Package 'cbioportalR'

October 5, 2023

Title Browse and Query Clinical and Genomic Data from cBioPortal **Version** 1.1.0

Description Provides R users with direct access to genomic and clinical data from the 'cBioPortal' web resource via user-friendly

functions that wrap 'cBioPortal's' existing API endpoints https://www.cbioportal.org/api/swagger-ui/index.html. Users can browse and query genomic data on mutations, copy number alterations and fusions, as well as data on tumor mutational burden ('TMB'), microsatellite instability status ('MSI'), 'FACETS' and select clinical data points (depending on the study).

See <https:

//www.cbioportal.org/> and Gao et al., (2013) <doi:10.1126/scisignal.2004088> for more information on the cBioPortal web resource.

```
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 $\pmb{BugReports} \ \text{https://github.com/karissawhiting/cbioportalR/issues}$

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```
available_clinical_attributes
```

Get all available clinical attribute IDs for a study

Description

Get all available clinical attribute IDs for a study

Usage

```
available_clinical_attributes(study_id = NULL, base_url = NULL)
```

Arguments

study_id cbioportal study ID

base_url The database URL to query. If NULL will default to URL set with set_cbioportal_db(<your_db>)

Value

a data frame of available clinical attributes for that study

Examples

```
## Not run:
available_clinical_attributes("acc_tcga", base_url = 'www.cbioportal.org/api')
## End(Not run)
```

 $available_gene_panels \quad \textit{Get Available Gene Panels For a Database}$

Description

Get Available Gene Panels For a Database

Usage

```
available_gene_panels(base_url = NULL)
```

Arguments

base_url

The database URL to query. If NULL will default to URL set with set_cbioportal_db(<your_db>)

Value

a dataframe of metadata regarding each available panel

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Examples

```
## Not run:
set_cbioportal_db("public")
available_gene_panels()
## End(Not run)
```

available_patients

Get All Patient IDs in a Study

Description

Get All Patient IDs in a Study

Usage

```
available_patients(study_id = NULL, base_url = NULL)
```

Arguments

study_id A character string indicating which study ID should be searched. Only 1 study

ID allowed.

base_url The database URL to query If NULL will default to URL set with set_cbioportal_db(<your_db>)

Value

A dataframe of patient_ids in a given study

Examples

```
## Not run:
set_cbioportal_db("public")
available_samples(study_id = "acc_tcga")
## End(Not run)
```

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available_profiles

Get Available Genomic Profiles For a Study or Database

Description

Get Available Genomic Profiles For a Study or Database

Usage

```
available_profiles(study_id = NULL, base_url = NULL)
```

Arguments

study_id A character vector of length 1 indicating study_id. See get_studies() to see

all available studies for your URL. If NULL, it will return all profiles for your

current database url

base_url The database URL to query. If NULL will default to URL set with set_cbioportal_db(<your_db>)

Value

A dataframe of available genetic profiles and their profile ids

Examples

```
## Not run:
set_cbioportal_db("public")
available_profiles()
available_profiles(study_id = "acc_tcga")
## End(Not run)
```

available_samples

Get All Sample IDs in a Study

Description

Pulls all available sample IDs for a given study ID or sample list ID. Either a study ID or sample list ID must be passed. If both sample_list and study_id are not NULL, sample_list ID will be searched and study_id will be ignored.

Usage

```
available_samples(study_id = NULL, sample_list_id = NULL, base_url = NULL)
```

Arguments

study_id A character string indicating which study ID should be searched. Only 1 study

ID allowed.

sample_list_id A character string indicating which sample list ID should be searched. Only 1

sample list ID allowed.

base_url The database URL to query If NULL will default to URL set with set_cbioportal_db(<your_db>)

Value

A dataframe of sample_ids in a given study

Examples

```
## Not run:
set_cbioportal_db("public")
available_samples(study_id = "acc_tcga")
available_samples(sample_list_id = "acc_tcga_cna")
## End(Not run)
```

```
available_sample_lists
```

