# Package 'hca'

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Title Exploring the Human Cell Atlas Data Coordinating Platform

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**Description** This package provides users with the ability to query the Human Cell Atlas data repository for single-cell experiment data. The `projects()`, `files()`, `samples()` and `bundles()` functions retrieve summary information on each of these indexes; corresponding `\*\_details()` are available for individual entries of each index. File-based resources can be downloaded using `files\_download()`. Advanced use of the package allows the user to page through large result sets, and to flexibly query the 'list-of-lists' structure representing query responses.

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.details

Single Entity Details

# Description

Single Entity Details

# Usage

```
.details(
  uuid = character(),
  catalog = NULL,
  view = c("projects", "files", "samples", "bundles")
)
```

# Arguments

uuid	character() unique *_id
catalog	$\mbox{character}(1)$ source of data. Use $\mbox{catalogs}()$ for possible values.
view	character() type of entity i.e. project, file, sample, or bundle

# Value

list-of-lists containing relevant details about the project, file, sample, or bundle

bundles

# Description

bundles() takes a list of user provided project titles to be used to query the HCA API for information about available bundles.

bundles\_detail() takes a unique bundle\_id and catalog for the bundle, and returns details about the specified bundle as a list-of-lists

# Usage

```
bundles(
  filters = NULL,
  size = 100L,
  sort = "projectTitle",
  order = c("asc", "desc"),
  catalog = NULL,
  as = c("tibble", "lol", "list", "tibble_expanded"),
  columns = bundles_default_columns("character")
)
bundles_facets(facet = character(), catalog = NULL)
bundles_default_columns(as = c("tibble", "character"))
```

```
bundles_detail(uuid, catalog = NULL)
```

# Arguments

filters	filter object created by filters(), or NULL (default; all projects).
size	integer(1) maximum number of results to return; default: all projects matching filter. The default (10000) is meant to be large enough to return all results.
sort	<pre>character(1) project facet (see facet_options()) to sort result; default: "projectTitle".</pre>
order	character(1) sort order. One of "asc" (ascending) or "desc" (descending).
catalog	character(1) source of data. Use catalogs() for possible values.
as	character(1) return format. One of "tibble" (default), "lol", "list", or "tibble_expanded", as described in the Details and Value sections of ?projects.
columns	named character() indicating the paths to be used for parsing the 'lol' returned from the HCA to a tibble. The names of columns are used as column names in the returned tibble. If the columns are unnamed, a name is derived from the ele- ments of path by removing hits[*] and all [*], e.g., a path hits[*].donorOrganisms[*].biological is given the name donorOrganisms.biologicalSex.

facet	character() of valid facet names. Summary results (see 'Value', below) are re-
	turned when missing or length greater than 1; details are returned when a single
	facet is specified.
uuid	character() unique identifier (e.g., projectId) of the object.

#### Value

bundles\_detail() returns a list-of-lists containing relevant details about the bundle

#### Examples

```
title <- paste(
    "Tabula Muris: Transcriptomic characterization of 20 organs and",
    "tissues from Mus musculus at single cell resolution"
)
filters <- filters( projectTitle = list(is = title) )
bundles(filters = filters)
bundles_facets()
bundle <- bundles(size = 1, as = "list")
bundle_uuid <- bundle[["hits"]][[1]][["entryId"]]
bundles_detail(uuid = bundle_uuid) |> lol() |>
lol_filter(is_leaf) |> print(n = Inf)
```

catalogs

Catalogs Available in the HCA

#### Description

catalogs() queries the API for all available project catalogs

#### Usage

```
catalogs(catalog = NULL)
```

#### Arguments

catalog character(1) default catalog. When missing or NULL, the catalog defined by the Human Cell Atlas API is used; this is usually the most recently available catalog. Providing a non-null argument changes the default globally; restore default order by explicitly defining the arugment catalog = NULL.

#### Value

character() vector of available catalogs. The first is the default, defined by the API or by the user with argument catalog.

