

Package ‘cellNexus’

June 23, 2026

Type Package

Title Queries the Human Cell Atlas

Version 0.99.23

Description Provides access to a copy of the Human Cell Atlas, but with harmonised metadata. This allows for uniform querying across numerous datasets within the Atlas using common fields such as cell type, tissue type, and patient ethnicity. Usage involves first querying the metadata table for cells of interest, and then downloading the corresponding cells into one of several supported formats, including SingleCellExperiment, Seurat, or pseudobulk SummarizedExperiment objects. Note: The Human Cell Atlas data accessed through this package is subject to its own licensing terms (typically Creative Commons Attribution or similar open data licenses as specified by the Human Cell Atlas Data Use Agreement), which differ from the package license (GPL-3). See the package documentation for details.

License GPL-3

Depends R (>= 4.5.0)

Imports dplyr, SummarizedExperiment, SingleCellExperiment, purrr (>= 1.0.0), BiocGenerics, glue, HDF5Array, DBI, tools, httr, curl, cli, SeuratObject, Seurat, methods, rlang, S4Vectors, tibble, utils, dbplyr (>= 2.3.0), duckdb, checkmate, shiny, shinyWidgets, zellkonverter, anndataR, stringr, Matrix, rclipboard

Suggests BiocStyle, knitr, rmarkdown, testthat, basilisk, arrow, reticulate, spelling, forcats, ggplot2, rprojroot, openssl, DelayedArray, tidybulk, BiocParallel, preprocessCore

Biarch true

biocViews AssayDomain, Infrastructure, SingleCell, RNASeq, DifferentialExpression, GeneExpression, Normalization, Clustering, QualityControl, Sequencing, Transcription, Transcriptomics, ShinyApps

Encoding UTF-8

RoxygenNote 7.3.3

URL <https://github.com/MangiolaLaboratory/cellNexus>,
<https://mangiolalaboratory.github.io/cellNexus/>

BugReports <https://github.com/MangiolaLaboratory/cellNexus/issues>

VignetteBuilder knitr

Roxygen list(markdown = TRUE)

Collate 'utils.R' 'counts.R' 'counts_per_million.R' 'data.R' 'dev.R'
'interface_app.R' 'metadata.R' 'seurat.R' 'unharmonised.R'
'zzz.R'

Language en-US

SystemRequirements Python

git_url <https://git.bioconductor.org/packages/cellNexus>

git_branch devel

git_last_commit d7f0137

git_last_commit_date 2026-05-26

Repository Bioconductor 3.24

Date/Publication 2026-06-23

Author Stefano Mangiola [aut, rev, ctb] (ORCID:
<<https://orcid.org/0000-0001-7474-836X>>),
Mengyuan Shen [aut, cre, rev] (ORCID:
<<https://orcid.org/0009-0008-0057-8239>>),
Michael Milton [aut, rev, ctb] (ORCID:
<<https://orcid.org/0000-0002-8965-2595>>),
Jared Andrews [aut, rev, ctb],
Juan Henao [aut, ctb],
Edward Yang [aut, ctb],
Julie Iskander [rev],
Silicon Valley Foundation CZF2019-002443 [fnd],
NIH NHGRI 5U24HG004059-18 [fnd],
Victoria Cancer Agency ECRF21036 [fnd],
NHMRC 1116955 [fnd]

Maintainer Mengyuan Shen <shen.m@wehi.edu.au>

Contents

cellNexus-package	3
create_interface_app	4
get_atlas_versions	5
get_cell_communication_strength	6
get_counts_per_million	7
get_default_cache_dir	8
get_metadata	8
get_metadata_url	10
get_pseudobulk	11
get_seurat	12
get_SingleCellExperiment	13
get_single_cell_experiment	15
get_unharmonised_dataset	16
join_census_table	17
keep_quality_cells	18
organize_inputs	19
pbmc3k_sce	20

SAMPLE_DATABASE_URL	21
ui_choices	22
upload_swift	23

Index	24
--------------	-----------

cellNexus-package	<i>cellNexus: Query Interface for the Human Cell Atlas</i>
-------------------	--

Description

cellNexus provides a query interface for programmatic exploration and retrieval of the harmonised, curated and reannotated CELLxGENE single-cell Human Cell Atlas. The package allows users to query metadata and download single-cell RNA sequencing data in various formats including SingleCellExperiment, Seurat, and pseudobulk SummarizedExperiment objects.

Value

The cellNexus package (invisibly).

