Package 'cBioPortalData'

June 19, 2022

Title Exposes and makes available data from the cBioPortal web resources

Version 2.8.2

Description The cBioPortalData R package accesses study datasets from the cBio Cancer Genomics Portal. It accesses the data either from the pre-packaged zip / tar files or from the API interface that was recently implemented by the cBioPortal Data Team. The package can provide data in either tabular format or with MultiAssayExperiment object that uses familiar Bioconductor data representations.

Depends R (>= 4.2.0), AnVIL (>= 1.7.1), MultiAssayExperiment

Imports BiocFileCache (>= 1.5.3), digest, dplyr, GenomeInfoDb, GenomicRanges, httr, IRanges, methods, readr, RaggedExperiment, RTCGAToolbox (>= 2.19.7), S4Vectors, SummarizedExperiment, stats, tibble, tidyr, TCGAutils (>= 1.9.4), utils

Suggests BiocStyle, knitr, survival, survminer, rmarkdown, testthat

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BugReports https://github.com/waldronlab/cBioPortalData/issues

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cBioCache

Manage cache / download directories for study data

Description

Managing data downloads is important to save disk space and re-downloading data files. This can be done effortlessly via the integrated BiocFileCache system.

Usage

```
cBioCache(...)
setCache(
  directory = tools::R_user_dir("cBioPortalData", "cache"),
  verbose = TRUE,
  ask = interactive()
)
removePackCache(cancer_study_id, dry.run = TRUE)
removeDataCache(
  api,
  studyId = NA_character_,
  genePanelId = NA_character_,
  genes = NA_character_,
 molecularProfileIds = NULL,
  sampleListId = NULL,
  sampleIds = NULL,
  by = c("entrezGeneId", "hugoGeneSymbol"),
  dry.run = TRUE,
)
```

Arguments

... For cBioCache, arguments passed to setCache

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directory The file location where the cache is located. Once set future downloads will go

to this folder.

verbose Whether to print descriptive messages

ask logical (default TRUE when interactive session) Confirm the file location of the

cache directory

cancer_study_id

character(1) The studyId from getStudies

dry.run logical Whether or not to remove cache files (default TRUE).

api An API object of class 'cBioPortal' from the 'cBioPortal' function

studyId character(1) Indicates the "studyId" as taken from 'getStudies'

genePanelId character(1) Identifies the gene panel, as obtained from the 'genePanels' func-

tion

genes character() Either Entrez gene identifiers or Hugo gene symbols. When in-

cluded, the 'by' argument indicates the type of identifier provided and 'genePan-

elId' is ignored. Preference is given to Entrez IDs due to faster query responses.

molecularProfileIds

character() A vector of molecular profile IDs

sampleListId character(1) A sample list identifier as obtained from 'sampleLists()"

sampleIds character() Sample identifiers

by character(1) Either 'entrezGeneId' or 'hugoGeneSymbol' for row metadata (de-

fault: 'entrezGeneId')

Value

cBioCache: The path to the cache location

cBioCache

Get the directory location of the cache. It will prompt the user to create a cache if not already created. A specific directory can be used via setCache.

setCache

Specify the directory location of the data cache. By default, it will go to the user directory as given by:

```
tools::R_user_dir("cBioPortalData", "cache")
```

removePackCache

Some files may become corrupt when downloading, this function allows the user to delete the tarball associated with a cancer_study_id in the cache. This only works for the cBioDataPack function. To remove the entire cBioPortalData cache, run unlink("~/.cache/cBioPortalData").

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Examples

```
cBioCache()
removePackCache("acc_tcga", dry.run = TRUE)
cbio <- cBioPortal()</pre>
cBioPortalData(
    cbio, by = "hugoGeneSymbol",
    studyId = "acc_tcga",
    genePanelId = "AmpliSeq",
   molecularProfileIds =
        c("acc_tcga_rppa", "acc_tcga_linear_CNA", "acc_tcga_mutations")
)
removeDataCache(
    cbio, by = "hugoGeneSymbol",
    studyId = "acc_tcga",
    genePanelId = "AmpliSeq",
   molecularProfileIds =
        c("acc_tcga_rppa", "acc_tcga_linear_CNA", "acc_tcga_mutations"),
    dry.run = TRUE
)
```

cBioDataPack

Obtain pre-packaged data from cBioPortal and represent as a Multi-AssayExperiment object

Description

The cBioDataPack function allows the user to download and process cancer study datasets found in MSKCC's cBioPortal. Output datasets use the MultiAssayExperiment data representation to faciliate analysis and data management operations.

