

# Package ‘clustifyrdatahub’

October 17, 2024

**Title** External data sets for clustifyr in ExperimentHub

**Version** 1.14.0

**Description** References made from external single-cell mRNA sequencing data sets, stored as average gene expression matrices. For use with clustifyr <<https://bioconductor.org/packages/clustifyr>> to assign cell type identities.

**License** MIT + file LICENSE

**URL** <https://rnabioco.github.io/clustifyrdatahub/>

**Depends** R (>= 4.0), ExperimentHub

**Imports** utils

**Suggests** clustifyr, Seurat, usethis, rmarkdown, knitr, tidyr,  
BiocStyle

**VignetteBuilder** knitr

**Encoding** UTF-8

**LazyData** true

**RoxygenNote** 7.2.1

**biocViews** SingleCellData, SequencingData, MicroarrayData,  
ExperimentHub, RNASeqData, PackageTypeData, ExpressionData

**BugReports** <https://github.com/rnabioco/clustifyrdatahub/issues>

**Config/Needs/website** pkgdown, rnabioco/rbitemplate

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## Contents

clustifyrdatahub	2
ref_cortex_dev	3
ref_hema_microarray	3
ref_immgem	4
ref_MCA	5
ref_moca_main	5
ref_mouse.rnaseq	6
ref_mouse_atlas	7
ref_pan_indrop	7
ref_pan_smartseq2	8
ref_tabula_muris_drop	9
ref_tabula_muris_facs	9

## Index

11

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clustifyrdatahub *clustifyrdatahub: reference scRNA-seq data for clustifyr*

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## Description

The `clustifyrdatahub` package provides access to the summarized reference count matrix data from multiple datasets used in the `clustifyr` manuscript and tutorial. They are provided for future users of `clustifyr`, for automated cell type classification of scRNA-seq experiments.

## Details

Available datasets:

- + Mouse Cell Atlas - `[ref_MCA()]` + Tabula Muris (10X) - `[ref_tabula_muris_drop()]` + Tabula Muris (SmartSeq2) - `[ref_tabula_muris_facs()]` + Mouse RNA-seq from 28 cell types - `[ref_mouse.rnaseq()]`
- + Mouse Organogenesis Cell Atlas (main cell types) - `[ref_moca_main()]` + Mouse sorted immune cells - `[ref_immgem()]` + Human hematopoietic cell microarray - `[ref_hema_microarray()]` + Human cortex development scRNA-seq - `[ref_cortex_dev()]` + Human pancreatic cell scRNA-seq (inDrop) - `[ref_pan_indrop()]` + Human pancreatic cell scRNA-seq (SmartSeq2) - `[ref_pan_smartseq2()]`

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ref\_cortex\_dev

*Matrix of average gene detection on cortex development cell types*

---

### Description

47 distinct annotated clusters of human neuronal subtypes

### Usage

```
ref_cortex_dev
```

### Format

An object of class function of length 1.

### Source

```
<https://cells.ucsc.edu/?ds=cortex-dev#>
```

### See Also

Other ref: [ref\\_MCA\(\)](#), [ref\\_hema\\_microarray\(\)](#), [ref\\_immgen\(\)](#), [ref\\_moca\\_main\(\)](#), [ref\\_mouse.rnaseq\(\)](#), [ref\\_mouse\\_atlas\(\)](#), [ref\\_pan\\_indrop\(\)](#), [ref\\_pan\\_smartseq2\(\)](#), [ref\\_tabula\\_muris\\_drop\(\)](#), [ref\\_tabula\\_muris\\_facs\(\)](#)

### Examples

```
ref_cortex_dev(metadata = TRUE)
```

---

ref\_hema\_microarray

*Matrix of average gene detection by microarray on sorted immune cell populations*

---

### Description

38 distinct purified populations of human hematopoietic cells

### Usage

```
ref_hema_microarray
```

### Format

An object of class function of length 1.