Getting Started

To get started with cellNexus, first load the package and retrieve the metadata:

```
library(cellNexus)
metadata <- get_metadata()
```

Then filter the metadata to find cells of interest and download the data:

```
filtered_metadata <- metadata |>
  dplyr::filter(
    tissue == "lung parenchyma" &
    cell_type %LIKE% "%CD4%"
  )

sce <- get_single_cell_experiment(filtered_metadata)
```

Main Functions

[get_metadata](#) Retrieve and query the harmonised metadata
[get_single_cell_experiment](#) Download data as SingleCellExperiment objects
[get_seurat](#) Download data as Seurat objects
[get_pseudobulk](#) Download aggregated pseudobulk data

Data Licensing

Important: The Human Cell Atlas data accessed through this package is subject to its own licensing terms, which differ from the package license. The cellNexus package itself is licensed under GPL-3. However, the underlying Human Cell Atlas data is typically licensed under Creative Commons Attribution (CC-BY) or similar open data licenses as specified by the Human Cell Atlas Data Use Agreement. Users should review the specific licensing terms for any datasets they access through this package. For more information, see the Human Cell Atlas Data Portal: <https://data.humancellatlas.org/about>

Vignettes

See vignette("cellNexus", package = "cellNexus") for a comprehensive introduction to using the package.

Source

[Mangiola et al.,2023](#)

References

Mangiola, S., M. Milton, N. Ranathunga, C. S. N. Li-Wai-Suen, A. Odainic, E. Yang, W. Hutchison et al. "A multi-organ map of the human immune system across age, sex and ethnicity." bioRxiv (2023): 2023-06. doi:10.1101/2023.06.08.542671.

create_interface_app *Create a Shiny app that allows users to generate filtering & retrieval code for cellNexus*

Description

Create a Shiny app that allows users to generate filtering & retrieval code for cellNexus

Usage

```
create_interface_app(ui_choices, return_as_list = FALSE)
```

Arguments

`ui_choices` A list of pre-computed unique values for each filterable column in the metadata.
`return_as_list` If TRUE, returns a list with 'ui' and 'server' components instead of a Shiny app object.

Value

A Shiny app that allows users to filter cellNexus metadata and generate code for retrieval in the selected format.

Author(s)

Jared Andrews

Source

[Mangiola et al.,2023](#)

Examples

```
get_default_cache_dir()

# Create the interface app with metadata
metadata <- get_metadata(cloud_metadata = SAMPLE_DATABASE_URL)
app <- create_interface_app(metadata)
# Run the app
shiny::runApp(app)
```

get_atlas_versions *Returns the atlas version changelog as a tibble*

Description

Downloads the atlas_versions.parquet registry from the cellNexus metadata store, caches it locally, and returns it as an in-memory tibble. Each row describes one atlas data release and its relationship to a CellxGene Census snapshot.

Usage

```
get_atlas_versions(cache = tempdir())
```

Arguments

cache Optional character scalar. A local directory used to cache the downloaded parquet file. Defaults to a temporary directory to separate from the main cache directory.

Details

The atlas_id column in this table corresponds directly to the atlas_id column returned by [get_metadata\(\)](#), so you can join them to find which Census snapshot any cell in your query came from.

Value

A tibble with columns:

atlas_id Atlas version identifier, e.g. "cellxgene_2024/0.1.0". Matches the atlas_id column in [get_metadata\(\)](#).

census_version The CellxGene Census snapshot this atlas was built from, e.g. "01-07-2024".

change_type One of "initial", "patch", "minor", or "major". See ATLAS_VERSIONS.md for the conventions standards.

description Summary text of what changed in this release.

modified_at Modification date as a character scalar ("YYYY-MM-DD"). By default use Sys.Date()

Source

[Mangiola et al.,2023](#)

References

Mangiola, S., M. Milton, N. Ranathunga, C. S. N. Li-Wai-Suen, A. Odainic, E. Yang, W. Hutchison et al. "A multi-organ map of the human immune system across age, sex and ethnicity." *bioRxiv* (2023): 2023-06. doi:10.1101/2023.06.08.542671.

See Also

- CellxGene Census data releases (LTS): https://chanzuckerberg.github.io/cellxgene-census/cellxgene_census_docsite_data_release_info.html

Examples

```
get_atlas_versions()
```

```
get_cell_communication_strength
```

Retrieve cellNexus cell communication ligand–receptor strength as a data frame.

Description

Downloads a parquet database of the cell communication strength to a local cache, and then opens it as a data frame. It can then be filtered.

Usage

```
get_cell_communication_strength(
  cloud_metadata =
    get_metadata_url("cellNexus_lr_signaling_pathway_strength_DEMO.parquet"),
  local_metadata = NULL,
  cache_directory = get_default_cache_dir(),
  use_cache = TRUE
)
```

Arguments

cloud_metadata	Character vector of any length. HTTP URL/URLs pointing to the name and location of parquet database/databases. By default, it points to cell communication metadata in cellNexus ARDC Nectar Research Cloud. Assign NULL to query local_metadata only if exists.
local_metadata	Optional character vector of any length representing the local path of parquet database(s).
cache_directory	Optional character vector of length 1. A file path on your local system to a directory (not a file) that will be used to store metadata.parquet
use_cache	Optional logical scalar. If TRUE (the default), and this function has been called before with the same parameters, then a cached reference to the table will be returned. If FALSE, a new connect138/4.7ion will be created no matter what.

Details

The returned table integrates three levels of cell communication inference from CellChat, for each sample: (i) ligand–receptor–level communication (lr_prob, lr_pval), (ii) pathway-level aggregated signaling (pathway_prob, pathway_pval), (iii) cell-pair–level summaries of communication breadth (interaction_count - number of significant LR interactions) and intensity (interaction_weight - overall communication strength).

