of eigenvalues. The Review of Economics and Statistics 92(4).

effective_cor

```
return = "full")
```

Correlation between facmat and groupfac should be about 0.9 cor(cout\$facmat, cout\$groupfac)

```
## Correlation between facmat and x should be about 0.9 * sqrt(2 / pi)
cor(cout$facmat, cout$x)
corvec * sqrt(2 / pi)
```

effective_cor Estimates the effective correlation.

Description

Will return the estimated correlation between the design matrix and the surrogate variables when you assign a target correlation. The method is described in detail in Gerard (2020).

Usage

```
effective_cor(
   design_perm,
   sv,
   target_cor,
   calc_first = c("cor", "mean"),
   method = c("hungarian", "marriage"),
   iternum = 1000
)
```

Arguments

design_perm	A numeric design matrix whose rows are to be permuted (thus controlling the
	amount by which they are correlated with the surrogate variables). The rows
	index the samples and the columns index the variables. The intercept should not
	be included (though see Section "Unestimable Components").

```
sv A matrix of surrogate variables
```

target_cor A numeric matrix of target correlations between the variables in design_perm and the surrogate variables. The rows index the observed covariates and the columns index the surrogate variables. That is, target_cor[i, j] specifies the target correlation between the ith column of design_perm and the jth surrogate variable. The surrogate variables are estimated either using factor analysis or surrogate variable analysis (see the parameter use_sva). The number of columns in target_cor specifies the number of surrogate variables. Set target_cor to NULL to indicate that design_perm and the surrogate variables are independent.

calc_first	Should we calculate the correlation of the mean design_perm and sv (calc_first = "mean"), or should we calculate the mean of the correlations between design_perm and sv (calc_first = "cor")? This should only be changed by expert users.
method	Should we use the Gale-Shapley algorithm for stable marriages ("marriage") (Gale and Shapley, 1962) as implemented in the matchingR package, or the Hungarian algorithm (Papadimitriou and Steiglitz, 1982) ("hungarian") as implemented in the clue package (Hornik, 2005)? The Hungarian method almost always works better, so is the default.
iternum	The total number of simulated correlations to consider.

Details

This function permutes the rows of design_perm many times, each time calculating the Pearson correlation between the columns of design_perm and the columns of sv. It then returns the averages of these Pearson correlations. The permutation is done using permute_design.

Value

A matrix of correlations. The rows index the observed covariates and the columns index the surrogate variables. Element (i, j) is the estimated correlation between the ith variable in design_perm and the jth variable in sv.

Author(s)

David Gerard

References

- Gale, David, and Lloyd S. Shapley. "College admissions and the stability of marriage." *The American Mathematical Monthly* 69, no. 1 (1962): 9-15. doi:10.1080/00029890.1962.11989827.
- Gerard, D (2020). "Data-based RNA-seq simulations by binomial thinning." BMC Bioinformatics. 21(1), 206. doi:10.1186/s1285902034509.
- Hornik K (2005). "A CLUE for CLUster Ensembles." Journal of Statistical Software, 14(12). doi:10.18637/jss.v014.i12. doi:10.18637/jss.v014.i12.
- C. Papadimitriou and K. Steiglitz (1982), Combinatorial Optimization: Algorithms and Complexity. Englewood Cliffs: Prentice Hall.

```
sv = sv,
target_cor = target_cor,
iternum = 10)
```

```
est_sv
```

Estimate the surrogate variables.

Description

This will use either sva or an SVD on the residuals of a regression of mat on design_obs to estimate the surrogate variables.

Usage

est_sv(mat, n_sv, design_obs, use_sva = FALSE)

Arguments

mat	A numeric matrix of RNA-seq counts. The rows index the genes and the columns index the samples.
n_sv	The number of surrogate variables.
design_obs	A numeric matrix of observed covariates that are NOT to be a part of the signal generating process. Only used in estimating the surrogate variables (if target_cor is not NULL). The intercept should <i>not</i> be included (it will sometimes produce an error if it is included).
use_sva	A logical. Should we use surrogate variable analysis (Leek and Storey, 2008) using design_obs to estimate the hidden covariates (TRUE) or should we just do an SVD on log2(mat + 0.5) after regressing out design_obs (FALSE)? Setting this to TRUE allows the surrogate variables to be correlated with the observed covariates, while setting this to FALSE assumes that the surrogate variables are orthogonal to the observed covariates. This option only matters if design_obs is not NULL. Defaults to FALSE.

Value

A matrix of estimated surrogate variables. The columns index the surrogate variables and the rows index the individuals. The surrogate variables are centered and scaled to have mean 0 and variance 1.

Author(s)

David Gerard

fix_cor

Description

Shrinks the target correlation using a uniform scaling factor so that the overall correlation matrix is positive semi-definite. The method is described in detail in Gerard (2020).

Usage

fix_cor(design_perm, target_cor, num_steps = 51)

Arguments

design_perm	A numeric design matrix whose rows are to be permuted (thus controlling the amount by which they are correlated with the surrogate variables). The rows index the samples and the columns index the variables. The intercept should <i>not</i> be included (though see Section "Unestimable Components").
target_cor	A numeric matrix of target correlations between the variables in design_perm and the surrogate variables. The rows index the observed covariates and the columns index the surrogate variables. That is, target_cor[i, j] specifies the target correlation between the ith column of design_perm and the jth surro- gate variable. The surrogate variables are estimated either using factor anal- ysis or surrogate variable analysis (see the parameter use_sva). The number of columns in target_cor specifies the number of surrogate variables. Set target_cor to NULL to indicate that design_perm and the surrogate variables are independent.
num_steps	The number of steps between 0 and 1 to take in the grid search for the shrinkage factor. The step-size would be 1 / (num_steps - 1).

Details

Let $W = cor(design_perm)$. Let $R = target_cor$. Then the overall correlation matrix is:

$$\left(\begin{array}{cc} W & R \\ R' & I_K \end{array}\right).$$

This function applies a multiplicative scaling factor to R until the above matrix is positive semidefinite. That is, it finds a between 0 and 1 such that

$$\left(\begin{array}{cc} W & aR \\ aR' & I_K \end{array}\right)$$

is positive semi-definite.

Value

A matrix of correlations the same dimension as target_cor. Actually, the returned matrix is a * target_cor, where a was determined to make the overall correlation matrix positive semi-definite.