**Source**

[<https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE24759>]

**See Also**

Other ref: [ref\\_MCA\(\)](#), [ref\\_cortex\\_dev\(\)](#), [ref\\_immgene\(\)](#), [ref\\_moca\\_main\(\)](#), [ref\\_mouse.rnaseq\(\)](#), [ref\\_mouse\\_atlas\(\)](#), [ref\\_pan\\_indrop\(\)](#), [ref\\_pan\\_smartseq2\(\)](#), [ref\\_tabula\\_muris\\_drop\(\)](#), [ref\\_tabula\\_muris\\_facs\(\)](#)

**Examples**

```
ref_hema_microarray(metadata = TRUE)
```

---

ref\_immgene

*Matrix of average gene expression from immgen data*

---

**Description**

253 mouse immune cell types

**Usage**

```
ref_immgene
```

**Format**

An object of class `function` of length 1.

**Source**

```
["SingleR"]
```

**See Also**

Other ref: [ref\\_MCA\(\)](#), [ref\\_cortex\\_dev\(\)](#), [ref\\_hema\\_microarray\(\)](#), [ref\\_moca\\_main\(\)](#), [ref\\_mouse.rnaseq\(\)](#), [ref\\_mouse\\_atlas\(\)](#), [ref\\_pan\\_indrop\(\)](#), [ref\\_pan\\_smartseq2\(\)](#), [ref\\_tabula\\_muris\\_drop\(\)](#), [ref\\_tabula\\_muris\\_facs\(\)](#)

**Examples**

```
ref_immgene(metadata = TRUE)
```

---

ref\_MCA

*Matrix of average gene expression for single-cell RNA-seq.*

---

### Description

from mouse cell atlas project

### Usage

`ref_MCA`

### Format

An object of class function of length 1.

### Source

[<https://figshare.com/s/865e694ad06d5857db4b>](https://figshare.com/s/865e694ad06d5857db4b)

### See Also

Other ref: [ref\\_cortex\\_dev\(\)](#), [ref\\_hema\\_microarray\(\)](#), [ref\\_immgene\(\)](#), [ref\\_moca\\_main\(\)](#), [ref\\_mouse.rnaseq\(\)](#), [ref\\_mouse\\_atlas\(\)](#), [ref\\_pan\\_indrop\(\)](#), [ref\\_pan\\_smartseq2\(\)](#), [ref\\_tabula\\_muris\\_drop\(\)](#), [ref\\_tabula\\_muris\\_facs\(\)](#)

### Examples

`ref_MCA(metadata = TRUE)`

---

ref\_moca\_main

*Matrix of average gene expression per cluster from single-cell sci-RNA-seq3 from Mouse Organogenesis Cell Atlas.*

---

### Description

100,000 cells

### Usage

`ref_moca_main`

### Format

An object of class function of length 1.

**Source**

<<http://atlas.gs.washington.edu/mouse-rna/>>

**See Also**

Other ref: [ref\\_MCA\(\)](#), [ref\\_cortex\\_dev\(\)](#), [ref\\_hema\\_microarray\(\)](#), [ref\\_immgene\(\)](#), [ref\\_mouse.rnaseq\(\)](#),  
[ref\\_mouse\\_atlas\(\)](#), [ref\\_pan\\_indrop\(\)](#), [ref\\_pan\\_smartseq2\(\)](#), [ref\\_tabula\\_muris\\_drop\(\)](#),  
[ref\\_tabula\\_muris\\_facs\(\)](#)

**Examples**

```
ref_moca_main(metadata = TRUE)
```

---

`ref_mouse.rnaseq`      *Matrix of average gene expression from general mouse cell type data*

---

**Description**

28 general cell types

**Usage**

```
ref_mouse.rnaseq
```

**Format**

An object of class `function` of length 1.

