Package 'Dino'

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Type Package

Title Normalization of Single-Cell mRNA Sequencing Data

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Description Dino normalizes single-cell, mRNA sequencing data to correct for technical variation, particularly sequencing depth, prior to downstream analysis. The approach produces a matrix of corrected expression for which the dependency between sequencing depth and the full distribution of normalized expression; many existing methods aim to remove only the dependency between sequencing depth and the mean of the normalized expression. This is particuarly useful in the context of highly sparse datasets such as those produced by 10X genomics and other uninque molecular identifier (UMI) based microfluidics protocols for which the depth-dependent proportion of zeros in the raw expression data can otherwise present a challenge.

Depends R (>= 4.0.0)

License GPL-3

Encoding UTF-8

LazyData false

RoxygenNote 7.1.1

Suggests testthat (>= 2.1.0), knitr, rmarkdown, BiocStyle, devtools, ggplot2, gridExtra, ggpubr, grid, magick, hexbin

VignetteBuilder knitr

Imports BiocParallel, BiocSingular, SummarizedExperiment, SingleCellExperiment, S4Vectors, Matrix, Seurat, matrixStats, parallel, scran, grDevices, stats, methods

URL https://github.com/JBrownBiostat/Dino

BugReports https://github.com/JBrownBiostat/Dino/issues

git_url https://git.bioconductor.org/packages/Dino

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Dino

Normalize scRNAseq data

Description

Dino removes cell-to-cell variation in observed counts due to the effects of sequencing depth from single-cell mRNA sequencing experiments. Dino was particularly designed with UMI based protocols in mind, but is applicable to non-UMI based chemistries in the library preparation stage of sequencing.

Usage

```
Dino(counts, nCores = 2, prec = 3, minNZ = 10,
    nSubGene = 1e4, nSubCell = 1e4, depth = NULL, slope = NULL,
    minSlope = 1/2, maxSlope = 2, clusterSlope = TRUE,
    returnMeta = FALSE, doRQS = FALSE,
    emPar = list(maxIter = 100, tol = 0.1, conPar = 15, maxK = 100), ...)
```

Arguments

counts	A numeric matrix object of expression counts - usually in dgCMatrix format for
	memory efficiency. Column names denote cells (samples or droplets) and row
	names denote genes.
nCores	A non-negative integer scalar denoting the number of cores which should be
	used. Setting nCores to 0 uses all cores as determined by running parallel::detectCores()

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A positive integer denoting the number of decimals to which to round depth (if estimated internally via depth = NULL) and normalized counts for computational efficiency.

A positive integer denoting the minimum number of non-zero counts for a gene

to be normalized by the Dino algorithm. It is recommended to pre-filter the *counts* matrix such that all genes meet this threshold. Otherwise, genes with fewer than *minNZ* non-zeros will be scaled by depth for normalization.

nSubGene A positive integer denoting the number of genes to subset for calculation of

slope.

minNZ

nSubCell A positive integer denoting the number of samples to subset for calculation of

slope and the EM algorithm.

depth A numeric vector of length equal to the columns of counts. depth denotes a

median-centered, log-scale measure of cell-wise sequencing depth. Dino defaults to defining depth as the (within-cell) sum of counts across genes, followed

by a log and median-centering transformation.

slope A numeric scalar denoting the count-depth relationship on the log-log scale.

Typical values are close to 1 (implying a unit increase in depth corresponds to a unit increase in expected counts on the log-log scale), but may be higher, particularly in the case of non-UMI protocols. Dino defaults to estimating *slope*

internally.

minSlope A numeric scalar denoting the minimum slope. Fitted slopes below this value

will return a warning and be set to 1

maxSlope A numeric scalar denoting the maximum slope. Fitted slopes above this value

will return a warning and be set to 1

clusterSlope A logical indicating whether cells should be pre-clustered prior to calculation of

slope. Under the default where cells are pre-clustered, cluster is used as a factor

in the regression.

returnMeta A logical indicating whether metadata (sequencing depth and slope) should be

returned.

doRQS A logical indicating how normalization resampling is to be done. By default

(F), normalization is done by resampling from the full posterior distribution. Alternately, restricted quantile sampling (RQS) can be performed to enforce stronger preservation of expression ranks in normalized data. Currently RQS is

considered experimental.

emPar A list of parameters to send to the EM algorithm. *maxIter* denotes the maximum

number of model updates. tol denotes the cutoff threshold for reductions in the log likelihood function. conPar denotes the concentration parameter for the resampling. conPar = 1 implies full resampling from the fitted distribution. As conPar increases, the normalized expression converges to the scale-factor normalized values. maxK denotes the maximum number of mixture components

in the mixture model.

... Additional parameters to pass to Scran::quickCluster.

Value

Dino by default returns a matrix of normalized expression with identical dimensions as *counts*. If *returnMeta* = *TRUE*, then Dino returns a list of normalized expression, sequencing depth, and slope.