Together, these metrics allow simultaneous assessment of signaling specificity, pathway dominance, and global communication structure between cell populations.

Value

A lazy data.frame subclass containing the metadata. You can interact with this object using most standard dplyr functions. For string matching, it is recommended that you use stringr::str_like to filter character columns, as stringr::str_match will not work.

Examples

```
# For fast build purpose only, you do not need to specify anything in the function.
communication_meta <- get_cell_communication_strength(
  cloud_metadata = get_metadata_url(
    "cellNexus_lr_signaling_pathway_strength_DEMO.parquet"
  )
)
```

```
get_counts_per_million
```

Generating counts per million from a SingleCellExperiment object

Description

Generating counts per million from a SingleCellExperiment object

Usage

```
get_counts_per_million(sce, output_file)
```

Arguments

sce	A SingleCellExperiment object
output_file	A character vector of CPM Anndata file path

Value

A directory stores counts per million Anndata

Examples

```
data(pbmc3k_sce)
get_counts_per_million(pbmc3k_sce, tempfile(fileext = ".h5ad"))
```

`get_default_cache_dir` *Returns the default cache directory with a version number*

Description

Returns the default cache directory with a version number

Usage

```
get_default_cache_dir()
```

Value

A length one character vector.

Source

[Mangiola et al.,2023](#)

References

Mangiola, S., M. Milton, N. Ranathunga, C. S. N. Li-Wai-Suen, A. Odainic, E. Yang, W. Hutchison et al. "A multi-organ map of the human immune system across age, sex and ethnicity." *bioRxiv* (2023): 2023-06. doi:10.1101/2023.06.08.542671.

Examples

```
get_metadata(cloud_metadata = SAMPLE_DATABASE_URL, cache_directory = get_default_cache_dir())
```

`get_metadata` *Gets the CellNexus metadata as a data frame.*

Description

Downloads a parquet database of the Human Cell Atlas metadata to a local cache, and then opens it as a data frame. It can then be filtered and passed into `get_single_cell_experiment()` to obtain a `SingleCellExperiment::SingleCellExperiment`

Usage

```
get_metadata(  
  cloud_metadata = get_metadata_url("cellnexus_metadata.2.3.0.parquet"),  
  local_metadata = NULL,  
  cache_directory = get_default_cache_dir(),  
  use_cache = TRUE  
)
```

Arguments

cloud_metadata	Optional character vector of any length. HTTP URL/URLs pointing to the name and location of parquet database/databases. By default, it points to cellNexus ARDC Nectar Research Cloud. Assign NULL to query local_metadata only if exists.
local_metadata	Optional character vector of any length representing the local path of parquet database(s).
cache_directory	Optional character vector of length 1. A file path on your local system to a directory (not a file) that will be used to store metadata.
use_cache	Optional logical scalar. If TRUE (the default), and this function has been called before with the same parameters, then a cached reference to the table will be returned. If FALSE, a new connection will be created no matter what.

Details

The metadata was collected from the Bioconductor package `cellxgenedp`. `vignette("using_cellxgenedp", package="cellxgenedp")` provides an overview of the columns in the metadata. The data for which the column `organism_name` included "Homo sapiens" was collected from `cellxgenedp`.

The columns `dataset_id` and `file_id_cellNexus_single_cell` link the datasets explorable through cellNexus and `cellxgenedp` to the CELLxGENE portal.

Our representation, harmonises the metadata at dataset, sample and cell levels, in a unique coherent database table.

Field definitions for the CELLxGENE schema follow the [CELLxGENE schema 5.1.0](#).

Through harmonisation and curation we introduced custom columns not present in the original CELLxGENE metadata:

`cell_count`: Number of cells in a dataset. `feature_count`: Number of genes in a dataset.
`age_days`: Donor age in days. `tissue_groups`: Coarse tissue grouping for analysis. `empty_droplet`: Whether a cell is called an empty droplet from expressed-gene count per sample (default threshold 200; targeted panels may differ). `alive`: Whether a cell passes viability / mitochondrial QC.
`scDb1Finder.class`: Doublet, singlet, or unknown (scDb1Finder default parameters). `cell_type_unified_ensemble`: Consensus immune identity from Azimuth and SingleR (Blueprint, Monaco). `cell_annotation_azimuth_l2`: Azimuth cell annotation. `cell_annotation_blueprint_singler`: SingleR annotation (Blueprint).
`cell_annotation_blueprint_monaco`: SingleR annotation (Monaco). `is_immune`: Whether a cell is an immune cell. `sample_heuristic`: Internal sample subdivision helper. `file_id_cellNexus_single_cell`: Internal file id for single-cell layers. `file_id_cellNexus_pseudobulk`: Internal file id for pseudobulk layers. `sample_id`: Harmonised sample identifier. `nCount_RNA`: Total RNA counts per cell (sample-aware). `nFeature_expressed_in_sample`: Number of expressed features per cell.
`ethnicity_flagging_score`: Supporting score for ethnicity imputation. `low_confidence_ethnicity`: Supporting flag for low-confidence ethnicity calls. `.aggregated_cells`: Post-QC cells combined into each pseudobulk sample. `imputed_ethnicity`: Imputed ethnicity label. `atlas_id`: cellNexus atlas release identifier (internal use).