**Source**

```
["SingleR"]
```

**See Also**

Other ref: [ref\\_MCA\(\)](#), [ref\\_cortex\\_dev\(\)](#), [ref\\_hema\\_microarray\(\)](#), [ref\\_immgene\(\)](#), [ref\\_moca\\_main\(\)](#),  
[ref\\_mouse\\_atlas\(\)](#), [ref\\_pan\\_indrop\(\)](#), [ref\\_pan\\_smartseq2\(\)](#), [ref\\_tabula\\_muris\\_drop\(\)](#),  
[ref\\_tabula\\_muris\\_facs\(\)](#)

**Examples**

```
ref_mouse.rnaseq(metadata = TRUE)
```

---

ref_mouse_atlas	<i>Atlas matrix of average gene expression aggregated from scRNA-seq data of mouse organisms</i>
-----------------	--

---

## Description

Data aggregated from NCBI Gene Expression Omnibus

## Usage

```
ref_mouse_atlas
```

## Format

An object of class function of length 1.

## Source

<<https://github.com/rnabioco/scRNA-seq-Cell-Ref-Matrix/blob/master/atlas/musMusculus/MouseAtlas.rds>>

## See Also

Other ref: [ref\\_MCA\(\)](#), [ref\\_cortex\\_dev\(\)](#), [ref\\_hema\\_microarray\(\)](#), [ref\\_immgene\(\)](#), [ref\\_moca\\_main\(\)](#), [ref\\_mouse.rnaseq\(\)](#), [ref\\_pan\\_indrop\(\)](#), [ref\\_pan\\_smartseq2\(\)](#), [ref\\_tabula\\_muris\\_drop\(\)](#), [ref\\_tabula\\_muris\\_facs\(\)](#)

## Examples

```
ref_mouse_atlas(metadata = TRUE)
```

---

ref_pan_indrop	<i>Matrix of average gene expression per cluster from single-cell RNA-seq pancreas data using indrop method.</i>
----------------	--

---

## Description

A sample of 8569 pancreas cells

## Usage

```
ref_pan_indrop
```

## Format

An object of class function of length 1.

**Source**

<<https://hemberg-lab.github.io/scRNA.seq.datasets/human/pancreas/>>

**See Also**

Other ref: [ref\\_MCA\(\)](#), [ref\\_cortex\\_dev\(\)](#), [ref\\_hema\\_microarray\(\)](#), [ref\\_immgene\(\)](#), [ref\\_moca\\_main\(\)](#),  
[ref\\_mouse.rnaseq\(\)](#), [ref\\_mouse\\_atlas\(\)](#), [ref\\_pan\\_smartseq2\(\)](#), [ref\\_tabula\\_muris\\_drop\(\)](#),  
[ref\\_tabula\\_muris\\_facs\(\)](#)

**Examples**

```
ref_pan_indrop(metadata = TRUE)
```

---

**ref\_pan\_smartseq2**      *Matrix of average gene expression per cluster from single-cell RNA-seq pancreas data using smartseq2 method.*

---

**Description**

A sample of 2209 pancreas cells

**Usage**

```
ref_pan_smartseq2
```

**Format**

An object of class `function` of length 1.

**Source**

<<https://hemberg-lab.github.io/scRNA.seq.datasets/human/pancreas/>>

**See Also**

Other ref: [ref\\_MCA\(\)](#), [ref\\_cortex\\_dev\(\)](#), [ref\\_hema\\_microarray\(\)](#), [ref\\_immgene\(\)](#), [ref\\_moca\\_main\(\)](#),  
[ref\\_mouse.rnaseq\(\)](#), [ref\\_mouse\\_atlas\(\)](#), [ref\\_pan\\_indrop\(\)](#), [ref\\_tabula\\_muris\\_drop\(\)](#), [ref\\_tabula\\_muris\\_facs\(\)](#)

**Examples**

```
ref_pan_smartseq2(metadata = TRUE)
```

---

`ref_tabula_muris_drop` *Matrix of average gene expression per cluster from single-cell RNA-seq from Tabula Muris (10x genomics).*

---

### Description

422,803 droplet libraries, 55,656 of which passed a QC cutoff of 500 genes and 1000 UMI

### Usage

```
ref_tabula_muris_drop
```

### Format

An object of class function of length 1.