Get All Sample Lists Available For a Study

Description

Get All Sample Lists Available For a Study

Usage

```
available_sample_lists(study_id = NULL, base_url = NULL)
```

Arguments

study_id A character string indicating which study ID should be searched. Only 1 study

ID allowed.

base_url The database URL to query If NULL will default to URL set with set_cbioportal_db(<your_db>)

Value

A dataframe of patient_ids in a given study

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Examples

```
## Not run:
set_cbioportal_db("public")
available_sample_lists(study_id = "acc_tcga")
## End(Not run)
```

available_studies

Get Metadata on All Available Studies in a Database

Description

Get Metadata on All Available Studies in a Database

Usage

```
available_studies(base_url = NULL)
```

Arguments

base_url

The database URL to query. If NULL will default to URL set with set_cbioportal_db(<your_db>)

Value

A dataframe of available studies and their metadata

Examples

```
## Not run:
set_cbioportal_db("public")
available_studies()
## End(Not run)
```

get_alias

Get Gene Name Alias for a Given Hugo Symbol

Description

This function grabs known gene aliases for a given Hugo Symbol. You may notice that genes - alias pairs are not always consistent. For example get_alias("KMT2D") will return "MLL2" but get_alias("MLL2") will not return "KMT2D" This function relies on the existing cBioPortal API which controls this database of aliases. Therefore, this is a convenience function but you may want to consider a more carefully curated alias list like cbioportalR::impact_gene_info

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Usage

```
get_alias(hugo_symbol = NULL, base_url = NULL)
```

Arguments

hugo_symbol a hugo symbol for which to return aliases base_url The database URL to query

Value

A character string with all aliases

Examples

```
## Not run:
get_alias(hugo_symbol = "FGFR3", base_url = 'www.cbioportal.org/api')
get_alias(hugo_symbol = c("FGFR3", "TP53"), base_url = 'www.cbioportal.org/api')
## End(Not run)
```

```
get_cbioportal_token Get cBioPortal Access Token
```

Description

Convenience function that retrieves cBioPortal token System Environment variable "CBIOPORTAL_TOKEN"

Usage

```
get_cbioportal_token()
```

Value

Returns a string with cBioPortal token if successfully authenticated, or a warning that token could not be found.

Author(s)

Karissa Whiting, Daniel D. Sjoberg

Examples

```
## Not run:
get_cbioportal_token()
## End(Not run)
```

get_clinical_by_patient

```
get_clinical_by_patient
```

Get clinical data by attribute, study ID and patient ID

Description

Get clinical data by attribute, study ID and patient ID

Usage

```
get_clinical_by_patient(
  study_id = NULL,
  patient_id = NULL,
  patient_study_pairs = NULL,
  clinical_attribute = NULL,
  base_url = NULL
)
```

Arguments

study_id

A string indicating the study ID from which to pull data. If no study ID, will guess the study ID based on your URL and inform. Only 1 study ID can be passed. If mutations/cna from more than 1 study needed, see sample_study_pairs

patient_id a cBioPortal patient_id
patient_study_pairs

A dataframe with columns: patient_id, study_id. Variations in capitalization of column names are accepted. This can be used in place of patient_id, study_id, arguments above if you need to pull samples from several different studies at once. If passed, this will take overwrite patient_id and study_id if they are also passed.

clinical_attribute

one or more clinical attributes for your study. If none provided, will return all attributes available for studies

base_url

The database URL to query If NULL will default to URL set with $set_cbioportal_db(<your_db>)$

Value

a dataframe of a specific clinical attribute

Examples

```
"P-0003238", "blca_nmibc_2017",
"P-0000004", "msk_impact_2017",
"P-0000023", "msk_impact_2017")

x <- get_clinical_by_patient(patient_study_pairs = ex,
    clinical_attribute = NULL, base_url = 'www.cbioportal.org/api')
## End(Not run)</pre>
```

get_clinical_by_sample

Get clinical data by attribute, study ID and sample ID

Description

Get clinical data by attribute, study ID and sample ID

Usage

```
get_clinical_by_sample(
  study_id = NULL,
  sample_id = NULL,
  sample_study_pairs = NULL,
  clinical_attribute = NULL,
  base_url = NULL
)
```