Author(s)

David Gerard

References

Gerard, D (2020). "Data-based RNA-seq simulations by binomial thinning." BMC Bioinformatics. 21(1), 206. doi:10.1186/s1285902034509.

Examples

```
n <- 10
design_perm <- matrix(rep(c(0, 1), length.out = n))
target_cor <- matrix(seq(1, 0, length.out = 10), nrow = 1)
new_cor <- seqgendiff:::fix_cor(design_perm = design_perm, target_cor = target_cor)
new_cor / target_cor
## In the case of one observed covariate, the requirement is just that
## the sum of squared correlations is less than or equal to one.
sum(target_cor ^ 2)
sum(new_cor ^ 2)
```

permute_design	Permute the design matrix so that it is approximately correlated with
	the surrogate variables.

Description

Permute the design matrix so that it is approximately correlated with the surrogate variables.

Usage

```
permute_design(
   design_perm,
   sv,
   target_cor,
   method = c("hungarian", "marriage")
)
```

Arguments

design_perm	A numeric design matrix whose rows are to be permuted (thus controlling the
	amount by which they are correlated with the surrogate variables). The rows
	index the samples and the columns index the variables. The intercept should not
	be included (though see Section "Unestimable Components").
sv	A matrix of surrogate variables

target_cor	A numeric matrix of target correlations between the variables in design_perm and the surrogate variables. The rows index the observed covariates and the columns index the surrogate variables. That is, target_cor[i, j] specifies the target correlation between the ith column of design_perm and the jth surro- gate variable. The surrogate variables are estimated either using factor anal- ysis or surrogate variable analysis (see the parameter use_sva). The number of columns in target_cor specifies the number of surrogate variables. Set target_cor to NULL to indicate that design_perm and the surrogate variables are independent.
method	Should we use the Gale-Shapley algorithm for stable marriages ("marriage") (Gale and Shapley, 1962) as implemented in the matchingR package, or the Hungarian algorithm (Papadimitriou and Steiglitz, 1982) ("hungarian") as implemented in the clue package (Hornik, 2005)? The Hungarian method almost always works better, so is the default.

Value

A list with two elements:

design_perm A row-permuted version of the user-provided design_perm.

latent_var A matrix of the latent variables on which design_perm was matched.

Author(s)

David Gerard

References

- Gale, David, and Lloyd S. Shapley. "College admissions and the stability of marriage." *The American Mathematical Monthly* 69, no. 1 (1962): 9-15. doi:10.1080/00029890.1962.11989827.
- C. Papadimitriou and K. Steiglitz (1982), Combinatorial Optimization: Algorithms and Complexity. Englewood Cliffs: Prentice Hall.
- Hornik K (2005). "A CLUE for CLUster Ensembles." *Journal of Statistical Software*, 14(12). doi:10.18637/jss.v014.i12. doi:10.18637/jss.v014.i12.

poisthin

Apply Poisson thinning to a matrix of count data.

Description

This is now defunct. Please try out select_counts and thin_2group.

poisthin

Usage

```
poisthin(
  mat,
  nsamp = nrow(mat),
  ngene = ncol(mat),
  gselect = c("max", "random", "rand_max", "custom", "mean_max"),
  gvec = NULL,
  skip_gene = 0L,
  signal_fun = stats::rnorm,
  signal_params = list(mean = 0, sd = 1),
  prop_null = 1,
  alpha = 0,
  group_assign = c("frac", "random", "cor"),
  group_prop = 0.5,
  corvec = NULL
)
```

Arguments

mat	A matrix of count data. The rows index the individuals and the columns index the genes.
nsamp	The number of samples to select from mat.
ngene	The number of genes to select from mat.
gselect	How should we select the subset of genes? Should we choose the ngene most median expressed genes ("max"), a random sample of the genes ("random"), a random sample of the most expressed genes ("rand_max"), a user-provided list ("custom"), or by maximum mean expression level ("mean_max")? If "custom", then gvec should be specified. Expression levels of a gene are measured by median expression across individuals with ties broken by mean expression.
gvec	A logical of length ncol(mat). A TRUE in position i indicates inclusion into the smaller dataset. Hence, sum(gvec) should equal ngene.
skip_gene	The number of maximally expressed genes to skip. Not used if gselect = "custom".
signal_fun	A function that returns the signal. This should take as input n for the number of samples to return and then return only a vector of samples.
signal_params	A list of additional arguments to pass to signal_fun.
prop_null	The proportion of genes that are null.
alpha	If b is an effect and s is an empirical standard deviation, then we model b/s^{α} as being exchangeable.
group_assign	How should we assign groups? Exactly specifying the proportion of individuals in each group ("frac"), with a Bernoulli distribution ("random"), or correlated with latent factors ("cor")? If group_assign = "cor", then you have to spec- ify corvec. If group_assign = "frac" or group_assign = "random", then the proportion of samples in each group is specified with the group_prop argument.

group_prop	The proportion of individuals that are in group 1. This proportion is determinis- tic if group_assign = "frac", and is the expected proportion if group_assign = "random". This argument is not used if group_assign = "cor".
corvec	A vector of correlations. corvec[i] is the correlation of the latent group as- signment vector with the ith latent confounder. Only used if group_assign = "cor". This vector is constrained so that crossprod(corvec) < 1. The number of latent factors is taken to be the length of corvec. Note that the correlations of the latent factors with the observed group-assignment vector (instead of the latent group-assignment vector) will be corvec * sqrt(2 / pi).

Details

Given a matrix of RNA-seq counts, this function will randomly select two groups of samples and add signal to a known proportion of the genes. This signal is the log (base 2) effect size of the group indicator in a linear model. The user may specify the distribution of the effects.

The Poisson thinning approach first randomly assigns samples to be in one of two groups. Then, given this assignment, will Binomially sample counts with a sample size of the gene expression counts and a probability that is a function of the effect size. For details, see Gerard and Stephens (2021).

Value

A list with the following elements:

Y A matrix of altered counts with nsamp rows and ngene columns.

X A design matrix. The first column contains a vector ones (for an intercept term) and the second column contains an indicator for group membership.

beta The approximately true effect sizes of $log(Y) X\beta$.

corassign The output from the call to corassign. Only returned if group_assign = "cor".