For all fields definitions, please refer to our [documentation site](#)

Possible cache path issues

If your default R cache path includes non-standard characters (e.g. dash because of your user or organisation name), the following error can occur.

```
Error in `db_query_fields.DBIconnection()` : ! Can't query fields. Caused by
error: ! Parser Error: syntax error at or near "/" LINE 2: FROM
/Users/bob/Library/Caches...
```

The solution is to choose a different cache, for example

```
get_metadata(cache_directory = path.expand('~'))
```

Value

A lazy data.frame subclass containing the metadata. You can interact with this object using most standard dplyr functions. For string matching, it is recommended that you use `stringr::str_like` to filter character columns, as `stringr::str_match` will not work.

Source

[Mangiola et al.,2023](#)

References

Mangiola, S., M. Milton, N. Ranathunga, C. S. N. Li-Wai-Suen, A. Odainic, E. Yang, W. Hutchison et al. "A multi-organ map of the human immune system across age, sex and ethnicity." *bioRxiv* (2023): 2023-06. doi:10.1101/2023.06.08.542671.

Examples

```
library(dplyr)
# For fast build purpose only, you do not need to specify anything in cloud_metadata.
filtered_metadata <- get_metadata(cloud_metadata = SAMPLE_DATABASE_URL) |>
  filter(
    self_reported_ethnicity == "African" &
    assay %LIKE% "%10x%" &
    tissue == "lung parenchyma" &
    cell_type %LIKE% "%CD4%"
  )
```

get_metadata_url	<i>Returns the URLs for all metadata files</i>
------------------	--

Description

Returns the URLs for all metadata files

Usage

```
get_metadata_url(
  databases = c("cellnexus_metadata.2.3.0.parquet", "census_cell_metadata.2.3.0.parquet")
)
```

Arguments

databases A character vector specifying the names of the metadata files. Download the specific metadata by defining the metadata version.

Value

A character vector of URLs to parquet files to download

Source

[Mangiola et al.,2023](#)

References

Mangiola, S., M. Milton, N. Ranathunga, C. S. N. Li-Wai-Suen, A. Odainic, E. Yang, W. Hutchison et al. "A multi-organ map of the human immune system across age, sex and ethnicity." *bioRxiv* (2023): 2023-06. doi:10.1101/2023.06.08.542671.

Examples

```
get_metadata_url("cellnexus_metadata.2.3.0.parquet")
```

get_pseudobulk	<i>Gets a Pseudobulk from curated metadata</i>
----------------	--

Description

Given a data frame of Curated Atlas metadata obtained from `get_metadata()`, returns a `SummarizedExperiment::Summ` object corresponding to the samples in that data frame

Usage

```
get_pseudobulk(
  data,
  assays = "counts",
  cell_aggregation = "pseudobulk",
  cache_directory = get_default_cache_dir(),
  repository = COUNTS_URL,
  features = NULL,
  as_SummarizedExperiment = FALSE
)
```

Arguments

<code>data</code>	A data frame containing, at minimum, <code>cell_id</code> , <code>file_id_cellNexus_pseudobulk</code> , <code>sample_id</code> , <code>cell_type_unified_ensemble</code> , <code>atlas_id</code> columns, which correspond to a single cell ID, file subdivision for internal use, a single cell sample ID, harmonised cell type, and atlas name in format (e.g <code>cellxgene_2024/0.1.0</code>) for internal use. They can be obtained from the <code>get_metadata()</code> function. Use <code>get_atlas_versions()</code> to download atlas versions data frame.
<code>assays</code>	A character vector specifying the desired assay(s) to be requested. The default setting retrieves only the counts assay.
<code>cell_aggregation</code>	A character vector that specifies which cell aggregation strategy should be applied. This will create a corresponding subdirectory in the cache directory.
<code>cache_directory</code>	An optional character vector of length one. If provided, it should indicate a local file path where any remotely accessed files should be copied.

repository	A character vector of length one. If provided, it should be an HTTP URL pointing to the location where the single cell data is stored.
features	An optional character vector of features (ie genes) to return the counts for. By default counts for all features will be returned. When provided, the returned object will contain exactly the requested features (row order preserved), and any experiments/samples that do not contain all requested features are dropped. This preserves the full set of requested features at the cost of potentially fewer samples. A warning is emitted when samples are dropped.
as_SummarizedExperiment	If TRUE, coerce the result to a SummarizedExperiment. Note that as(x, "SummarizedExperiment") drops feature rownames; get_pseudobulk() restores them after coercion.

Value

By default, a SingleCellExperiment object. If as_SummarizedExperiment is TRUE, a SummarizedExperiment object.