Arguments

study_id

A string indicating the study ID from which to pull data. If no study ID, will guess the study ID based on your URL and inform. Only 1 study ID can be passed. If mutations/cna from more than 1 study needed, see sample_study_pairs

sample_id

a vector of sample IDs (character)

sample_study_pairs

A dataframe with columns: sample_id, study_id and molecular_profile_id (optional). Variations in capitalization of column names are accepted. This can be used in place of sample_id, study_id, molecular_profile_id arguments above if you need to pull samples from several different studies at once. If passed this will take overwrite sample_id, study_id, molecular_profile_id if also passed.

clinical_attribute

one or more clinical attributes for your study. If none provided, will return all attributes available for studies

base_url

The database URL to query If NULL will default to URL set with set_cbioportal_db(<your_db>)

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Value

a dataframe of a specific clinical attribute

Examples

```
## Not run:
get_clinical_by_sample(study_id = "acc_tcga", sample_id = "TCGA-OR-A5J2-01",
    clinical_attribute = "CANCER_TYPE", base_url = 'www.cbioportal.org/api')

ex <- tibble::tribble(
    ~sample_id, ~study_id,
    "P-0001453-T01-IM3", "blca_nmibc_2017",
    "P-0002166-T01-IM3", "blca_nmibc_2017",
    "P-0003238-T01-IM5", "blca_nmibc_2017",
    "P-0000004-T01-IM3", "msk_impact_2017",
    "P-00000023-T01-IM3", "msk_impact_2017")

x <- get_clinical_by_sample(sample_study_pairs = ex,
    clinical_attribute = NULL, base_url = 'www.cbioportal.org/api')

## End(Not run)</pre>
```

get_clinical_by_study Get all available clinical data for a specified study

Description

Returns all sample-level and patient-level clinical data for a given study

Usage

```
get_clinical_by_study(
   study_id = NULL,
   clinical_attribute = NULL,
   base_url = NULL
)
```

Arguments

Value

a dataframe of all available clinical attributes and their values

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Examples

```
## Not run:
get_clinical_by_study(study_id = "acc_tcga",
    clinical_attribute = "CANCER_TYPE", base_url = 'www.cbioportal.org/api')
get_clinical_by_study(study_id = "acc_tcga", base_url = 'www.cbioportal.org/api')
## End(Not run)
```

get_cna_by_sample

Get CNA By Sample ID

Description

```
Get CNA By Sample ID
```

Usage

```
get_cna_by_sample(
   sample_id = NULL,
   study_id = NULL,
   molecular_profile_id = NULL,
   sample_study_pairs = NULL,
   genes = NULL,
   panel = NULL,
   add_hugo = TRUE,
   base_url = NULL
)
```

Arguments

sample_id

a vector of sample IDs (character)

study_id

A string indicating the study ID from which to pull data. If no study ID, will guess the study ID based on your URL and inform. Only 1 study ID can be passed. If mutations/cna from more than 1 study needed, see sample_study_pairs

molecular_profile_id

A string indicating the molecular profile ID from which to pull data. If ID supplied, will guess the molecular profile ID based on the study ID. Only 1 molecular profile ID can be passed. If mutations from more than 1 study needed, see sample_study_pairs

sample_study_pairs

A dataframe with columns: sample_id, study_id and molecular_profile_id (optional). Variations in capitalization of column names are accepted. This can be used in place of sample_id, study_id, molecular_profile_id arguments above if you need to pull samples from several different studies at once. If passed this will take overwrite sample_id, study_id, molecular_profile_id if also passed.

