

# Package ‘HCADatA’

January 22, 2026

**Type** Package

**Title** Accessing The Datasets Of The Human Cell Atlas in R/Bioconductor

**Version** 1.27.0

**Description** This package allows a direct access to the dataset generated by the Human Cell Atlas project for further processing in R and Bioconductor, in the comfortable format of SingleCellExperiment objects (available in other formats here: <http://preview.data.humancellatlas.org/>).

**License** MIT + file LICENSE

**Encoding** UTF-8

**biocViews** RNASeqData, SingleCellData, ExperimentData, ExpressionData, ExperimentHub

**Depends** R (>= 3.6), SingleCellExperiment

**Imports** ExperimentHub, AnnotationHub, HDF5Array, utils

**Suggests** knitr, rmarkdown, BiocStyle, scran (>= 1.11.4), BiocSingular, scater, scuttle, Rtsne, igraph, iSEE, testthat

**URL** <https://github.com/federicomarini/HCADatA>

**VignetteBuilder** knitr

**RoxygenNote** 7.3.2

**Roxygen** list(markdown = TRUE)

**git\_url** <https://git.bioconductor.org/packages/HCADatA>

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HCAData	<i>Download data from the HCA via ExperimentHub</i>
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### Description

Download HDF5 (dense assay) and RDS (row and column annotations) files from the HCA via ExperimentHub, composing them together as a SingleCellExperiment object

### Usage

```
HCAData(dataset = NULL, as.sparse = TRUE)
```

### Arguments

dataset	A character string: which dataset should be retrieved?
as.sparse	Logical, specifies whether the underlying HDF5 dataset should be treated as sparse or not - will be passed to the call to <code>HDF5Array()</code> . Defaults to TRUE, i.e. by using the <code>DelayedArray</code> infrastructure.

### Details

This current release includes the following datasets:

**Census of Immune Cells - Umbilical cord blood** UMI counts from the 10x (droplet) single-cell RNA-seq data. The object contains counts for 33694 genes in 384000 cells.

**Census of Immune Cells - Bone marrow** UMI counts from the 10x (droplet) single-cell RNA-seq data. The object contains counts for 33694 genes in 378000 cells.

### Value

A SingleCellExperiment object with a `HDF5Matrix` in the `counts` assay, containing the UMI counts for each gene in each cell. Row- and column-level metadata are also provided.

### Examples

```
HCAData()
sce_cordblood <- HCAData("ica_cord_blood")
```

**Description**

HCAData is an ExperimentHub package which provides access to the single-cell RNA-seq data from the Human Cell Atlas project (<https://www.humancellatlas.org>)

**Author(s)**

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**References**

If you use the data in this package, please refer to the original sources (Human Cell Atlas Data Portal, <https://preview.data.humancellatlas.org>) as well (plus the related publications, which will be listed here when they will be out), which are licensed under a Creative Commons Attribution 4.0 International License.

**See Also**

Useful links:

- <https://github.com/federicomarini/HCAData>

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