Source

[Mangiola et al.,2023](#)

References

Mangiola, S., M. Milton, N. Ranathunga, C. S. N. Li-Wai-Suen, A. Odainic, E. Yang, W. Hutchison et al. "A multi-organ map of the human immune system across age, sex and ethnicity." *bioRxiv* (2023): 2023-06. doi:10.1101/2023.06.08.542671.

Examples

```
# Use the lightweight sample database URL (for fast checks during development only)
meta <- get_metadata(cloud_metadata = cellNexus::SAMPLE_DATABASE_URL) |>
  keep_quality_cells() |>
  dplyr::filter(cell_type_unified_ensemble == "epithelial")
pseudobulk <- meta |> get_pseudobulk()
```

get_seurat	<i>Given a data frame of HCA metadata, returns a Seurat object corresponding to the samples in that data frame</i>
------------	--

Description

Given a data frame of HCA metadata, returns a Seurat object corresponding to the samples in that data frame

Usage

```
get_seurat(...)
```

Arguments

... Arguments passed on to [get_single_cell_experiment](#)

data A data frame containing, at minimum, `cell_id`, `file_id_cellNexus_single_cell` and `atlas_id` columns, which correspond to a single cell ID, file subdivision for internal use, and atlas name in format (e.g `cellxgene_2024/0.1.0`) for internal use. They can be obtained from the [get_metadata\(\)](#) function. Use [get_atlas_versions\(\)](#) to download atlas versions data frame.

assays A character vector specifying the desired assay(s) to be requested. Valid elements include "counts", "cpm", "rank", and "sct" for single-cell analyses. The default setting retrieves only the counts assay. If your analysis involves a smaller set of genes, consider using the "cpm" assay. The "rank" assay is suited for signature calculations across millions of cells.

cell_aggregation A character vector that specifies which cell aggregation strategy should be applied. This will create a corresponding subdirectory in the cache directory. Single cell level is applied by default.

cache_directory An optional character vector of length one. If provided, it should indicate a local file path where any remotely accessed files should be copied.

repository A character vector of length one. If provided, it should be an HTTP URL pointing to the location where the single cell data is stored.

features An optional character vector of features (ie genes) to return the counts for. By default counts for all features will be returned.

Value

A Seurat object containing the same data as a call to [get_single_cell_experiment\(\)](#)

Source

[Mangiola et al.,2023](#)

References

Mangiola, S., M. Milton, N. Ranathunga, C. S. N. Li-Wai-Suen, A. Odainic, E. Yang, W. Hutchison et al. "A multi-organ map of the human immune system across age, sex and ethnicity." *bioRxiv* (2023): 2023-06. doi:10.1101/2023.06.08.542671.

Examples

```
# Use the lightweight sample database URL (for fast checks during development only)
meta <- get_metadata(cloud_metadata = cellNexus::SAMPLE_DATABASE_URL) |> head(2)
seurat <- get_seurat(meta)
```

get_SingleCellExperiment

Gets a SingleCellExperiment from curated metadata

Description

Given a data frame of Curated Atlas metadata obtained from [get_metadata\(\)](#), returns a `SingleCellExperiment::SingleCellExperiment` object corresponding to the samples in that data frame

Usage

```
get_SingleCellExperiment(...)
```

Arguments

... Arguments passed on to [get_single_cell_experiment](#)

data A data frame containing, at minimum, `cell_id`, `file_id_cellNexus_single_cell` and `atlas_id` columns, which correspond to a single cell ID, file subdivision for internal use, and atlas name in format (e.g `cellxgene_2024/0.1.0`) for internal use. They can be obtained from the [get_metadata\(\)](#) function. Use [get_atlas_versions\(\)](#) to download atlas versions data frame.

assays A character vector specifying the desired assay(s) to be requested. Valid elements include "counts", "cpm", "rank", and "sct" for single-cell analyses. The default setting retrieves only the counts assay. If your analysis involves a smaller set of genes, consider using the "cpm" assay. The "rank" assay is suited for signature calculations across millions of cells.

cell_aggregation A character vector that specifies which cell aggregation strategy should be applied. This will create a corresponding subdirectory in the cache directory. Single cell level is applied by default.

cache_directory An optional character vector of length one. If provided, it should indicate a local file path where any remotely accessed files should be copied.

repository A character vector of length one. If provided, it should be an HTTP URL pointing to the location where the single cell data is stored.

features An optional character vector of features (ie genes) to return the counts for. By default counts for all features will be returned.

Value

A `SingleCellExperiment` object.

Source

[Mangiola et al., 2023](#)

References

Mangiola, S., M. Milton, N. Ranathunga, C. S. N. Li-Wai-Suen, A. Odainic, E. Yang, W. Hutchison et al. "A multi-organ map of the human immune system across age, sex and ethnicity." *bioRxiv* (2023): 2023-06. doi:10.1101/2023.06.08.542671.