Author(s)

David Gerard

References

• Gerard, D., and Stephens, M. (2021). "Unifying and Generalizing Methods for Removing Unwanted Variation Based on Negative Controls." Statistica Sinica, 31(3), 1145-1166 doi:10.5705/ss.202018.0345.

```
## Simulate data from given matrix of counts
## In practice, you would obtain Y from a real dataset, not simulate it.
set.seed(1)
nsamp <- 10
ngene <- 1000
Y <- matrix(stats::rpois(nsamp * ngene, lambda = 50), nrow = ngene)</pre>
```

```
## Apply thinning
poisout <- poisthin(mat</pre>
                                  = t(Y),
                                  = 9,
                    nsamp
                    ngene
                                  = 999,
                    signal_fun = stats::rnorm,
                    signal_params = list(mean = 0, sd = 1),
                    prop_null
                                  = 0.9)
## Dimension of count matrix is smaller.
dim(poisout$Y)
## Can verify signal was added by estimating it with lm().
betahat <- coef(lm(log2(poisout$Y + 1) ~ poisout$X[, 2]))[2, ]</pre>
plot(poisout$beta, betahat, xlab = "Coefficients", ylab = "Estimates")
abline(0, 1, col = 2, lty = 2)
```

```
select_counts
```

Subsample the rows and columns of a count matrix.

Description

It is a good idea to subsample (each iteration) the genes and samples from a real RNA-seq dataset prior to applying thin_diff (and related functions) so that your conclusions are not dependent on the specific structure of your dataset. This function is designed to efficiently do this for you.

Usage

```
select_counts(
  mat,
  nsamp = ncol(mat),
  ngene = nrow(mat),
  gselect = c("random", "max", "mean_max", "custom"),
  gvec = NULL,
  filter_first = FALSE,
  nskip = 0L
)
```

Arguments

mat	A numeric matrix of RNA-seq counts. The rows index the genes and the columns index the samples.
nsamp	The number of samples (columns) to select from mat.
ngene	The number of genes (rows) to select from mat.
gselect	How should we select the subset of genes? Options include:
	random Randomly select the genes, with each gene having an equal probability of being included in the subsampled matrix.

	max Choose the ngene most median-expressed genes. Ties are broken by mean- expression.
	mean_max Choose the ngene most mean-expressed genes.
	<pre>custom A user-specified list of genes. If gselect = "custom" then gvec needs to be non-NULL.</pre>
gvec	A logical vector of length nrow(mat). A TRUE in position i indicates inclusion into the smaller dataset. Hence, sum(gvec) should equal ngene.
filter_first	Should we first filter genes by the method of Chen et al. (2016) (TRUE) or not (FALSE)? If TRUE then the edgeR package should be installed.
nskip	The number of median-maximally expressed genes to skip. Not used if gselect = "custom".

Details

The samples (columns) are chosen randomly, with each sample having an equal probability of being in the sub-matrix. The genes are selected according to one of four schemes (see the description of the gselect argument).

If you have edgeR installed, then some functionality is provided for filtering out the lowest expressed genes prior to applying subsampling (see the filter_first argument). This filtering scheme is described in Chen et al. (2016). If you want more control over this filtering, you should use the filterByExpr function from edgeR directly. You can install edgeR by following instructions at doi:10.18129/B9.bioc.edgeR.

Value

A numeric matrix, which is a ngene by nsamp sub-matrix of mat. If rownames(mat) is NULL, then the row names of the returned matrix are the indices in mat of the selected genes. If colnames(mat) is NULL, then the column names of the returned matrix are the indices in mat of the selected samples.

Author(s)

David Gerard

References

 Chen, Yunshun, Aaron TL Lun, and Gordon K. Smyth. "From reads to genes to pathways: differential expression analysis of RNA-Seq experiments using Rsubread and the edgeR quasilikelihood pipeline." *F1000Research* 5 (2016). doi:10.12688/f1000research.8987.2.

```
## Simulate data from given matrix of counts
## In practice, you would obtain mat from a real dataset, not simulate it.
set.seed(1)
n <- 100
p <- 1000
mat <- matrix(stats::rpois(n * p, lambda = 50), nrow = p)</pre>
```

summary. ThinData

```
submat <- select_counts(mat = mat, nsamp = 10, ngene = 100)
thout <- thin_2group(mat = submat, prop_null = 0.5)
## The rownames and colnames (if NULL in mat) tell you which genes/samples
## were selected.
rownames(submat)
colnames(submat)</pre>
```

summary.ThinData *Provide summary output of a ThinData S3 object.*

Description

Provide summary output of a ThinData S3 object.

Usage

S3 method for class 'ThinData'
summary(object, ...)

Arguments

object	A ThinData S3 object. This is generally output by either thin_diff, thin_2group,
	or thin_lib.
	Not used.

Value

Returns nothing. Prints out some summary information on object.

Author(s)

David Gerard

ThinDataToDESeqDataSet

Converts a ThinData S3 object into a DESeqDataSet S4 object.

Description

The design formula in the resulting DESeqDataSet is just the sum of all variables in designmat from the ThinData object (except the intercept term). You should change this design formula if you want to study other models.

Usage

ThinDataToDESeqDataSet(obj)

Arguments

obj	A ThinData S3 object. This is generally output by either thin_diff, thin_2group
	thin_lib, thin_gene, or thin_all.

Value

A DESeqDataSet S4 object. This will allow you to insert the simulated data directly into DESeq2.

Author(s)

David Gerard

Examples

```
## Generate simulated data and modify using thin_diff().
## In practice, you would use real data, not simulated.
set.seed(1)
n <- 10
p <- 1000
Z <- matrix(abs(rnorm(n, sd = 4)))</pre>
alpha <- matrix(abs(rnorm(p, sd = 1)))</pre>
mat <- round(2^(alpha %*% t(Z) + abs(matrix(rnorm(n * p, sd = 5),</pre>
                                                 nrow = p,
                                                 ncol = n))))
design_perm <- cbind(rep(c(0, 1), length.out = n), runif(n))</pre>
coef_perm <- matrix(rnorm(p * ncol(design_perm), sd = 6), nrow = p)</pre>
design_obs <- matrix(rnorm(n), ncol = 1)</pre>
target_cor <- matrix(c(0.9, 0))</pre>
thout <- thin_diff(mat</pre>
                                     = mat,
                     design_perm = design_perm,
                     coef_perm = coef_perm,
target_cor = target_cor,
design_obs = design_obs,
                     permute_method = "hungarian")
## Convert ThinData object to DESeqDataSet object.
seobj <- ThinDataToDESeqDataSet(thout)</pre>
class(seobj)
## The "01" variable in the colData corresponds to design_obs.
## The "P1" and "P2" variables in colData correspond to design_perm.
seobj
```

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ThinDataToSummarizedExperiment

Converts a ThinData S3 object into a SummarizedExperiment S4 object.

