# Package 'LRcellTypeMarkers'

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Type Package

Title Marker gene information for LRcell R Bioconductor package Version 1.12.0 Date 2020-02-25 BugReports https://github.com/marvinquiet/LRcellTypeMarkers/issues GitURL https://github.com/marvinquiet/LRcellTypeMarkers Description This is an external ExperimentData package for LRcell. This data package contains the gene enrichment scores calculated from scRNA-seq dataset which indicates the gene enrichment of each cell type in certain brain region. LRcell package is used to identify specific sub-cell types that drives the changes observed in a bulk RNA-seq differential gene expression experiment. For more details, please visit: https://github.com/marvinquiet/LRcell. License MIT + file LICENSE **Encoding UTF-8** biocViews ExperimentData, ExperimentHub, RNASeqData, SingleCellData, ExpressionData, PackageTypeData **Depends** R (>= 4.1)Imports ExperimentHub Suggests LRcell, BiocStyle, knitr, rmarkdown, roxygen2, testthat VignetteBuilder knitr RoxygenNote 7.1.1 git\_url https://git.bioconductor.org/packages/LRcellTypeMarkers git\_branch RELEASE\_3\_19 git\_last\_commit baceedc git\_last\_commit\_date 2024-04-30 Repository Bioconductor 3.19 Date/Publication 2024-06-11 **Author** Wenjing Ma [cre, aut] (<a href="https://orcid.org/0000-0001-8757-651X">https://orcid.org/0000-0001-8757-651X</a>)

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#### Description

This is an external ExperimentData package for LRcell. This data package contains the gene enrichment scores calculated from scRNA-seq dataset which indicates the gene enrichment of each cell type in certain brain region. LRcell package is used to identify specific sub-cell types that drives the changes observed in a bulk RNA-seq differential gene expression experiment. For more details, please visit: https://github.com/marvinquiet/LRcell.

#### **Details**

See the vignette for examples of extracting and using these data.

browseVignettes("LRcellTypeMarkers")

Details of how these data were created are in the scripts/ directory of the source package.

The current release includes the following datasets:

Mouse Whole Brain Marker Genes Gene enrichment scores calculated from Mouse Adult Whole Brain scRNA-seq dataset. The dataset contains 15,976 common genes among 9 brain regions. According to original cell type annotation, we computed the gene enrichment scores for each brain region. The 9 brain regions are: Frontal Cortex (EH4548), Cerebellum (EH4549), Entopeduncular (EH4550), Globus Pallidus (EH4551), Posterior Cortex (EH4552), Striatum (EH4553), Substantia Nigra (EH4554), Thalamus (EH4555) and Hippocampus (EH4556).

**Human Prefrontal Cortex Marker Genes** Gene enrichment scores calculated from healthy human prefrontal cortex scRNA-seq dataset (EH4557). Original paper annotated the cell types based on prior knowledge which contains 26 valid sub-cell types (clusters).

#### References

Saunders et al. (2018). Molecular Diversity and Specializations among the Cells of the Adult Mouse Brain. *Cell* 174(4), 1015-1030

Nagy et al. (2020) Single-nucleus transcriptomics of the prefrontal cortex in major depressive disorder implicates oligodendrocyte precursor cells and excitatory neurons. *Nature Neuroscience* 1-11

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## Examples

```
library(ExperimentHub)
eh <- ExperimentHub()
myfiles <- query(eh, "LRcellTypeMarkers")
## download resource
myfiles[[1]] ## load the first resource
myfiles[['EH4548']] ## load by EH id</pre>
```

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