# files

# Examples

catalogs()

files

HCA File Querying

#### Description

files() takes a list of user provided project titles to be used to query the HCA API for information about available files.

files\_download() takes a tibble of files and a directory location as arguments to download the files of the tibble into the specified directory.

files\_detail() takes a unique file\_id and catalog for the file, and returns details about the specified file as a list-of-lists

files\_cache() is the default location of the cache of downloaded files.

# Usage

```
files(
    filters = NULL,
    size = 1000L,
    sort = "projectTitle",
    order = c("asc", "desc"),
    catalog = NULL,
    as = c("tibble", "lol", "list", "tibble_expanded"),
    columns = files_default_columns("character")
)
files_default_columns(as = c("tibble", "character"))
files_default_columns(as = c("tibble", "character"))
files_default_columns(as = null)
files_facets(facet = character(), catalog = null)
files_detail(uuid, catalog = null)
files_cache(create = FALSE)
```

## Arguments

filters	filter object created by filters(), or NULL (default; all projects).
size	integer(1) maximum number of results to return; default: all projects matching filter. The default (10000) is meant to be large enough to return all results.
sort	character(1) project facet (see facet_options()) to sort result; default: "projectTitle".

order	character(1) sort order. One of "asc" (ascending) or "desc" (descending).
catalog	character(1) source of data. Use catalogs() for possible values.
as	character(1) return format. One of "tibble" (default), "lol", "list", or "tibble_expanded", as described in the Details and Value sections of ?projects.
columns	named character() indicating the paths to be used for parsing the 'lol' returned from the HCA to a tibble. The names of columns are used as column names in the returned tibble. If the columns are unnamed, a name is derived from the ele- ments of path by removing hits[*] and all [*], e.g., a path hits[*].donorOrganisms[*].biological is given the name donorOrganisms.biologicalSex.
tbl	tibble of files (result of files())
destination	character() vector name of temporary directory to use for file downloads, or NULL
facet	character() of valid facet names. Summary results (see 'Value', below) are re- turned when missing or length greater than 1; details are returned when a single facet is specified.
uuid	character() unique identifier (e.g., projectId) of the object.
create	logical(1) create the default cache location, if it does not yet exist.

# Details

files\_cache() can be useful when it is necessary to 'clean up' the cache, e.g., BiocFileCache::cleanbfc()
or more dramatically unlink(files\_cache(), recursive = TRUE).

# Value

files\_download() returns a character() vector of file destinations

files\_detail() returns a list-of-lists containing relevant details about the file.

files\_cache() returns the path to the default cache. Use this as the cache= argument to BiocFileCache().

```
title <- paste(
    "Tabula Muris: Transcriptomic characterization of 20 organs and",
    "tissues from Mus musculus at single cell resolution"
)
filters <- filters( projectTitle = list(is = title) )
files(filters = filters)
files_filter <- filters(
    projectId = list(is = "cddab57b-6868-4be4-806f-395ed9dd635a"),
    fileFormat = list(is = "loom")
)
files_tbl <- files(filter = files_filter)
## Not run: files_download(files_tbl, destination = tempdir())
files_facets()
files_facets("fileFormat")</pre>
```

# filters

```
file <- files(size = 1, as = "list")
file_uuid <- file[["hits"]][[1]][["entryId"]]
files_detail(uuid = file_uuid)
files_cache(create = FALSE)</pre>
```

filters

HCA Filter Construction

# Description

facet\_options() returns a character vector of possible facets to use during filtering.

filters() takes user input to be used as query filters. Each named argument is a list with a name specifying a verb (e.g., "is") and a character vector of allowed values, as in the examples. This input is then validated, reformatted to JSON, and encoded into a properly formatted URL.

## Usage

facet\_options()

filters(...)

## S3 method for class 'filters'
length(x)

## S3 method for class 'filters'
print(x, ...)

# Arguments

	named arguments, each of which is a list() specifying a query facet and its
	corresponding value to be used in the query
x	for length() and print(), an object of class filters.

# Value

facet\_options() returns a vector of all permissible query facets for the HCA api.

filters() returns a filters object representing validated filters in a format suitable for use in projects() and related functions.