Description

This only keeps the mat, design_obs, designmat, and coefmat elements of the ThinData object.

Usage

ThinDataToSummarizedExperiment(obj)

Arguments

obj A ThinData S3 object. This is generally output by either thin_diff, thin_2group, thin_lib, thin_gene, or thin_all.

Value

A SummarizedExperiment S4 object. This is often used in Bioconductor when performing differential expression analysis.

Author(s)

David Gerard

```
## Generate simulated data and modify using thin_diff().
## In practice, you would use real data, not simulated.
set.seed(1)
n <- 10
p <- 1000
Z <- matrix(abs(rnorm(n, sd = 4)))</pre>
alpha <- matrix(abs(rnorm(p, sd = 1)))</pre>
mat <- round(2^{(alpha \%\% t(Z) + abs(matrix(rnorm(n * p, sd = 5)),
                                              nrow = p,
                                              ncol = n))))
design_perm <- cbind(rep(c(0, 1), length.out = n), runif(n))</pre>
coef_perm <- matrix(rnorm(p * ncol(design_perm), sd = 6), nrow = p)</pre>
design_obs <- matrix(rnorm(n), ncol = 1)</pre>
target_cor <- matrix(c(0.9, 0))</pre>
thout <- thin_diff(mat</pre>
                                   = mat,
                    design_perm = design_perm,
                    coef_perm
                                 = coef_perm,
                    target_cor
                                 = target_cor,
                    design_obs
                                 = design_obs,
                    permute_method = "hungarian")
```

```
## Convert ThinData object to SummarizedExperiment object.
seobj <- ThinDataToSummarizedExperiment(thout)
class(seobj)
## The "01" variable in the colData corresponds to design_obs.
## The "P1" and "P2" variables in colData correspond to design_perm.
seobj
```

thin_2group

```
Binomial thinning in the two-group model.
```

Description

Given a matrix of real RNA-seq counts, this function will randomly assign samples to one of two groups, draw the log2-fold change in expression between two groups for each gene, and add this signal to the RNA-seq counts matrix. The user may specify the proportion of samples in each group, the proportion of null genes (where the log2-fold change is 0), and the signal function. This is a specific application of the general binomial thinning approach implemented in thin_diff.

Usage

```
thin_2group(
  mat,
  prop_null = 1,
  signal_fun = stats::rnorm,
  signal_params = list(mean = 0, sd = 1),
  group_prop = 0.5,
  corvec = NULL,
  alpha = 0,
  permute_method = c("hungarian", "marriage"),
  type = c("thin", "mult")
)
```

Arguments

mat	A numeric matrix of RNA-seq counts. The rows index the genes and the columns index the samples.
prop_null	The proportion of genes that are null.
signal_fun	A function that returns the signal. This should take as input n for the number of samples to return and then return only a vector of samples. Additional parameters may be passed through signal_params.
signal_params	A list of additional arguments to pass to signal_fun.
group_prop	The proportion of individuals that are in group 1.

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corvec	A vector of target correlations. corvec[i] is the target correlation of the latent group assignment vector with the ith surrogate variable. The default is to set this to NULL, in which case group assignment is made independently of any unobserved confounding.
alpha	The scaling factor for the signal distribution from Stephens (2016). If $x_1, x_2,, x_n$ are drawn from signal_fun, then the signal is set to $x_1s_1^{\alpha}, x_2s_2^{\alpha},, x_ns_n^{\alpha}$, where s_g is the empirical standard deviation of gene g. Setting this to 0 means that the effects are exchangeable, setting this to 1 corresponds to the p-value prior of Wakefield (2009). You would rarely set this to anything but 0 or 1.
permute_method	Should we use the Gale-Shapley algorithm for stable marriages ("marriage") (Gale and Shapley, 1962) as implemented in the matchingR package, or the Hungarian algorithm (Papadimitriou and Steiglitz, 1982) ("hungarian") as implemented in the clue package (Hornik, 2005)? The Hungarian method almost always works better, so is the default.
type	Should we apply binomial thinning (type = "thin") or just naive multiplication of the counts (type = "mult"). You should always have this set to "thin".

Details

The specific application of binomial thinning to the two-group model was used in Gerard and Stephens (2018) and Gerard and Stephens (2021). This is a specific case of the general method described in Gerard (2020).

Value

A list-like S3 object of class ThinData. Components include some or all of the following:

mat The modified matrix of counts.

- designmat The design matrix of variables used to simulate signal. This is made by column-binding design_fixed and the permuted version of design_perm.
- coefmat A matrix of coefficients corresponding to designmat.
- design_obs Additional variables that should be included in your design matrix in downstream fittings. This is made by column-binding the vector of 1's with design_obs.
- sv A matrix of estimated surrogate variables. In simulation studies you would probably leave this out and estimate your own surrogate variables.
- cormat A matrix of target correlations between the surrogate variables and the permuted variables in the design matrix. This might be different from the target_cor you input because we pass it through fix_cor to ensure positive semi-definiteness of the resulting covariance matrix.

Author(s)

David Gerard

References

- Gale, David, and Lloyd S. Shapley. "College admissions and the stability of marriage." *The American Mathematical Monthly* 69, no. 1 (1962): 9-15. doi:10.1080/00029890.1962.11989827.
- Gerard, D., and Stephens, M. (2021). "Unifying and Generalizing Methods for Removing Unwanted Variation Based on Negative Controls." *Statistica Sinica*, 31(3), 1145-1166 doi:10.5705/ss.202018.0345.
- David Gerard and Matthew Stephens (2018). "Empirical Bayes shrinkage and false discovery rate estimation, allowing for unwanted variation." *Biostatistics*, doi:10.1093/biostatistics/kxy029.
- Gerard, D (2020). "Data-based RNA-seq simulations by binomial thinning." BMC Bioinformatics. 21(1), 206. doi:10.1186/s1285902034509.
- Hornik K (2005). "A CLUE for CLUster Ensembles." *Journal of Statistical Software*, 14(12). doi:10.18637/jss.v014.i12. doi:10.18637/jss.v014.i12.
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- Stephens, Matthew. "False discovery rates: a new deal." *Biostatistics* 18, no. 2 (2016): 275-294. doi:10.1093/biostatistics/kxw041.
- Wakefield, Jon. "Bayes factors for genome-wide association studies: comparison with P-values." *Genetic epidemiology* 33, no. 1 (2009): 79-86. doi:10.1002/gepi.20359.