Examples

```
# Use the lightweight sample database URL (for fast checks during development only)
meta <- get_metadata(cloud_metadata = cellNexus::SAMPLE_DATABASE_URL) |> head(2)
sce <- get_single_cell_experiment(meta)
```

`get_single_cell_experiment`*Gets a SingleCellExperiment from curated metadata*

Description

Given a data frame of Curated Atlas metadata obtained from `get_metadata()`, returns a `SingleCellExperiment::SingleCellExperiment` object corresponding to the samples in that data frame

Usage

```
get_single_cell_experiment(  
  data,  
  assays = "counts",  
  cell_aggregation = "",  
  cache_directory = get_default_cache_dir(),  
  repository = COUNTS_URL,  
  features = NULL  
)
```

Arguments

<code>data</code>	A data frame containing, at minimum, <code>cell_id</code> , <code>file_id_cellNexus_single_cell</code> and <code>atlas_id</code> columns, which correspond to a single cell ID, file subdivision for internal use, and atlas name in format (e.g <code>cellxgene_2024/0.1.0</code>) for internal use. They can be obtained from the <code>get_metadata()</code> function. Use <code>get_atlas_versions()</code> to download atlas versions data frame.
<code>assays</code>	A character vector specifying the desired assay(s) to be requested. Valid elements include "counts", "cpm", "rank", and "sct" for single-cell analyses. The default setting retrieves only the counts assay. If your analysis involves a smaller set of genes, consider using the "cpm" assay. The "rank" assay is suited for signature calculations across millions of cells.
<code>cell_aggregation</code>	A character vector that specifies which cell aggregation strategy should be applied. This will create a corresponding subdirectory in the cache directory. Single cell level is applied by default.
<code>cache_directory</code>	An optional character vector of length one. If provided, it should indicate a local file path where any remotely accessed files should be copied.
<code>repository</code>	A character vector of length one. If provided, it should be an HTTP URL pointing to the location where the single cell data is stored.
<code>features</code>	An optional character vector of features (ie genes) to return the counts for. By default counts for all features will be returned.

Value

A `SingleCellExperiment` object.

Source

[Mangiola et al.,2023](#)

References

Mangiola, S., M. Milton, N. Ranathunga, C. S. N. Li-Wai-Suen, A. Odainic, E. Yang, W. Hutchison et al. "A multi-organ map of the human immune system across age, sex and ethnicity." *bioRxiv* (2023): 2023-06. doi:10.1101/2023.06.08.542671.

Examples

```
# Use the lightweight sample database URL (for fast checks during development only)
meta <- get_metadata(cloud_metadata = cellNexus::SAMPLE_DATABASE_URL) |> head(2)
sce <- get_single_cell_experiment(meta)
```

```
get_unharmonised_dataset
```

Returns unharmonised metadata for selected datasets.

Description

Various metadata fields are *not* common between datasets, so it does not make sense for these to live in the main metadata table. This function is a utility that allows easy fetching of this data if necessary.

Usage

```
get_unharmonised_dataset(
  dataset_id,
  cells = NULL,
  conn = dbConnect(drv = duckdb(), read_only = TRUE),
  remote_url = UNHARMONISED_URL,
  cache_directory = get_default_cache_dir()
)
```

Arguments

dataset_id	A character vector, where each entry is a dataset ID obtained from the <code>\$file_id_cellNexus_single</code> column of the table returned from <code>get_metadata()</code>
cells	An optional character vector of cell IDs. If provided, only metadata for those cells will be returned.
conn	An optional DuckDB connection object. If provided, it will re-use the existing connection instead of opening a new one.
remote_url	Optional character vector of length 1. An HTTP URL pointing to the root URL under which all the unharmonised dataset files are located.
cache_directory	Optional character vector of length 1. A file path on your local system to a directory (not a file) that will be used to store the unharmonised metadata files.

Value

A named list, where each name is a dataset file ID, and each value is a "lazy data frame", ie a tbl.

Source

Mangiola et al.,2023

References

Mangiola, S., M. Milton, N. Ranathunga, C. S. N. Li-Wai-Suen, A. Odainic, E. Yang, W. Hutchison et al. "A multi-organ map of the human immune system across age, sex and ethnicity." *bioRxiv* (2023): 2023-06. doi:10.1101/2023.06.08.542671.

Examples

```
## Not run:
dataset <- "838ea006-2369-4e2c-b426-b2a744a2b02b"
harmonised_meta <- get_metadata() |>
  dplyr::filter(file_id_cellNexus_single_cell == dataset) |>
  dplyr::collect()
unharmonised_meta <- get_unharmonised_dataset(dataset)
unharmonised_tbl <- dplyr::collect(unharmonised_meta[[dataset]])
dplyr::left_join(harmonised_meta, unharmonised_tbl,
  by = c("file_id_cellNexus_single_cell", "cell_id")
)

## End(Not run)
```

join_census_table	<i>Join Census metadata to an existing data frame</i>
-------------------	---

Description

Downloads and joins the Census metadata with cellNexus metadata. This function creates indexed tables for efficient joining and returns a data frame.