Usage

```
cBioDataPack(
  cancer_study_id,
  use_cache = TRUE,
  names.field = c("Hugo_Symbol", "Entrez_Gene_Id", "Gene"),
  cleanup = TRUE,
  ask = interactive()
)
```

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Arguments

cancer_study_id

character(1) The study identifier from cBioPortal as in https://cbioportal.

org/webAPI

use_cache logical(1) (default TRUE) create the default cache location and use it to track

downloaded data. If data found in the cache, data will not be re-downloaded. A

path can also be provided to data cache location.

names.field character() Possible column names for the column that will used to label ranges

from data such as mutations or copy number (default: c("Hugo_Symbol", "Entrez_Gene_Id",

"Gene")). Values are cycled through and eliminated when no data present, or duplicates are found. Values in the corresponding column must be unique in

each row.

cleanup logical(1) whether to delete the untar-red contents from the exdir folder (de-

fault TRUE)

ask logical(1) Whether to prompt the the user before downloading and loading study

MultiAssayExperiment. Set to interactive() by default; the user will be

prompted to continue for studies that are not currently building as MultiAssayExperiment

based on previous testing (in a non-interactive session, data download will be at-

tempted; equivalent to ask = FALSE)

Details

The full list of study identifiers (studyIds) can obtained from getStudies(). Currently, only ~ 72% of datasets can be represented as MultiAssayExperiment data objects from the data tarballs. Refer to getStudies(..., buildReport = TRUE) and its "pack_build" column to see which study identifiers are not building. Users who would like to prioritize particular datasets should open GitHub issues at the URL in the DESCRIPTION file. For a more fine-grained approach to downloading data from the cBioPortal API, refer to the cBioPortalData function.

Value

A MultiAssayExperiment object

cBio_URL

The cBioDataPack function accesses data from the cBio_URL option. By default, it points to an Amazon S3 bucket location. Previously, it pointed to 'http://download.cbioportal.org'. This recent change (> 2.1.17) should provide faster and more reliable downloads for all users. See the URL using cBioPortalData:::.url_location. This can be changed if there are mirrors that host this data by setting the cBio_URL option with getOption("cBio_URL", "https://some.url.com/") before running the function.

Author(s)

Levi Waldron, Marcel R., Ino dB.

See Also

https://www.cbioportal.org/datasets, cBioPortalData, removePackCache

Examples

```
cbio <- cBioPortal()
head(getStudies(cbio)[["studyId"]])
mae <- cBioDataPack("acc_tcga")</pre>
```

cBioPortal

The R interface to the cBioPortal API Data Service

Description

This section of the documentation lists the functions that allow users to access the cBioPortal API. The main representation of the API can be obtained from the 'cBioPortal' function. The supporting functions listed here give access to specific parts of the API and allow the user to explore the API with individual calls. Many of the functions here are listed for documentation purposes and are recommended for advanced usage only. Users should only need to use the 'cBioPortalData' main function to obtain data.

Usage

```
cBioPortal(
  hostname = "www.cbioportal.org",
  protocol = "https",
  api. = "/api/api-docs",
  token = character()
)
getStudies(api, buildReport = FALSE)
clinicalData(api, studyId = NA_character_)
molecularProfiles(
  api,
  studyId = NA_character_,
 projection = c("SUMMARY", "ID", "DETAILED", "META")
mutationData(
  api,
 molecularProfileIds = NA_character_,
  entrezGeneIds = NULL,
  sampleIds = NULL
)
molecularData(
```

```
molecularProfileIds = NA_character_,
  entrezGeneIds = NULL,
  sampleIds = NULL
)
searchOps(api, keyword)
samplesInSampleLists(api, sampleListIds = NA_character_)
sampleLists(api, studyId = NA_character_)
allSamples(api, studyId = NA_character_)
getSampleInfo(
  api,
  studyId = NA_character_,
  sampleListIds = NULL,
  projection = c("SUMMARY", "ID", "DETAILED", "META")
genePanels(api)
getGenePanel(api, genePanelId = NA_character_)
genePanelMolecular(
  api,
 molecularProfileId = NA_character_,
 sampleListId = NULL,
  sampleIds = NULL
)
getGenePanelMolecular(api, molecularProfileIds = NA_character_, sampleIds)
geneTable(api, pageSize = 1000, pageNumber = 0, ...)
queryGeneTable(
  api,
  by = c("entrezGeneId", "hugoGeneSymbol"),
  genes = NA_character_,
  genePanelId = NA_character_
)
getDataByGenes(
  api,
  studyId = NA_character_,
  genes = NA_character_,
  genePanelId = NA_character_,
```

```
by = c("entrezGeneId", "hugoGeneSymbol"),
molecularProfileIds = NULL,
sampleListId = NULL,
sampleIds = NULL,
...
)
```