See Also

select_counts For subsampling the rows and columns of your real RNA-seq count matrix prior to applying binomial thinning.

thin_diff For the more general thinning approach.

ThinDataToSummarizedExperiment For converting a ThinData object to a SummarizedExperiment object.

ThinDataToDESeqDataSet For converting a ThinData object to a DESeqDataSet object.

```
## Simulate data from given matrix of counts
## In practice, you would obtain Y from a real dataset, not simulate it.
set.seed(1)
nsamp <- 10
ngene <- 1000
Y <- matrix(stats::rpois(nsamp * ngene, lambda = 50), nrow = ngene)
                                 = Y,
thinout <- thin_2group(mat</pre>
                       prop_null
                                     = 0.9,
                       signal_fun = stats::rexp,
                       signal_params = list(rate = 0.5))
## 90 percent of genes are null
mean(abs(thinout$coef) < 10^-6)</pre>
## Check the estimates of the log2-fold change
Ynew <- log2(t(thinout$mat + 0.5))</pre>
```

thin_all

```
X <- thinout$designmat
Bhat <- coef(lm(Ynew ~ X))["X", ]
plot(thinout$coefmat,
        Bhat,
        xlab = "log2-fold change",
        ylab = "Estimated log2-fold change")
abline(0, 1, col = 2, lwd = 2)
```

thin_all

Binomial thinning for altering read-depth.

Description

Given a matrix of real RNA-seq counts, this function will apply a thinning factor uniformly to every count in this matrix. This uniformly lowers the read-depth for the entire dataset. The thinning factor should be provided on the log2-scale. This is a specific application of the binomial thinning approach in thin_diff. Though this particular form of thinning was used by Robinson and Storey (2014) in the context of deriving read-depth suggestions. It is also described in detail in Gerard (2020).

Usage

thin_all(mat, thinlog2, type = c("thin", "mult"))

Arguments

mat	A numeric matrix of RNA-seq counts. The rows index the genes and the columns index the samples.
thinlog2	A numeric scalar. This is the amount to shrink each count in mat (on the log2-scale). For example, a value of 0 means that we do not thin, a value of 1 means that we thin by a factor of 2, a value of 2 means we thin by a factor of 4, etc.
type	Should we apply binomial thinning (type = "thin") or just naive multiplication of the counts (type = "mult"). You should always have this set to "thin".

Value

A list-like S3 object of class ThinData. Components include some or all of the following:

- mat The modified matrix of counts.
- designmat The design matrix of variables used to simulate signal. This is made by column-binding design_fixed and the permuted version of design_perm.
- coefmat A matrix of coefficients corresponding to designmat.
- design_obs Additional variables that should be included in your design matrix in downstream fittings. This is made by column-binding the vector of 1's with design_obs.
- sv A matrix of estimated surrogate variables. In simulation studies you would probably leave this out and estimate your own surrogate variables.

- cormat A matrix of target correlations between the surrogate variables and the permuted variables in the design matrix. This might be different from the target_cor you input because we pass it through fix_cor to ensure positive semi-definiteness of the resulting covariance matrix.

Author(s)

David Gerard

References

- Gerard, D (2020). "Data-based RNA-seq simulations by binomial thinning." BMC Bioinformatics. 21(1), 206. doi:10.1186/s1285902034509.
- Robinson, David G., and John D. Storey. "subSeq: determining appropriate sequencing depth through efficient read subsampling." *Bioinformatics* 30, no. 23 (2014): 3424-3426. doi:10.1093/bioinformatics/btu552.

See Also

- select_counts For subsampling the rows and columns of your real RNA-seq count matrix prior to applying binomial thinning.
- thin_diff For the more general thinning approach.
- thin_lib For thinning sample-wise.
- thin_gene For thinning gene-wise.
- ThinDataToSummarizedExperiment For converting a ThinData object to a SummarizedExperiment object.
- ThinDataToDESeqDataSet For converting a ThinData object to a DESeqDataSet object.

```
## Generate count data and set thinning factor
## In practice, you would obtain mat from a real dataset, not simulate it.
set.seed(1)
n <- 10
p <- 100
lambda <- 1000
mat <- matrix(lambda, ncol = n, nrow = p)
thinlog2 <- 1
## Thin read-depths
thout <- thin_all(mat = mat, thinlog2 = thinlog2)
## Compare empirical and theoretical proportions
mean(thout$mat) / lambda
2 ^ -thinlog2
```

thin_base

Description

Given a matrix of counts (Y) where $log_2(E[Y]) = Q$, a design matrix (X), and a matrix of coefficients (B), thin_diff will generate a new matrix of counts such that $log_2(E[Y]) = BX' + u1' + Q$, where u is some vector of intercept coefficients. This function is used by all other thinning functions. The method is described in detail in Gerard (2020).

Usage

```
thin_base(mat, designmat, coefmat, relative = TRUE, type = c("thin", "mult"))
```

Arguments

mat	A numeric matrix of RNA-seq counts. The rows index the genes and the columns index the samples.
designmat	A design matrix. The rows index the samples and the columns index the variables. The intercept should <i>not</i> be included.
coefmat	A matrix of coefficients. The rows index the genes and the columns index the samples.
relative	A logical. Should we apply relative thinning (TRUE) or absolute thinning (FALSE). Only experts should change the default.
type	Should we apply binomial thinning (type = "thin") or just naive multiplication of the counts (type = "mult"). You should always have this set to "thin".

Value

A matrix of new RNA-seq read-counts. This matrix has the signal added from designmat and coefmat.

Author(s)

David Gerard

References

• Gerard, D (2020). "Data-based RNA-seq simulations by binomial thinning." *BMC Bioinformatics*. 21(1), 206. doi:10.1186/s1285902034509.