Usage

```
join_census_table(
  tbl,
  cloud_metadata = get_metadata_url("census_cell_metadata.2.3.0.parquet"),
  cache_directory = get_default_cache_dir(),
  join_keys = c("sample_id", "dataset_id", "observation_joinid")
)
```

Arguments

tbl	A tbl_sql object (from get_metadata) or a database connection.
cloud_metadata	HTTP URL/URLs pointing to the census parquet database name and location.
cache_directory	A character string specifying the local cache directory where remote parquet files will be stored. Defaults to <code>get_default_cache_dir()</code> .
join_keys	A character vector of column names used for the join. Defaults to <code>c("sample_id", "dataset_id", "observation_joinid")</code> .

Value

A lazy SQL table with Census metadata joined to the cellNexus metadata.

Examples

```
library(dplyr)
get_metadata(cloud_metadata = SAMPLE_DATABASE_URL) |> head(2) |>
  # You do not need to specify anything in cloud_metadata
  join_census_table(
    cloud_metadata = SAMPLE_DATABASE_URL,
    cache_directory = tempdir()
  )
```

keep_quality_cells	<i>Keep high-quality cells based on QC columns</i>
--------------------	--

Description

Keep high-quality cells based on QC columns

Usage

```
keep_quality_cells(
  data,
  empty_droplet_col = "empty_droplet",
  alive_col = "alive",
  doublet_col = "scDbfFinder.class"
)
```

Arguments

data	A data frame or tibble containing single-cell metadata.
empty_droplet_col	A string specifying the column name that indicates empty droplets (default: "empty_droplet"). Expected logical vector
alive_col	A string specifying the column name that indicates whether cells are alive (default: "alive"). Expected logical vector
doublet_col	A string specifying the column name that indicates doublets (default: "scDbfFinder.class"). Expected character vector: "doublet" and/or "singlet" and/or "unknown".

Value

A filtered data frame containing only cells that pass all QC checks.

Source

[Mangiola et al.,2023](#)

Examples

```
get_metadata(cloud_metadata = SAMPLE_DATABASE_URL, cache_directory = tempdir()) |>
  head(2) |>
  keep_quality_cells()
```

organize_inputs	<i>Organize arbitrary Shiny inputs into a grid layout</i>
-----------------	---

Description

Organize arbitrary Shiny inputs into a grid layout

Usage

```
organize_inputs(  
  tag.list,  
  id = NULL,  
  title = NULL,  
  tack = NULL,  
  columns = NULL,  
  rows = NULL  
)
```

Arguments

<code>tag.list</code>	A tagList containing UI inputs or a named list containing multiple tagLists containing UI inputs.
<code>id</code>	An optional ID for the tabsetPanel if a named list is provided.
<code>title</code>	An optional title for the grid, should be a UI element, e.g. <code>h3("Title")</code> .
<code>tack</code>	An optional UI input to tack onto the end of the grid.
<code>columns</code>	Number of columns.
<code>rows</code>	Number of rows.

Value

A Shiny tagList with inputs organized into a grid, optionally nested inside a tabsetPanel.

Author(s)

Jared Andrews

Source

[Mangiola et al.,2023](#)

Examples

```
library(shiny)  
# Example 1: Basic usage with a simple grid  
ui.inputs <- tagList(  
  textInput("name", "Name"),  
  numericInput("age", "Age", value = 30),  
  selectInput("gender", "Gender", choices = c("Male", "Female", "Other"))  
)  
organize_inputs(ui.inputs, columns = 2, rows = 2)
```

```

# Example 2: Using a named list to create tabs
ui.inputs.tabs <- list(
  Personal = tagList(
    textInput("firstname", "First Name"),
    textInput("lastname", "Last Name")
  ),
  Settings = tagList(
    checkboxInput("newsletter", "Subscribe to newsletter", value = TRUE),
    sliderInput("volume", "Volume", min = 0, max = 100, value = 50)
  )
)
organize_inputs(ui.inputs.tabs, columns = 2)

# Example 3: Adding an additional UI element with 'tack'
additional.ui <- actionButton("submit", "Submit")
organize_inputs(ui.inputs, tack = additional.ui, columns = 3)

# Example 4: Handling a case with more inputs than grid cells
many.inputs <- tagList(replicate(10, textInput("input", "Input")))
organize_inputs(many.inputs, columns = 3) # Creates more than one row

```

pbmc3k_sce

Sample SingleCellExperiment Object

Description

A sample `SingleCellExperiment` object created from the pbmc3k dataset for testing and demonstration purposes. The dataset contains 500 cells with gene expression data mapped to Ensembl gene IDs and formatted with cellNexus-compatible metadata structure.

Format

An object of class `SingleCellExperiment` with:

assays Gene expression matrix with Ensembl gene IDs as rownames

colData Cell metadata including `sample_id`, `cell_type_unified_ensemble`, `nCount_RNA`, etc.

metadata List containing 'data' field with cellNexus-formatted metadata including:

- `cell_id`: Unique cell identifier
- `sample_id`: Sample identifier
- `cell_type_unified_ensemble`: Cell type annotation
- `nCount_RNA`: Number of RNA molecules per cell
- `ident`: Seurat cluster identity
- `dataset_id`: Dataset identifier
- `file_id_cellNexus_single_cell`: Generated file ID for cellNexus
- `atlas_id`: Atlas identifier with date

Details

See `dev/create_pbmc3k_sce.R` for the complete creation script.