### Source

<<https://tabula-muris.ds.czbiohub.org/>>

### See Also

Other ref: [ref\\_MCA\(\)](#), [ref\\_cortex\\_dev\(\)](#), [ref\\_hema\\_microarray\(\)](#), [ref\\_immmgen\(\)](#), [ref\\_moca\\_main\(\)](#), [ref\\_mouse.rnaseq\(\)](#), [ref\\_mouse\\_atlas\(\)](#), [ref\\_pan\\_indrop\(\)](#), [ref\\_pan\\_smartseq2\(\)](#), [ref\\_tabula\\_muris\\_facs\(\)](#)

### Examples

```
ref_tabula_muris_drop(metadata = TRUE)
```

---

`ref_tabula_muris_facs` *Matrix of average gene expression per cluster from single-cell RNA-seq from Tabula Muris (SMART-Seq2).*

---

### Description

53,760 cells from 20 tissues from 8 mice

### Usage

```
ref_tabula_muris_facs
```

### Format

An object of class function of length 1.

**Source**

<<https://tabula-muris.ds.czbiohub.org/>>

**See Also**

Other ref: [ref\\_MCA\(\)](#), [ref\\_cortex\\_dev\(\)](#), [ref\\_hema\\_microarray\(\)](#), [ref\\_immgene\(\)](#), [ref\\_moca\\_main\(\)](#),  
[ref\\_mouse\\_rnaseq\(\)](#), [ref\\_mouse\\_atlas\(\)](#), [ref\\_pan\\_indrop\(\)](#), [ref\\_pan\\_smartseq2\(\)](#), [ref\\_tabula\\_muris\\_drop\(\)](#)

**Examples**

```
ref_tabula_muris_facs(metadata = TRUE)
```

# Index

- \* **datasets**
  - ref\_cortex\_dev, [3](#)
  - ref\_hema\_microarray, [3](#)
  - ref\_immgem, [4](#)
  - ref\_MCA, [5](#)
  - ref\_moca\_main, [5](#)
  - ref\_mouse.rnaseq, [6](#)
  - ref\_mouse\_atlas, [7](#)
  - ref\_pan\_indrop, [7](#)
  - ref\_pan\_smartseq2, [8](#)
  - ref\_tabula\_muris\_drop, [9](#)
  - ref\_tabula\_muris\_facs, [9](#)
- \* **internal**
  - clustifyrdatahub, [2](#)
- \* **ref**
  - ref\_cortex\_dev, [3](#)
  - ref\_hema\_microarray, [3](#)
  - ref\_immgem, [4](#)
  - ref\_MCA, [5](#)
  - ref\_moca\_main, [5](#)
  - ref\_mouse.rnaseq, [6](#)
  - ref\_mouse\_atlas, [7](#)
  - ref\_pan\_indrop, [7](#)
  - ref\_pan\_smartseq2, [8](#)
  - ref\_tabula\_muris\_drop, [9](#)
  - ref\_tabula\_muris\_facs, [9](#)
- clustifyrdatahub, [2](#)
  - ref\_cortex\_dev, [3, 4–10](#)
  - ref\_hema\_microarray, [3, 3, 4–10](#)
  - ref\_immgem, [3, 4, 4, 5–10](#)
  - ref\_MCA, [3, 4, 5, 6–10](#)
  - ref\_moca\_main, [3–5, 5, 6–10](#)
  - ref\_mouse.rnaseq, [3–6, 6, 7–10](#)
  - ref\_mouse\_atlas, [3–6, 7, 8–10](#)
  - ref\_pan\_indrop, [3–7, 7, 8–10](#)
  - ref\_pan\_smartseq2, [3–8, 8, 9, 10](#)
  - ref\_tabula\_muris\_drop, [3–8, 9, 10](#)
  - ref\_tabula\_muris\_facs, [3–9, 9](#)