#### Examples

```
facet_options()
```

filters()

filters(
 organ = list(is = "pancreas")

```
)
filters(
    organ = list(is = "pancreas"),
    genusSpecies = list(is = "Homo sapiens")
)
filters(
    fileFormat = list(is = c("fastq", "fastq.gz"))
)
```

hca\_next

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# Page through HCA results

# Description

hca\_next() retrieves the next 'page' of results from a query of projects(), samples(), files(),
or bundles().

hca\_prev() returns the previous 'page' of results.

#### Usage

```
hca_next(x, size)
```

hca\_prev(x, size)

# Arguments

X	a 'tibble' or 'lol' object returned by projects(), samples(), files(), or bundles().
size	the (non-negative integer) number of elements to retrieve in the page request. The default is the number of elements requested in x.

# Value

hca\_next() returns the next page of results as a 'tibble' or 'lol' hcl\_prev() returns the previous page of results.

files <- files(size = 5)	# results 1–5, as a tibble
<pre>next_files &lt;- hca_next(files) next_files</pre>	# results 6-10
<pre>hca_prev(next_files)</pre>	<pre># previous results, i.e., files 1-5</pre>

hca\_next.list\_hca 'list' representation of HCA query results

# Description

projects(), samples(), files() and bundles() return results for the number of records indicated by the size= argument. Use as = "list" to return results as a "list\_hca" list.

hca\_next() returns a list containing the next 'page' of results.

hca\_prev() returns a list containing the previous 'page' of results.

#### Usage

```
## S3 method for class 'list_hca'
hca_next(x, size)
```

## S3 method for class 'list\_hca' hca\_prev(x, size)

# Arguments

х	a 'list' returned by projects(), samples(), files(), or bundles().
size	the (non-negative integer) number of elements to retrieve in the page request. The default is the number of elements requested in x.

# Value

hca\_next() returns a list containing the next 'page' of results.

hca\_prev() returns a list containing the previous 'page' of results.

```
projects <- projects(size = 5, as = "list") # projects 1-5
next_projects <- hca_next(projects) # projects 6-10
hca_prev(next_projects) # projects 1-5</pre>
```

hca\_next.tbl\_hca 'tibble' representation of HCA query results

# Description

projects(), samples(), files(), and bundles() return, by default, a 'tibble' representation of the query.

hca\_next() returns the next 'page' of results, if available.

hca\_prev() returns the previous 'page' of results.

# Usage

## S3 method for class 'tbl\_hca' hca\_next(x, size)

## S3 method for class 'tbl\_hca' hca\_prev(x, size)

# Arguments

х	a 'tibble' returned by projects(), samples(), files(), or bundles().
size	the (non-negative integer) number of elements to retrieve in the page request. The default is the number of elements requested in $x$ .

#### Value

hca\_next() returns a tibble, with the same columns as x, containing the next 'page' of results.

hca\_prev() returns a tibble with the same columns as x, containing the previous 'page' of results.

```
projects <- projects(size = 5)  # projects 1-5
next_projects <- hca_next(projects)  # projects 6-10
hca_prev(next_projects)  # projects 1-5
```

hca\_view

#### Description

View and select table rows interactively

# Usage

```
hca_view(tbl)
```

#### Arguments

tbl

a 'tibble' of projects(), samples(), bundles(), or files().

#### Value

hca\_view() returns a tibble filtered to reflect the rows selected in the interface.

#### Examples

```
if (interactive()) {
    p <- projects(size = 100)
    p1 <- hca_view(p) # interactive table browser; filtered results
}</pre>
```

lol

Representing and manipulating list-of-list data structures.

#### Description

lol() constructs an indexed representation of an R 'list-of-lists', typically from JSON queries. The object is conveniently manipulated by other functions on this page to filter and select subsets of the structure, and to pull individual paths from across the list-of-lists.

lol\_filter() filters available paths based on selections in ..., e.g., n (number of matching elements) or is\_leaf (is the element a 'leaf' in the list-of-lists representation?).

lol\_lpull() returns a list containing elements corresponding to a single path.

lol\_pull() tries to simplify the list-of-lists structure returned by lol\_lpull() to a vector.

lol\_path() returns a tibble representing the paths through the list-of-lists, without the underlying list-of-list data.

as.list() returns a list-of-lists representation of the data returned by projects(), etc.

hca\_next() returns the next 'page' of results, if available.

hca\_prev() returns the previous 'page' of results.

lol\_hits\_lpull() and lol\_hits\_pull() are variants of lol\_lpull() and lol\_pull() that retain the original geometry of hits[\*], even when the mapping between hits[\*] and path is not 1:1.