Arguments

hostname character(1) The internet location of the service (default: 'www.cbioportal.org')
protocol character(1) The internet protocol used to access the hostname (default: 'https')
api. character(1) The directory location of the API protocol within the hostname

(default: '/api/api-docs')

token character(1) The Authorization Bearer token e.g., "63eba81c-2591-4e15-9d1c-

fb6e8e51e35d" or a path to text file.

api An API object of class 'cBioPortal' from the 'cBioPortal' function

buildReport logical(1) Indicates whether to append the build information to the 'getStudies()'

table (default FALSE)

studyId character(1) Indicates the "studyId" as taken from 'getStudies'

projection character(default: "SUMMARY") Specify the projection type for data retrieval

for details see API documentation

molecularProfileIds

character() A vector of molecular profile IDs

entrezGeneIds numeric() A vector indicating entrez gene IDs

sampleIds character() Sample identifiers

keyword character(1) Keyword or pattern for searching through available operations sampleListIds character() A vector of 'sampleListId' as obtained from 'sampleLists'

genePanelId character(1) Identifies the gene panel, as obtained from the 'genePanels' func-

tion

molecularProfileId

character(1) Indicates a molecular profile ID

sampleListId character(1) A sample list identifier as obtained from 'sampleLists()"

pageSize numeric(1) The number of rows in the table to return

pageNumber numeric(1) The pagination page number

... Additional arguments to lower level API functions

by character(1) Either 'entrezGeneId' or 'hugoGeneSymbol' for row metadata (de-

fault: 'entrezGeneId')

genes character() Either Entrez gene identifiers or Hugo gene symbols. When in-

cluded, the 'by' argument indicates the type of identifier provided and 'genePanelId' is ignored. Preference is given to Entrez IDs due to faster query responses.

Value

cBioPortal: An API object of class 'cBioPortal'

cBioPortalData: A data object of class 'MultiAssayExperiment'

API Metadata

* getStudies - Obtain a table of studies and associated metadata and optionally include a 'buildReport' status (default FALSE) for each study. When enabled, the 'api_build' and 'pack_build' columns will be added to the table and will show if 'MultiAssayExperiment' objects can be generated for that particular study identifier ('studyId'). The 'api_build' column corresponds to datasets obtained with 'cBioPortalData' and the 'pack_build' column corresponds to datsets loaded via 'cBioDataPack'.

- * searchOps Search through API operations with a keyword
- * sampleLists obtain all 'sampleListIds' for a particular 'studyId'
- * allSamples obtain all samples within a particular 'studyId'
- * genePanels Show all available gene panels
- * geneTable Get a table of all genes by 'entrezGeneId' and 'hugoGeneSymbol'
- * queryGeneTable Get a table for only the 'genes' or 'genePanelId' of interest. Gene inputs are identified with the 'by' argument

Patient Data

* clinicalData - Obtain clinical data for a particular study identifier ('studyId')

Molecular Profiles

- * molecularProfiles Produce a molecular profiles dataset for a given study identifier ('studyId')
- * molecularData Produce a dataset of molecular profile data based on 'molecularProfileId', 'entrezGeneIds', and 'sampleIds'

Mutation Data

* mutationData - Produce a dataset of mutation data using 'molecularProfileId', 'entrezGeneIds', and 'sampleIds'

Sample Data

- * samplesInSampleLists get all samples associated with a 'sampleListId'
- * getSampleInfo Obtain sample metadata for a particular 'studyId' or 'sampleListId'

Gene Panels

- * getGenePanels Obtain the gene panel for a particular 'genePanelId'
- * genePanelMolecular get gene panel data for a particular 'molecularProfileId' and either a vector of 'sampleListId' or 'sampleId'
- * getGenePanelMolecular get gene panel data for multiple 'molecularProfileId's and a vector of 'sampleIds'

Genes

* getDataByGenes - Download data for a number of genes within 'molecularProfileId' indicators, optionally a 'sampleListId' can be provided.