Dino_SCE

Author(s)

Jared Brown

References

Brown, J., Ni, Z., Mohanty, C., Bacher, R. and Kendziorski, C. (2020) "Normalization by distributional resampling of high throughput single-cell RNA-sequencing data." bioRxiv. https://doi.org/10.1101/2020.10.28.359901

Examples

```
# raw data
data("pbmcSmall")
str(pbmcSmall)

# run Dino on raw expression matrix
pbmcSmall_Norm <- Dino(pbmcSmall)
str(pbmcSmall_Norm)</pre>
```

Dino_SCE

Run Dino normalization on a SingleCellExperiment dataset

Description

Dino_SCE is a wrapper simplifying the application of the *Dino* method to data formatted as a *Sin-gleCellExperiment*

Usage

```
Dino_SCE(SCE, ...)
```

Arguments

SCE

A SingleCellExperiment object with unnormalized count data (eg. raw UMIs)

in the assays slot under the name counts.

... Further arguments to pass to *Dino*

Value

Dino_SCE returns a *SingleCellExperiment* object using Dino normalized expression in the *assays* slot under the *normcounts* name for downstream analysis.

If returnMeta = T is passed to Dino, then depth and slope results are stored in the metadata slot under the names depth and slope respectively.

Author(s)

Jared Brown

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References

Brown, J., Ni, Z., Mohanty, C., Bacher, R. and Kendziorski, C. (2020). "Normalization by distributional resampling of high throughput single-cell RNA-sequencing data." bioRxiv. https://doi.org/10.1101/2020.10.28.359901

Amezquita, R.A., Lun, A.T.L., Becht, E., Carey, V.J., Carpp, L.N., Geistlinger, L., Marini, F., Rue-Albrecht, K., Risso, D., Soneson, C., et al. (2020). "Orchestrating single-cell analysis with Bioconductor." Nat. Methods, 17, 137–145. https://doi.org/10.1038/s41592-019-0654-x

Examples

```
# raw data
data("pbmcSmall")
str(pbmcSmall)

# format as SingleCellExperiment
library(SingleCellExperiment)
pbmc_SCE <- SingleCellExperiment(assays = list("counts" = pbmcSmall))

# Run Dino
pbmc_SCE <- Dino_SCE(pbmc_SCE)
str(pbmc_SCE)
str(normcounts(pbmc_SCE))</pre>
```

multimodalDat

Plot data from simulated expression

Description

This data is used in the vignette to demonstrate the flexibility of the Dino model to smoothly estimate arbitrary latent multimodal expression distributions. These data are intended for internal use only.

Usage

```
data("multimodalDat")
```

Format

Object of class "gtable".

Examples

```
data("multimodalDat")
```

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pbmcSmall	Subset of 500 peripheral blood mononuclear cells (PBMCs) from a healthy donor

Description

This dataset derives from the "3k PBMCs from a Healthy Donor" public dataset from 10X Genomics.

Usage

```
data("pbmcSmall")
```

Format

An object of class "dgCMatrix".

Source

3k PBMCs from a Healthy Donor

Examples

```
data("pbmcSmall")
str(pbmcSmall)
```

SeuratFromDino

Create Seurat object from Dino normalized data

Description

SeuratFromDino is a wrapper simplifying the export of Dino normalized counts to a *Seurat* object for secondary analysis.

Usage

```
SeuratFromDino(counts, doNorm = TRUE, doLog = TRUE, ...)
```

Arguments

counts	A numeric matrix of count data, either raw (eg. UMIs) or normalized expression.
doNorm	A logical indicating whether to normalize the input <i>counts</i> data before exporting results to a <i>Seurat</i> object. By default, it is assumed that the contents of <i>counts</i> raw expression which should be normalized.
doLog	A logical indicating whether normalized counts should be log transformed with a psuedocount of 1 prior to export.
	Further arguments to pass to Dino

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Value

SeuratFromDino returns a Seurat object using Dino normalized and log transformed expression (default) for downstream analysis in the Seurat pipeline.

If returnMeta = T is passed to Dino, then depth and slope results are stored in the Misc slot under the names depth and slope respectively.

Author(s)

Jared Brown

References

Brown, J., Ni, Z., Mohanty, C., Bacher, R. and Kendziorski, C. (2020). "Normalization by distributional resampling of high throughput single-cell RNA-sequencing data." bioRxiv. https://doi.org/10.1101/2020.10.28.359901 Satija, R., Farrell, J.A., Gennert, D., Schier, A.F. and Regev, A. (2015). "Spatial reconstruction of single-cell gene expression data." Nat. Biotechnol., 33, 495–502. https://doi.org/10.1038/nbt.3192

Examples

```
# raw data
data("pbmcSmall")
str(pbmcSmall)

# run Dino on raw expression matrix, output Seurat object
pbmcSmall_Seurat <- SeuratFromDino(pbmcSmall)
str(pbmcSmall_Seurat)</pre>
```

unimodalDat

Plot data from simulated expression

Description

This data is used in the vignette to demonstrate the flexibility of the Dino model to smoothly estimate arbitrary latent uninmodal expression distributions. These data are intended for internal use only.

Usage

```
data("unimodalDat")
```

Format

Object of class "gtable".

Examples

```
data("unimodalDat")
```

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