#### Usage

```
lol(x = list())
lol_select(x, path = character())
lol_filter(x, ...)
lol_lpull(x, path)
lol_pull(x, path)
lol_path(x)
## S3 method for class 'lol'
as.list(x, ...)
## S3 method for class 'lol'
print(x, ...)
## S3 method for class 'lol_hca'
hca_next(x, size)
## S3 method for class 'lol_hca'
hca_prev(x, size)
lol_hits_lpull(x, path)
lol_hits_pull(x, path)
```

#### Arguments

х	a 'list-of-lists' returned by projects(), samples(), files(), or bundles()
path	character(1) from the tibble returned by lol_path(x).
	for lol_filter(), named filter expressions evaluating to a logical vector with length equal to the number of rows in lol_path().
size	the (non-negative integer) number of elements to retrieve in the page request. The default is the number of elements requested in $x$ .

# Value

lol() returns a representation of the list-of-lists. The list has been processed to a dictionary with entries to all paths through the list, as well as a tibble summarizing the path, number of occurrences, and leaf status of each unique path.

lol\_select() returns an object of class "lol" subset to contain just the elements matching path as 'top-level' elements of the list-of-lists.

lol\_filter() returns an object of class lol, filtered to contain elements consistent with the filter criteria.

#### manifest

lol\_lpull() returns a list, where each element corresponds to an element found at path in the list-of-lists structure x.

lol\_pull() returns an unnamed vector of elements matching key.

hca\_next() returns a list-of-lists containing the next 'page' of results.

hca\_prev() returns a tibble with the same columns as x, containing the previous 'page' of results.

# Examples

```
plol <- projects(size = 5, as = "lol")</pre>
plol
plol |> lol_select("hits[*].projects[*]")
plol |>
   lol_select("hits[*].projects[*]") |>
   lol_filter(n == 44, is_leaf)
plol |>
    lol_pull("hits[*].entryId") |>
    head()
plol |> lol_path()
projects <- projects(size = 5, as = "lol")</pre>
                                                 # projects 1-5
next_projects <- hca_next(projects)</pre>
                                                 # projects 6-10
hca_prev(next_projects)
                                                  # projects 1-5
```

manifest

HCA File Querying

#### Description

manifest() takes a list of user provided project titles to be used to query the HCA API for information about available manifest files.

manifest\_cache() is the default location of the cache of downloaded manifest.

#### Usage

manifest(filters = NULL, catalog = NULL, update\_cache = FALSE)

```
manifest_cache(create = FALSE)
```

#### Arguments

filters	hca filter object
catalog	character() name of catalog
update_cache	logical(1) when TRUE, update an existing cached resource by querying the HCA data server.
create	logical(1) create the default cache location, if it does not yet exist.

# Details

manifest\_cache() can be useful when it is necessary to 'clean up' the cache, e.g., BiocFileCache::cleanbfc()
or more dramatically unlink(manifest\_cache(), recursive = TRUE).

# Value

manifest\_cache() returns the path to the default cache. Use this as the cache= argument to BiocFileCache().

#### Examples

```
manifest_filter <- hca::filters(
    projectId = list(is = "4a95101c-9ffc-4f30-a809-f04518a23803"),
    fileFormat = list(is = "loom"),
    workflow = list(is = c("optimus_v4.2.2", "optimus_v4.2.3"))
)
## Not run:
result <- manifest(manifest_filter)
result
## End(Not run)
manifest_cache(create = FALSE)</pre>
```

optimus\_loom\_annotation

HCA loom file annotation

#### Description

optimus\_loom\_annotation() takes the file path location of a .loom file generated by the Optimus pipeline, for which additional data will be extracted from the appropriate manifest. The .loom file will be imported as a LoomExperiment object, and the additional manifest information will be added to the object for return.