See Also

- select_counts For subsampling the rows and columns of your real RNA-seq count matrix prior to applying binomial thinning.
- thin_diff For the function most users should be using for general-purpose binomial thinning.

thin_2group For the specific application of thinning in the two-group model.

thin_lib For the specific application of library size thinning.

thin_gene For the specific application of total gene expression thinning.

thin_all For the specific application of thinning all counts uniformly.

Examples

```
## Simulate data from given matrix of counts
## In practice, you would obtain Y from a real dataset, not simulate it.
set.seed(1)
nsamp <- 10
ngene <- 1000
Y <- matrix(stats::rpois(nsamp * ngene, lambda = 100), nrow = ngene)
X <- matrix(rep(c(0, 1), length.out = nsamp))
B <- matrix(seq(3, 0, length.out = ngene))
Ynew <- thin_base(mat = Y, designmat = X, coefmat = B)</pre>
```

```
## Demonstrate how the log2 effect size is B
Bhat <- coefficients(lm(t(log2(Ynew)) ~ X))["X", ]
plot(B, Bhat, xlab = "Coefficients", ylab = "Coefficient Estimates")
abline(0, 1, col = 2, lwd = 2)</pre>
```

thin_diff

Binomial thinning for differential expression analysis.

Description

Given a matrix of real RNA-seq counts, this function will add a known amount of signal to the count matrix. This signal is given in the form of a Poisson / negative binomial / mixture of negative binomials generalized linear model with a log (base 2) link. The user may specify any arbitrary design matrix and coefficient matrix. The user may also control for the amount of correlation between the observed covariates and any unobserved surrogate variables. The method is described in detail in Gerard (2020).

Usage

```
thin_diff(
  mat,
  design_fixed = NULL,
  coef_fixed = NULL,
  design_perm = NULL,
```

thin_diff

```
coef_perm = NULL,
target_cor = NULL,
use_sva = FALSE,
design_obs = NULL,
relative = TRUE,
change_colnames = TRUE,
permute_method = c("hungarian", "marriage"),
type = c("thin", "mult")
)
```

Arguments

mat	A numeric matrix of RNA-seq counts. The rows index the genes and the columns index the samples.
design_fixed	A numeric design matrix whose rows are fixed and not to be permuted. The rows index the samples and the columns index the variables. The intercept should <i>not</i> be included (though see Section "Unestimable Components").
coef_fixed	A numeric matrix. The coefficients corresponding to design_fixed. The rows index the genes and the columns index the variables.
design_perm	A numeric design matrix whose rows are to be permuted (thus controlling the amount by which they are correlated with the surrogate variables). The rows index the samples and the columns index the variables. The intercept should <i>not</i> be included (though see Section "Unestimable Components").
coef_perm	A numeric matrix. The coefficients corresponding to design_perm. The rows index the genes and the columns index the variables.
target_cor	A numeric matrix of target correlations between the variables in design_perm and the surrogate variables. The rows index the observed covariates and the columns index the surrogate variables. That is, target_cor[i, j] specifies the target correlation between the ith column of design_perm and the jth surro- gate variable. The surrogate variables are estimated either using factor anal- ysis or surrogate variable analysis (see the parameter use_sva). The number of columns in target_cor specifies the number of surrogate variables. Set target_cor to NULL to indicate that design_perm and the surrogate variables are independent.
use_sva	A logical. Should we use surrogate variable analysis (Leek and Storey, 2008) using design_obs to estimate the hidden covariates (TRUE) or should we just do an SVD on log2(mat + 0.5) after regressing out design_obs (FALSE)? Setting this to TRUE allows the surrogate variables to be correlated with the observed covariates, while setting this to FALSE assumes that the surrogate variables are orthogonal to the observed covariates. This option only matters if design_obs is not NULL. Defaults to FALSE.
design_obs	A numeric matrix of observed covariates that are NOT to be a part of the signal generating process. Only used in estimating the surrogate variables (if target_cor is not NULL). The intercept should <i>not</i> be included (it will sometimes produce an error if it is included).
relative	A logical. Should we apply relative thinning (TRUE) or absolute thinning (FALSE). Only experts should change the default.

change_colnames	3
	A logical. Should we change the column-names of the design matrices (TRUE) or not (FALSE)? Each new column name begins with either "O" (observed), "P" (permuted), or "F" (fixed), followed by a number. The letters correspond to whether the variables come from design_obs, design_perm, or design_fixed. Setting this to TRUE also changes the column-names of the corresponding coefficient matrices. Defaults to TRUE.
permute_method	Should we use the Gale-Shapley algorithm for stable marriages ("marriage") (Gale and Shapley, 1962) as implemented in the matchingR package, or the Hungarian algorithm (Papadimitriou and Steiglitz, 1982) ("hungarian") as implemented in the clue package (Hornik, 2005)? The Hungarian method almost always works better, so is the default.
type	Should we apply binomial thinning (type = "thin") or just naive multiplication of the counts (type = "mult"). You should always have this set to "thin".

Value

A list-like S3 object of class ThinData. Components include some or all of the following:

- mat The modified matrix of counts.
- designmat The design matrix of variables used to simulate signal. This is made by column-binding design_fixed and the permuted version of design_perm.
- coefmat A matrix of coefficients corresponding to designmat.
- design_obs Additional variables that should be included in your design matrix in downstream fittings. This is made by column-binding the vector of 1's with design_obs.
- sv A matrix of estimated surrogate variables. In simulation studies you would probably leave this out and estimate your own surrogate variables.
- cormat A matrix of target correlations between the surrogate variables and the permuted variables in the design matrix. This might be different from the target_cor you input because we pass it through fix_cor to ensure positive semi-definiteness of the resulting covariance matrix.

Mathematical Formulation

Let

- N Be the number of samples.
- G Be the number of genes.
- Y Be an G by N matrix of real RNA-seq counts. This is mat.
- X_1 Be an N by P_1 user-provided design matrix. This is design_fixed.
- X_2 Be an N by P_2 user-provided design matrix. This is design_perm.
- X_3 Be an N by P_3 matrix of known covariates. This is design_obs.
- Z Be an N by K matrix of unobserved surrogate variables. This is estimated when target_cor is not NULL.

M Be a G by N of additional (unknown) unwanted variation.