Source

Created from pbmc3k dataset (SeuratData package)

References

Mangiola, S., M. Milton, N. Ranathunga, C. S. N. Li-Wai-Suen, A. Odainic, E. Yang, W. Hutchison et al. "A multi-organ map of the human immune system across age, sex and ethnicity." bioRxiv (2023): 2023-06. doi:10.1101/2023.06.08.542671.

Examples

```
data(pbmc3k_sce)
pbmc3k_sce
# Access metadata
S4Vectors::metadata(pbmc3k_sce)$data
```

SAMPLE_DATABASE_URL *URL pointing to the sample metadata file, which is smaller and for test, demonstration, and vignette purposes only*

Description

URL pointing to the sample metadata file, which is smaller and for test, demonstration, and vignette purposes only

Usage

```
SAMPLE_DATABASE_URL
```

Format

An object of class character of length 2.

Value

Character scalar consisting of the URL/URLs

Source

[Mangiola et al.,2023](#)

References

Mangiola, S., M. Milton, N. Ranathunga, C. S. N. Li-Wai-Suen, A. Odainic, E. Yang, W. Hutchison et al. "A multi-organ map of the human immune system across age, sex and ethnicity." bioRxiv (2023): 2023-06. doi:10.1101/2023.06.08.542671.

Examples

```
get_metadata(cloud_metadata = SAMPLE_DATABASE_URL["cellnexus"], cache_directory = tempdir())
```

`ui_choices`*Pre-computed UI Choices for Interface App*

Description

A list of unique values for each filterable column used in the cellNexus Shiny interface app. Pre-computing these choices avoids slow metadata queries when the app starts.

Usage

```
data(ui_choices)
```

Format

A named list where each element contains unique values for a column:

cell_type_unified_ensemble Character vector of unified cell type labels

cell_type Character vector of original cell type labels

alive Logical values for cell viability

scDbfFinder.class Character vector of doublet classification results

is_immune Logical values for immune cell classification

empty_droplet Logical values for empty droplet detection

development_stage Character vector of developmental stages

disease Character vector of disease states

self_reported_ethnicity Character vector of ethnicity labels

sex Character vector of sex labels

tissue Character vector of tissue types

tissue_groups Character vector of tissue group labels

Details

See `dev/generate_ui_choices.R` for the creation script. Run this script to regenerate the choices when metadata columns change.

Source

Generated from cellNexus metadata

upload_swift	<i>Upload a file to the Nectar object store</i>
--------------	---

Description

Upload a file to the Nectar object store

Usage

```
upload_swift(  
    source,  
    container,  
    name = basename(source),  
    credential_id = NULL,  
    credential_secret = NULL  
)
```

Arguments

source	A character scalar indicating the local path to the file to upload
container	A character scalar indicating the name of the container to upload to
name	An optional character scalar indicating the name the file should have after being uploaded. Defaults to being the basename of the source file.
credential_id	The OpenStack application credential secret as a character scalar

Value

NULL, invisibly

Examples

```
## Not run:  
upload_swift(  
    "/vast/projects/cellxgene_curated/metadata_parquet_0.2",  
    "cellNexus-metadata",  
    credential_id = "ABCDEFGHIJK",  
    credential_secret = "ABCD1234EFGH-5678IJK"  
)  
  
## End(Not run)
```

Index

- * **datasets**
 - pbmc3k_sce, [20](#)
 - SAMPLE_DATABASE_URL, [21](#)
 - ui_choices, [22](#)
- * **internal**
 - get_unharmonised_dataset, [16](#)
 - upload_swift, [23](#)
- cellNexus (cellNexus-package), [3](#)
- cellNexus-package, [3](#)
- create_interface_app, [4](#)
- get_atlas_versions, [5](#)
- get_cell_communication_strength, [6](#)
- get_counts_per_million, [7](#)
- get_default_cache_dir, [8](#)
- get_default_cache_dir(), [17](#)
- get_metadata, [3](#), [8](#)
- get_metadata(), [5](#), [11](#), [13–16](#)
- get_metadata_url, [10](#)
- get_pseudobulk, [3](#), [11](#)
- get_seurat, [3](#), [12](#)
- get_single_cell_experiment, [3](#), [13](#), [14](#), [15](#)
- get_single_cell_experiment(), [8](#), [13](#)
- get_SingleCellExperiment, [13](#)
- get_unharmonised_dataset, [16](#)
- join_census_table, [17](#)
- keep_quality_cells, [18](#)
- organize_inputs, [19](#)
- pbmc3k_sce, [20](#)
- SAMPLE_DATABASE_URL, [21](#)
- SingleCellExperiment::SingleCellExperiment,
[8](#), [13](#), [15](#)
- SummarizedExperiment::SummarizedExperiment,
[11](#)
- ui_choices, [22](#)
- upload_swift, [23](#)