## projects

## Usage

```
optimus_loom_annotation(loom, catalog = NULL)
## S3 method for class 'character'
optimus_loom_annotation(loom, catalog = NULL)
## S3 method for class 'LoomExperiment'
optimus_loom_annotation(loom, catalog = NULL)
```

# Arguments

loom	Either a character(1) file path to a loom file on user's system, or a loom file ob-
	tained from the HCA and imported into R using LoomExperiment::import().
catalog	character() HCA catalog from which the .loom file originated.

#### Value

A 'LoomExperiment' object annotated with additional metadata() and colData() derived from the manifest file describing samples in the object.

#### See Also

manifest() and related functions for working with data returned from the \*/manifest/\* HCA API endpoints.

projects HCA Project Querying
-------------------------------

# Description

projects() takes user input to be used to query the HCA API for information about available projects.

projects\_facets() summarizes facets and terms used by all records in the projects index.
\*\_columns() returns a tibble or named character vector describing the content of the tibble returned
by projects(), files(), samples(), or bundles().

projects\_detail() takes a unique project\_id and catalog for the project, and returns details about the specified project as a list-of-lists

# Usage

```
projects(
  filters = NULL,
  size = 1000L,
  sort = "projectTitle",
  order = c("asc", "desc"),
  catalog = NULL,
```

```
as = c("tibble", "lol", "list", "tibble_expanded"),
columns = projects_default_columns("character")
)
projects_facets(facet = character(), catalog = NULL)
projects_default_columns(as = c("tibble", "character"))
projects_detail(uuid, catalog = NULL)
```

#### Arguments

filters	filter object created by filters(), or NULL (default; all projects).
size	integer(1) maximum number of results to return; default: all projects matching filter. The default (10000) is meant to be large enough to return all results.
sort	character(1) project facet (see facet_options()) to sort result; default: "projectTitle".
order	character(1) sort order. One of "asc" (ascending) or "desc" (descending).
catalog	character(1) source of data. Use catalogs() for possible values.
as	character(1) return format. One of "tibble" (default), "lol", "list", or "tibble_expanded", as described in the Details and Value sections of ?projects.
columns	named character() indicating the paths to be used for parsing the 'lol' returned from the HCA to a tibble. The names of columns are used as column names in the returned tibble. If the columns are unnamed, a name is derived from the ele- ments of path by removing hits[*] and all [*], e.g., a path hits[*].donorOrganisms[*].biological is given the name donorOrganisms.biologicalSex.
facet	character() of valid facet names. Summary results (see 'Value', below) are re- turned when missing or length greater than 1; details are returned when a single facet is specified.
uuid	character() unique identifier (e.g., projectId) of the object.

#### Details

The as argument determines the object returned by the function. Possible values are:

- "tibble" (default) A tibble (data.frame) summarizing essential elements of projects, samples, bundles, or files.
- "lol" A 'list-of-lists' representation of the JSON returned by the query as a 'list-of-lists' data structure, indexed and presented to enable convenient filtering, selection, and extraction. See ?lol.
- "list" An R list (typically, highly recursive) containing detailed project information, constructed from the JSON response to the original query.
- "tibble\_expanded" A tibble (data.frame) containing (almost) all information for each project, sample, bundle, or file. The exception is user-contributed matrices present in projects() records; these must be accessed using the "lol" format to extract specific paths as a standard "tibble".

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#### projects

#### Value

When as = "tibble" or as = "tibble\_expanded", a tibble with each row representing an HCA object (project, sample, bundle, or file, depending on the function invoked), and columns summarizing the object. "tibble\_expanded" columns contains almost all information about the object, except as noted in the Details section.

When as = "lol", a list-of-lists data structure representing detailed information on each object.