We assume that Y is Poisson distributed given X_3 and Z such that

$$\log_2(EY) = \mu 1'_N + B_3 X'_3 + AZ' + M.$$

thin_diff() will take as input X_1, X_2, B_1, B_2 , and will output a \tilde{Y} and W such that \tilde{Y} is Poisson distributed given X_1, X_2, X_3, W, Z , and M such that

$$\log_2(EY) \approx \tilde{\mu} 1'_N + B_1 X'_1 + B_2 X'_2 W' + B_3 X'_3 + AZ' + M,$$

where W is an N by N permutation matrix. W is randomly drawn so that WX_2 and Z are correlated approximately according to the target correlation matrix.

The Poisson assumption may be generalized to a mixture of negative binomials.

Unestimable Components

It is possible to include an intercept term or a column from design_obs into either design_fixed or design_perm. This will not produce an error and the specified thinning will be applied. However, If any column of design_fixed or design_perm is a vector of ones or contains a column from design_obs, then the corresponding columns in coef_fixed or coef_perm cannot be estimated by *any* method. This is represented in the output by having duplicate columns in design_obs.

Including duplicate columns in design_fixed and design_perm is also allowed but, again, will produce unestimable coefficients.

Including an intercept term in design_obs will produce an error if you are specifying correlated surrogate variables.

Author(s)

David Gerard

References

- Gale, David, and Lloyd S. Shapley. "College admissions and the stability of marriage." *The American Mathematical Monthly* 69, no. 1 (1962): 9-15. doi:10.1080/00029890.1962.11989827.
- Gerard, D (2020). "Data-based RNA-seq simulations by binomial thinning." BMC Bioinformatics. 21(1), 206. doi:10.1186/s1285902034509.
- Hornik K (2005). "A CLUE for CLUster Ensembles." *Journal of Statistical Software*, 14(12). doi:10.18637/jss.v014.i12.
- Leek, Jeffrey T., and John D. Storey. "A general framework for multiple testing dependence." *Proceedings of the National Academy of Sciences* 105, no. 48 (2008): 18718-18723. doi:10.1073/pnas.0808709105.
- C. Papadimitriou and K. Steiglitz (1982), Combinatorial Optimization: Algorithms and Complexity. Englewood Cliffs: Prentice Hall.

See Also

- select_counts For subsampling the rows and columns of your real RNA-seq count matrix prior to applying binomial thinning.
- thin_2group For the specific application of thin_diff to the two-group model.
- thin_lib For the specific application of thin_diff to library size thinning.
- thin_gene For the specific application of thin_diff to total gene expression thinning.
- thin_all For the specific application of thin_diff to thinning all counts uniformly.
- thin_base For the underlying thinning function used in thin_diff.
- sva For the implementation of surrogate variable analysis.
- ThinDataToSummarizedExperiment For converting a ThinData object to a SummarizedExperiment object.
- ThinDataToDESeqDataSet For converting a ThinData object to a DESeqDataSet object.

```
## Generate simulated data with surrogate variables
## In practice, you would obtain mat from a real dataset, not simulate it.
set.seed(1)
n <- 10
p <- 1000
Z <- matrix(abs(rnorm(n, sd = 4)))</pre>
alpha <- matrix(abs(rnorm(p, sd = 1)))</pre>
mat <- round(2^(alpha %*% t(Z) + abs(matrix(rnorm(n * p, sd = 5),</pre>
                                              nrow = p,
                                              ncol = n))))
## Choose simulation parameters
design_perm <- cbind(rep(c(0, 1), length.out = n), runif(n))</pre>
coef_perm <- matrix(rnorm(p * ncol(design_perm), sd = 6), nrow = p)</pre>
## Specify one surrogate variable (number of columns in taget_cor),
## highly correlated with first observed covariate and uncorrelated
## with second observed covariate
target_cor <- matrix(c(0.9, 0))</pre>
## Thin
thout <- thin_diff(mat = mat,</pre>
                   design_perm = design_perm,
                   coef_perm = coef_perm,
                    target_cor = target_cor)
## target_cor approximates correlation between estimated surrogate variable
## and matching variable.
cor(thout$matching_var, thout$sv)
## Estimated surrogate variable is associated with true surrogate variable
## (because the signal is strong in this case)
plot(Z, thout$sv, xlab = "True SV", ylab = "Estimated SV")
```

thin_gene

```
## So target_cor approximates correlation between surrogate variable and
## matching variables
cor(thout$matching_var, Z)
## Correlation between permuted covariates and surrogate variables are less
## close to target_cor
cor(thout$designmat, Z)
## Estimated signal is correlated to true single. First variable is slightly
## biased because the surrogate variable is not included.
Ynew <- log2(t(thout$mat) + 0.5)
X <- thout$designmat
coef_est <- t(coef(lm(Ynew ~ X))[2:3, ])</pre>
plot(thout$coefmat[, 1], coef_est[, 1],
     main = "First Variable",
     xlab = "Coefficient",
     ylab = "Estimated Coefficient")
abline(0, 1, col = 2, lwd = 2)
plot(thout$coefmat[, 2], coef_est[, 2],
     main = "Second Variable",
     xlab = "Coefficient",
     ylab = "Estimated Coefficient")
abline(0, 1, col = 2, lwd = 2)
## But estimated coefficient of the first variable is slightly closer when
## the surrogate variable is included.
Ynew <- log2(t(thout$mat) + 0.5)
X <- cbind(thout$designmat, thout$sv)</pre>
coef_est <- t(coef(lm(Ynew ~ X))[2:3, ])</pre>
plot(thout$coefmat[, 1], coef_est[, 1],
     main = "First Variable",
     xlab = "Coefficient",
    ylab = "Estimated Coefficient")
abline(0, 1, col = 2, lwd = 2)
plot(thout$coefmat[, 2], coef_est[, 2],
     main = "Second Variable",
     xlab = "Coefficient",
     ylab = "Estimated Coefficient")
abline(0, 1, col = 2, lwd = 2)
```

thin_gene

Description

Given a matrix of real RNA-seq counts, this function will apply a separate, user-provided thinning factor to each gene. This uniformly lowers the counts for all samples in a gene. The thinning factor should be provided on the log2-scale. This is a specific application of the binomial thinning approach in thin_diff. The method is described in detail in Gerard (2020).

