Package 'TooManyCellsR'

October 12, 2022

| tooManyCells | Title An R wrapper for ToolManyCells | | | |
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| Maintainer Gregory W. Schwartz <gsch@pennmedicine.upenn.edu> Description An R wrapper for using 'TooManyCells', a command line program for clustering, visualizing, and quantifying cell clade relationships. See https://gregoryschwartz.github.io/too-many-cells/ for more details. Imports Matrix, imager, ggplot2, cowplot, jsonlite SystemRequirements 'TooManyCells'</gsch@pennmedicine.upenn.edu> | Version 0.1.1.0 | | | |
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2 importResults

importResults

Import some 'too-many-cells make-tree' outputs into a data frame.

Description

This function will import some of the files resulting from a 'too-many-cells make-tree' run into R as data frames. Does not include cluster list. Look at the main tooManyCells function for the cluster list.

Usage

```
importResults(dir = "out")
```

Arguments

dir

The output directory of a 'too-many-cells' run.

Value

A list of each output. Reads the following files, see https://gregoryschwartz.github.io/too-many-cells/ for more details: "dendrogram.svg", "clumpiness.pdf", "projection.pdf", "label projection.pdf", "clumpiness.csv", "cluster info.csv", "node info.csv", and "cluster diversity.csv".

```
input <- system.file("extdata", "mat.csv", package="TooManyCellsR")</pre>
inputLabels <- system.file("extdata", "labels.csv", package="TooManyCellsR")</pre>
df = read.csv(input, row.names = 1, header = TRUE)
mat = Matrix::Matrix(as.matrix(df), sparse = TRUE)
labelsDf = read.csv(inputLabels, header = TRUE)
# Here we draw this small toy example with no filter or normalization, and
# decrease the size of the branches and increase the size of the leaf nodes.
# With non-toy real world single cell data, these options should not be
# necessary.
## Not run:
tooManyCells( mat, labels = labelsDf
            , args = c( "make-tree"
                      , "--no-filter"
                        "--normalization", "NoneNorm"
                        "--draw-max-node-size", "40"
                        "--draw-max-leaf-node-size", "70"
res = importResults("out")
plot(res$treePlot, axes = FALSE)
## End(Not run)
```

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| tooManyCells Execute too-many-cells. | tooManyCells | Execute 'too-many-cells'. |
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Description

This function will run 'too-many-cells' on a Matrix. Requires 'TooManyCells' to be installed (follow instructions at https://gregoryschwartz.github.io/too-many-cells/).

Usage

```
tooManyCells(mat, args = c("make-tree"), labels = NULL,
  output = "out", prior = NULL, docker = NULL, mounts = c())
```

Arguments

| mat | The input Matrix with gene row names and cell barcode column names. |
|--------|---|
| args | The arguments to give to the command line program. See https://gregoryschwartz.github.io/too-many-cells/ for more information. Defaults to "make-tree". |
| labels | The input labels data frame with item (cell barcodes) and label (whatever labels you want to give them, such as tissue of origin, celltype, etc.) columns. Optional. |
| output | The output folder for the 'too-many-cells' process. Defaults to "out". |
| prior | The location of the tree that was already made (previous 'too-many-cells' output) so quick visual or pruning changes can be made without remaking the tree (can potentially save hours). |
| docker | If using 'too-many-cells' with docker, use this argument as the command to call. For instance, if version 0.2.1.0 was pulled from Docker Hub, set to "gregoryschwartz/too-many-cells:0.2.1.0". |
| mounts | Additional directories to mount if needed for docker. The 'prior' argument will automatically mount if specified. |

Value

A list of each output, including the stdout. Reads the following files, see https://gregoryschwartz.github.io/too-many-cells/ for more details: "dendrogram.svg", "clumpiness.pdf", "projection.pdf", "label_projection.pdf", "clumpiness.csv", "cluster_info.csv", "node_info.csv", and "cluster_diversity.csv".

```
input <- system.file("extdata", "mat.csv", package="TooManyCellsR")
inputLabels <- system.file("extdata", "labels.csv", package="TooManyCellsR")
df = read.csv(input, row.names = 1, header = TRUE)
mat = Matrix::Matrix(as.matrix(df), sparse = TRUE)
labelsDf = read.csv(inputLabels, header = TRUE)
# Here we draw this small toy example with no filter or normalization, and</pre>
```

4 tryFunc

tryFunc

Different error for importing data.

Description

This function will fail gracefully instead of stopping the program with an error.

Usage

```
tryFunc(f, file)
```

Arguments

f The function to use.

file The input file to be read.

Value

The imported data frame or NULL if an error occurred.

```
input <- system.file("extdata", "mat.csv", package="TooManyCellsR")
fail = tryFunc(read.csv, "fail.csv")
fail
success = tryFunc(read.csv, input)
success</pre>
```

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| writeMatrixFiles | Write a Matrix to a folder. |
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Description

This function will write a Matrix from the Matrix library to a temporary directory containing matrix.mtx, genes.tsv, barcodes.tsv, and optionally a labels.csv file.

Usage

```
writeMatrixFiles(mat, labels = NULL)
```

Arguments

mat The input Matrix with gene row names and cell barcode column names.

labels The input labels data frame with item (cell barcodes) and label (whatever la-

bels you want to give them, such as tissue of origin, celltype, etc.) columns.

Optional.

Value

None

```
input <- system.file("extdata", "mat.csv", package="TooManyCellsR")
df = read.csv(input, row.names = 1, header = TRUE)
mat = Matrix::Matrix(as.matrix(df), sparse = TRUE)
writeMatrixFiles(mat)</pre>
```

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