When as = "list", projects() returns an R list, typically containing other lists or atomic vectors, representing detailed information on each project.

projects\_facets() invoked with no facet= argument returns a tibble summarizing terms available as projects() return values, and for use in filters. The tibble contains columns

- facet: the name of the facet.
- n\_terms: the number of distinct values the facet can take.
- n\_values: the number of occurrences of the facet term in the entire catalog.

projects\_facets() invoked with a scalar value for facet= returns a tibble summarizing terms
used in the facet, and the number of occurrences of the term in the entire catalog.
\*\_columns() returns a tibble with column name containing the column name used in the tibble
returned by projects(), files(), samples(), or bundles(), and path the path (see lol\_hits())
to the data in the list-of-lists by the same functions when as = "lol". When as = "character", the
return value is a named list with paths as elements and abbreviations as names.

list-of-lists containing relevant details about the project.

#### See Also

lol() and other lol\_\*() functions for working with the list-of-list data structure returned when as = "lol".

#### Examples

```
projects(filters(), size = 100)
```

projects\_facets()
projects\_facets("genusSpecies")

```
projects_default_columns()
```

project <- projects(size = 1, as = "list")
project\_uuid <- project[["hits"]][[1]][["entryId"]]
projects\_detail(uuid = project\_uuid)</pre>

samples

# Description

samples() takes a list of user provided project titles to be used to query the HCA API for information about available samples.

samples\_detail() takes a unique sample\_id and catalog for the sample, and returns details about the specified sample as a list-of-lists

# Usage

```
samples(
    filters = NULL,
    size = 1000L,
    sort = "projectTitle",
    order = c("asc", "desc"),
    catalog = NULL,
    as = c("tibble", "lol", "list", "tibble_expanded"),
    columns = samples_default_columns("character")
)
samples_facets(facet = character(), catalog = NULL)
samples_default_columns(as = c("tibble", "character"))
samples_detail(uuid, catalog = NULL)
```

# Arguments

filters	filter object created by filters(), or NULL (default; all projects).
size	integer(1) maximum number of results to return; default: all projects matching filter. The default (10000) is meant to be large enough to return all results.
sort	<pre>character(1) project facet (see facet_options()) to sort result; default: "projectTitle".</pre>
order	character(1) sort order. One of "asc" (ascending) or "desc" (descending).
catalog	character(1) source of data. Use catalogs() for possible values.
as	character(1) return format. One of "tibble" (default), "lol", "list", or "tibble_expanded", as described in the Details and Value sections of ?projects.
columns	named character() indicating the paths to be used for parsing the 'lol' returned from the HCA to a tibble. The names of columns are used as column names in the returned tibble. If the columns are unnamed, a name is derived from the ele- ments of path by removing hits[*] and all [*], e.g., a path hits[*].donorOrganisms[*].biological is given the name donorOrganisms.biologicalSex.

#### summary

facet	character() of valid facet names. Summary results (see 'Value', below) are re- turned when missing or length greater than 1; details are returned when a single facet is specified.
uuid	character() unique identifier (e.g., projectId) of the object.

# Value

samples\_detail() returns a list-of-lists containing relevant details about the sample

# Examples

```
title <- paste(
    "Tabula Muris: Transcriptomic characterization of 20 organs and",
    "tissues from Mus musculus at single cell resolution"
)
filters <- filters( projectTitle = list(is = title) )
samples(filters = filters)
samples_facets()
sample <- samples(size = 1, as = "list")
sample_uuid <- sample[["hits"]][[1]][["entryId"]]
samples_detail(uuid = sample_uuid)</pre>
```

summary Repository summary statistics	
---------------------------------------	--

# Description

summary() provides numerical summaries of catalog content

#### Usage

```
summary(
  filters = NULL,
  type = c("overview", "fileTypeSummaries", "cellCountSummaries", "organTypes", "list"),
  catalog = NULL
)
```

#### Arguments

filters	filter object created by filters(), or NULL (default; all projects).
type	character(1) type of summary to return. Possible values include "overview", "fileTypeSummaries", "cellCountSummaries", "organType", and a "list" off all summary statistics.
catalog	character(1) source of data. Use catalogs() for possible values.

# Value

summary() returns a tibble or (for type = "list") a list-of-lists of summary statistics.

```
summary()
filter <- filters(
    organ = list(is = c("brain", "heart")),
    genusSpecies = list(is = "Homo sapiens")
)
summary(filter)
summary(filter, "fileTypeSummaries")
summary(filter, "cellCountSummaries")</pre>
```

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