Usage

thin_gene(mat,	thinlog2,	relative	= FALSE,	type	= c(["thin",	, "mult")))
----------------	-----------	----------	----------	------	------	----------	-----------	----

Arguments

mat	A numeric matrix of RNA-seq counts. The rows index the genes and the columns index the samples.
thinlog2	A vector of numerics. Element i is the amount to thin (on the log2 scale) for gene i. For example, a value of 0 means that we do not thin, a value of 1 means that we thin by a factor of 2, a value of 2 means we thin by a factor of 4, etc.
relative	A logical. Should we apply relative thinning (TRUE) or absolute thinning (FALSE). Only experts should change the default.
type	Should we apply binomial thinning (type = "thin") or just naive multiplication of the counts (type = "mult"). You should always have this set to "thin".

Value

A list-like S3 object of class ThinData. Components include some or all of the following:

mat The modified matrix of counts.

- designmat The design matrix of variables used to simulate signal. This is made by column-binding design_fixed and the permuted version of design_perm.
- coefmat A matrix of coefficients corresponding to designmat.
- design_obs Additional variables that should be included in your design matrix in downstream fittings. This is made by column-binding the vector of 1's with design_obs.
- sv A matrix of estimated surrogate variables. In simulation studies you would probably leave this out and estimate your own surrogate variables.
- cormat A matrix of target correlations between the surrogate variables and the permuted variables in the design matrix. This might be different from the target_cor you input because we pass it through fix_cor to ensure positive semi-definiteness of the resulting covariance matrix.

Author(s)

David Gerard

References

Gerard, D (2020). "Data-based RNA-seq simulations by binomial thinning." BMC Bioinformatics. 21(1), 206. doi:10.1186/s1285902034509.

thin_lib

See Also

select_counts For subsampling the rows and columns of your real RNA-seq count matrix prior to applying binomial thinning.

thin_diff For the more general thinning approach.

thin_lib For thinning sample-wise instead of gene-wise.

thin_all For thinning all counts uniformly.

ThinDataToSummarizedExperiment For converting a ThinData object to a SummarizedExperiment object.

ThinDataToDESeqDataSet For converting a ThinData object to a DESeqDataSet object.

Examples

```
## Generate count data and thinning factors
## In practice, you would obtain mat from a real dataset, not simulate it.
set.seed(1)
n <- 10
p <- 1000
lambda <- 1000
mat <- matrix(lambda, ncol = n, nrow = p)</pre>
thinlog2 <- rexp(n = p, rate = 1)
## Thin total gene expressions
thout <- thin_gene(mat = mat, thinlog2 = thinlog2)</pre>
## Compare empirical thinning proportions to specified thinning proportions
empirical_propvec <- rowMeans(thout$mat) / lambda</pre>
specified_propvec <- 2 ^ (-thinlog2)</pre>
plot(empirical_propvec, specified_propvec,
     xlab = "Empirical Thinning Proportion",
     ylab = "Specified Thinning Proportion")
abline(0, 1, col = 2, lwd = 2)
```

thin_lib

Binomial thinning for altering library size.

Description

Given a matrix of real RNA-seq counts, this function will apply a separate, user-provided thinning factor to each sample. This uniformly lowers the counts for all genes in a sample. The thinning factor should be provided on the log2-scale. This is a specific application of the binomial thinning approach in thin_diff. The method is described in detail in Gerard (2020).

Usage

```
thin_lib(mat, thinlog2, relative = FALSE, type = c("thin", "mult"))
```

Arguments

mat	A numeric matrix of RNA-seq counts. The rows index the genes and the columns index the samples.
thinlog2	A vector of numerics. Element i is the amount to thin (on the log2-scale) for sample i. For example, a value of 0 means that we do not thin, a value of 1 means that we thin by a factor of 2, a value of 2 means we thin by a factor of 4, etc.
relative	A logical. Should we apply relative thinning (TRUE) or absolute thinning (FALSE). Only experts should change the default.
type	Should we apply binomial thinning (type = "thin") or just naive multiplication of the counts (type = "mult"). You should always have this set to "thin".

Value

A list-like S3 object of class ThinData. Components include some or all of the following:

mat The modified matrix of counts.

- designmat The design matrix of variables used to simulate signal. This is made by column-binding design_fixed and the permuted version of design_perm.
- coefmat A matrix of coefficients corresponding to designmat.
- design_obs Additional variables that should be included in your design matrix in downstream fittings. This is made by column-binding the vector of 1's with design_obs.
- sv A matrix of estimated surrogate variables. In simulation studies you would probably leave this out and estimate your own surrogate variables.
- cormat A matrix of target correlations between the surrogate variables and the permuted variables in the design matrix. This might be different from the target_cor you input because we pass it through fix_cor to ensure positive semi-definiteness of the resulting covariance matrix.

Author(s)

David Gerard

References

Gerard, D (2020). "Data-based RNA-seq simulations by binomial thinning." BMC Bioinformatics. 21(1), 206. doi:10.1186/s1285902034509.

See Also

- select_counts For subsampling the rows and columns of your real RNA-seq count matrix prior to applying binomial thinning.
- thin_diff For the more general thinning approach.
- thin_gene For thinning gene-wise instead of sample-wise.
- thin_all For thinning all counts uniformly.

uncorassign

- ThinDataToSummarizedExperiment For converting a ThinData object to a SummarizedExperiment object.
- ThinDataToDESeqDataSet For converting a ThinData object to a DESeqDataSet object.

Examples

```
## Generate count data and thinning factors
## In practice, you would obtain mat from a real dataset, not simulate it.
set.seed(1)
n <- 10
p <- 1000
lambda <- 1000
mat <- matrix(lambda, ncol = n, nrow = p)
thinlog2 <- rexp(n = n, rate = 1)
## Thin library sizes
thout <- thin_lib(mat = mat, thinlog2 = thinlog2)
## Compare empirical thinning proportions to specified thinning proportions
empirical_propvec <- colMeans(thout$mat) / lambda
specified_propvec</pre>
```

uncorassign Group assignment independent of anything.

Description

Group assignment independent of anything.

Usage

uncorassign(n, return = c("group", "full"))

Arguments

n	The sample size.
return	Should we just return a list with just the vector of assignment ("group") or a list
	with the vector of assignments and the vector of latent variables ("full")?

Value

A list with some or all of the following elements.

- x The group assignment. 1L for one group and 0L for the other group.
- w The latent assignment vector (only returned if return = "full"). Negative corresponds to one group and positive corresponds to the other group.

uncorassign

Author(s)

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