Examples

```
cbio <- cBioPortal()</pre>
getStudies(api = cbio)
searchOps(api = cbio, keyword = "molecular")
## obtain clinical data
acc_clin <- clinicalData(api = cbio, studyId = "acc_tcga")</pre>
acc_clin
molecularProfiles(api = cbio, studyId = "acc_tcga")
genePanels(cbio)
(gp <- getGenePanel(cbio, "AmpliSeq"))</pre>
muts <- mutationData(</pre>
    api = cbio,
    molecularProfileIds = "acc_tcga_mutations",
    entrezGeneIds = 1:1000,
    sampleIds = c("TCGA-OR-A5J1-01", "TCGA-OR-A5J2-01")
exps <- molecularData(</pre>
    api = cbio,
    molecularProfileIds = c("acc_tcga_rna_seq_v2_mrna", "acc_tcga_rppa"),
    entrezGeneIds = 1:1000,
    sampleIds = c("TCGA-OR-A5J1-01", "TCGA-OR-A5J2-01")
)
sampleLists(api = cbio, studyId = "acc_tcga")
samplesInSampleLists(
    api = cbio,
    sampleListIds = c("acc_tcga_rppa", "acc_tcga_cnaseq")
genePanels(api = cbio)
getGenePanel(api = cbio, genePanelId = "IMPACT341")
queryGeneTable(api = cbio, by = "entrezGeneId", genes = 7157)
getDataByGenes(
    cbio, studyId = "acc_tcga", genes = 1:3,
    by = c("entrezGeneId", "hugoGeneSymbol"),
    molecularProfileId = "acc_tcga_rppa",
    sampleListId = "acc_tcga_rppa"
)
```

cBioPortal-class

cBioPortal-class

A class for representing the cBioPortal API protocol

Description

The cBioPortal class is a representation of the cBioPortal API protocol that directly inherits from the Service class in the AnVIL package. For more information, see the AnVIL package.

Usage

```
## S4 method for signature 'cBioPortal'
operations(x, ..., .deprecated = FALSE)
```

Arguments

x A Service instance or API representation as given by the cBioPortal function.

... additional arguments passed to methods or, for 'operations, Service-method', to

the internal 'get_operation()' function.

. deprecated optional logical(1) include deprecated operations?

Details

This class takes the static API as provided at https://www.cbioportal.org/api/api-docs and creates an R object with the help from underlying infrastructure (i.e., rapiclient and AnVIL) to give the user a unified representation of the API specification provided by the cBioPortal group. Users are not expected to interact with this class other than to use it as input to the functionality provided by the rest of the package.

Functions

• operations,cBioPortal-method:

See Also

```
cBioPortal, Service
```

Examples

```
cBioPortal()
```

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cBioPortalData

Download data from the cBioPortal API

Description

Obtain a MultiAssayExperiment object for a particular gene panel, studyId, molecularProfileIds, and sampleListIds combination. Default molecularProfileIds and sampleListIds are set to NULL for including all data. This option is best for users who wish to obtain a section of the study data that pertains to a specific molecular profile and gene panel combination. For users looking to download the entire study data as provided by the https://cbioportal.org/datasets, refer to cBioDataPack.

Usage

```
cBioPortalData(
    api,
    studyId = NA_character_,
    genePanelId = NA_character_,
    genes = NA_character_,
    molecularProfileIds = NULL,
    sampleListId = NULL,
    sampleIds = NULL,
    by = c("entrezGeneId", "hugoGeneSymbol"),
    check_build = TRUE
)
```

Arguments

api An API object of class 'cBioPortal' from the 'cBioPortal' function

studyId character(1) Indicates the "studyId" as taken from 'getStudies'

genePanelId character(1) Identifies the gene panel, as obtained from the 'genePanels' func-

tion

genes character() Either Entrez gene identifiers or Hugo gene symbols. When in-

cluded, the 'by' argument indicates the type of identifier provided and 'genePanelId' is ignored. Preference is given to Entrez IDs due to faster query responses.

molecularProfileIds

character() A vector of molecular profile IDs

sampleListId character(1) A sample list identifier as obtained from 'sampleLists()"

sampleIds character() Sample identifiers

by character(1) Either 'entrezGeneId' or 'hugoGeneSymbol' for row metadata (de-

fault: 'entrezGeneId')

check_build logical(1L) Whether to check the build status of the studyId using an internal

dataset. This argument should be set to FALSE if using alternative hostnames,

e.g., 'pedcbioportal.kidsfirstdrc.org'

