

Package ‘bodenmiller’

August 29, 2016

Type Package

Title Profiling of Peripheral Blood Mononuclear Cells using CyTOF

Version 0.1

Depends R (>= 3.1.0)

Suggests ggplot2, dplyr, reshape2, RColorBrewer, knitr, rmarkdown

Description

This data package contains a subset of the Bodenmiller et al, Nat Biotech 2012 dataset for testing single cell, high dimensional analysis and visualization methods.

License CC0

LazyData TRUE

URL <https://github.com/yannabraham/bodenmiller>

BugReports <https://github.com/yannabraham/bodenmiller/issues>

VignetteBuilder knitr

Author Yann Abraham [aut, cre],
Bernd Bodenmiller [aut]

Maintainer Yann Abraham <yann.abraham@gmail.com>

NeedsCompilation no

Repository CRAN

Date/Publication 2015-12-18 22:45:47

R topics documented:

refAnnots	2
refFuncMat	2
refPhenoMat	3
untreatedAnnots	4
untreatedFuncMat	4
untreatedPhenoMat	5

Index

7

refAnnots	<i>Reference Annotation</i>
-----------	-----------------------------

Description

A `data.frame` containing the source file and cell type for every cell in `refPhenoMat` and `refFuncMat`. See [Bodenmiller et al 2012](#) for details.

Usage

```
refAnnots
```

Format

a `data.frame` with 15792 rows and 2 variables:

- `Source`the name of the source (fcs) file
- `Cell`the cell type for the corresponding row in `refPhenoMat` or `refFuncMat`

Source

<http://reports.cytobank.org/105/v2>

refFuncMat	<i>Reference Matrix (Functional)</i>
------------	--------------------------------------

Description

a `matrix` containing the signal intensity for functional markers measured using CyTOF on different cell types. See [Bodenmiller et al 2012](#) for details.

Usage

```
refFuncMat
```

Format

a `matrix` with 15792 rows and 14 variables:

- `pStat1`
- `pSlp76`
- `pBtk`
- `pPlcg2`
- `pErk`
- `pLat`

- pS6
- pNFkB
- pp38
- pStat5
- pAkt
- pSHP2
- pZap70
- pStat3

Source

<http://reports.cytobank.org/105/v2>

refPhenoMat

Reference Matrix (Phenotypic)

Description

a matrix containing the signal intensity for phenotypic markers measured using CyTOF on different cell types. See [Bodenmiller et al 2012](#) for details.

Usage

refPhenoMat

Format

a matrix with 15792 rows and 9 variables:

- CD20
- IgM
- CD4
- CD33
- HLA-DR
- CD14
- CD7
- CD3
- CD123

Source

<http://reports.cytobank.org/105/v2>

untreatedAnnots*Untreated Samples with 4 Different Treatments (Annotation)*

Description

A `data.frame` containing the source file, cell type and stimulation for every cell in `refPhenoMat` and `refFuncMat`. Stimulations include BCR/FcR-XL, PMA/Ionomycin and vanadate. See [Bodenmiller et al 2012](#) for details.

Usage

```
untreatedAnnots
```

Format

a `data.frame` with 15792 rows and 2 variables:

- `Source`the name of the source (fcs) file
- `Treatment`the treatment for the corresponding row in `refPhenoMat` or `refFuncMat`
- `Cell`the cell type for the corresponding row in `refPhenoMat` or `refFuncMat`

Source

<http://reports.cytobank.org/105/v2>

untreatedFuncMat*Untreated Samples with 4 Different Treatments (Functional)*

Description

a `matrix` containing the signal intensity for functional markers measured using CyTOF on different cell types, after stimulation with BCR/FcR-XL, PMA/Ionomycin and vanadate. See [Bodenmiller et al 2012](#) for details.

Usage

```
untreatedFuncMat
```

Format

a matrix with 15792 rows and 14 variables:

- pStat1
- pSlp76
- pBtk
- pPlcg2
- pErk
- pLat
- pS6
- pNFkB
- pp38
- pStat5
- pAkt
- pSHP2
- pZap70
- pStat3

Source

<http://reports.cytobank.org/105/v2>

untreatedPhenoMat

Untreated Samples with 4 Different Treatments (Phenotypic)

Description

a matrix containing the signal intensity for phenotypic markers measured using CyTOF on different cell types, after stimulation with BCR/FcR-XL, PMA/Ionomycin and vanadate. See [Bodenmiller et al 2012](#) for details.

Usage

untreatedPhenoMat

Format

a matrix with 15792 rows and 9 variables:

- CD20
- IgM
- CD4
- CD33

- HLA-DR
- CD14
- CD7
- CD3
- CD123

Source

<http://reports.cytobank.org/105/v2>

Index

*Topic **datasets**

- refAnnots, [2](#)
- refFuncMat, [2](#)
- refPhenoMat, [3](#)
- untreatedAnnots, [4](#)
- untreatedFuncMat, [4](#)
- untreatedPhenoMat, [5](#)

- refAnnots, [2](#)
- refFuncMat, [2](#), [2](#), [4](#)
- refPhenoMat, [2](#), [3](#), [4](#)
- untreatedAnnots, [4](#)
- untreatedFuncMat, [4](#)
- untreatedPhenoMat, [5](#)