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Details

We are able to succesfully represent 98 percent of the study identifiers as MultiAssayExperiment objects as obtained via cBioPortalData with the IMPACT341 genePanelId as the example gene panel. Datasets that currently fail to import can be seen in the getStudies(..., buildReport = TRUE) dataset under the "api_build" column. Note that changes to the cBioPortal API may affect this rate at any time. If you encounter any issues, please open a GitHub issue at the https://github.com/waldronlab/cBioPortalData/issues/ page with a fully reproducible example.

Value

A MultiAssayExperiment object

See Also

cBioDataPack, removeDataCache

Examples

```
cbio <- cBioPortal()

samps <- samplesInSampleLists(cbio, "acc_tcga_rppa")[[1]]

getGenePanelMolecular(
    cbio, molecularProfileIds = c("acc_tcga_rppa", "acc_tcga_linear_CNA"),
    samps
)

acc_tcga <- cBioPortalData(
    cbio, by = "hugoGeneSymbol",
    studyId = "acc_tcga",
    genePanelId = "AmpliSeq",
    molecularProfileIds =
        c("acc_tcga_rppa", "acc_tcga_linear_CNA", "acc_tcga_mutations")
)</pre>
```

downloadStudy

Manually download, untar, and load study tarballs

Description

Note that these functions should be used when a particular study is *not* currently available as a MultiAssayExperiment representation. Otherwise, use cBioDataPack. Provide a cancer_study_id from getStudies and retrieve the study tarball from the cBio Genomics Portal. These functions are used by cBioDataPack under the hood to download,untar, and load the tarball datasets with caching. As stated in ?cBioDataPack, not all studies are currently working as MultiAssayExperiment objects. As of July 2020, about ~80% of datasets can be successfully imported into the MultiAssayExperiment data class. Please open an issue if you would like the team to prioritize a study. You may also check getStudies(buildReport = TRUE)\$pack_build for the current status.

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Usage

```
downloadStudy(
  cancer_study_id,
  use_cache = TRUE,
  force = FALSE,
  url_location = getOption("cBio_URL", .url_location)
)

untarStudy(cancer_study_file, exdir = tempdir())

loadStudy(
  filepath,
  names.field = c("Hugo_Symbol", "Entrez_Gene_Id", "Gene"),
  cleanup = TRUE
)
```

Arguments

cancer_study_id

character(1) The study identifier from cBioPortal as in https://cbioportal.

org/webAPI

use_cache logical(1) (default TRUE) create the default cache location and use it to track

downloaded data. If data found in the cache, data will not be re-downloaded. A

path can also be provided to data cache location.

force logical(1) (default FALSE) whether to force re-download data from remote lo-

cation

url_location character(1) (default "https://cbioportal-datahub.s3.amazonaws.com") the URL

location for downloading packaged data. Can be set using the 'cBio_URL' op-

tion (see ?cBioDataPack for more details)

cancer_study_file

character(1) indicates the on-disk location of the downloaded tarball

exdir character(1) indicates the folder location to *put* the contents of the tarball (de-

fault tempdir(); see also ?untar)

filepath character(1) indicates the folder location where the contents of the tarball are

located (usually the same as exdir)

names.field character() Possible column names for the column that will used to label ranges

from data such as mutations or copy number (default: c("Hugo_Symbol", "Entrez_Gene_Id",

"Gene")). Values are cycled through and eliminated when no data present, or duplicates are found. Values in the corresponding column must be unique in

each row.

cleanup logical(1) whether to delete the untar-red contents from the exdir folder (de-

fault TRUE)

Details

When attempting to load a dataset using loadStudy, note that the cleanup argument is set to TRUE by default. Change the argument to FALSE if you would like to keep the untarred data in the exdir

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location. downloadStudy and untarStudy are not affected by this change. The tarball of the downloaded data is cached via BiocFileCache when use_cache is TRUE.

Value

- downloadStudy The file location of the data tarball
- untarStudy The directory location of the contents
- loadStudy A MultiAssayExperiment-class object

See Also

cBioDataPack, MultiAssayExperiment

Examples

```
(acc_file <- downloadStudy("acc_tcga"))
(file_dir <- untarStudy(acc_file, tempdir()))
loadStudy(file_dir)</pre>
```

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