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genes A vector of Entrez ids or Hugo symbols. If Hugo symbols are supplied, they will

be converted to entrez ids using the get_entrez_id() function. If panel and genes are both supplied, genes from both arguments will be returned. If both are NULL (default), it will return gene results for all available genomic data for

that sample.

panel One or more panel IDs to query (e.g. 'IMPACT468'). If panel and genes are

both supplied, genes from both arguments will be returned. If both are NULL (default), it will return gene results for all available genomic data for that sample.

add_hugo Logical indicating whether HugoGeneSymbol should be added to your resulting

data frame, if not already present in raw API results. Argument is TRUE by default. If FALSE, results will be returned as is (i.e. any existing Hugo Symbol

columns in raw results will not be removed).

base_url The database URL to query If NULL will default to URL set with set_cbioportal_db(<your_db>)

Value

A data frame of CNAs

Examples

get_cna_by_study

Get CNA By Study

Description

Get CNA By Study

Usage

```
get_cna_by_study(
  study_id = NULL,
  molecular_profile_id = NULL,
  add_hugo = TRUE,
  base_url = NULL
)
```

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Arguments

study_id A study ID to query mutations. If NULL, guesses study ID based on molecu-

lar_profile_id.

molecular_profile_id

a molecular profile to query mutations. If NULL, guesses molecular_profile_id

based on study ID.

add_hugo Logical indicating whether HugoGeneSymbol should be added to your resulting

data frame, if not already present in raw API results. Argument is TRUE by default. If FALSE, results will be returned as is (i.e. any existing Hugo Symbol

columns in raw results will not be removed).

base_url The database URL to query If NULL will default to URL set with set_cbioportal_db(<your_db>)

Value

A dataframe of CNAs

Examples

```
## Not run:
get_cna_by_study(study_id = "prad_msk_2019")
get_cna_by_study(molecular_profile_id = "prad_msk_2019_cna")
## End(Not run)
```

get_entrez_id

Get Entrez Gene ID for a given set of Hugo Symbols

Description

Get Entrez Gene ID for a given set of Hugo Symbols

Usage

```
get_entrez_id(hugo_symbol = NULL, base_url = NULL)
```

Arguments

hugo_symbol a character vector of Hugo Symbols

base_url The database URL to query

Value

A dataframe with Entrez Gene IDs and Hugo Symbols

Examples

```
## Not run:
get_entrez_id(hugo_symbol = "TAP1", base_url = 'www.cbioportal.org/api')
get_entrez_id(hugo_symbol = c("FGFR1", "TP53") , base_url = 'www.cbioportal.org/api')
## End(Not run)
```

get_fusions_by_sample Get Fusions By Sample ID

Description

Get Fusions By Sample ID

Usage

```
get_fusions_by_sample(
  sample_id = NULL,
  study_id = NULL,
 molecular_profile_id = NULL,
  sample_study_pairs = NULL,
  genes = NULL,
 panel = NULL,
 base_url = NULL
)
get_structural_variants_by_sample(
  sample_id = NULL,
  study_id = NULL,
 molecular_profile_id = NULL,
  sample_study_pairs = NULL,
  genes = NULL,
  panel = NULL,
 base_url = NULL
)
```

Arguments

sample_id a vector of sample IDs (character)

study_id A string indicating the study ID from which to pull data. If no study ID, will

guess the study ID based on your URL and inform. Only 1 study ID can be passed. If mutations/cna from more than 1 study needed, see sample_study_pairs

molecular_profile_id

A string indicating the molecular profile ID from which to pull data. If ID supplied, will guess the molecular profile ID based on the study ID. Only 1 molecular profile ID can be passed. If mutations from more than 1 study needed, see sample_study_pairs

sample_study_pairs

A dataframe with columns: sample_id, study_id and molecular_profile_id (optional). Variations in capitalization of column names are accepted. This can be used in place of sample_id, study_id, molecular_profile_id arguments above if you need to pull samples from several different studies at once. If passed this will take overwrite sample_id, study_id, molecular_profile_id if also passed.

genes

A vector of Entrez ids or Hugo symbols. If Hugo symbols are supplied, they will be converted to entrez ids using the get_entrez_id() function. If panel and genes are both supplied, genes from both arguments will be returned. If both are NULL (default), it will return gene results for all available genomic data for that sample.

panel

One or more panel IDs to query (e.g. 'IMPACT468'). If panel and genes are both supplied, genes from both arguments will be returned. If both are NULL (default), it will return gene results for all available genomic data for that sample.

base_url

The database URL to query If NULL will default to URL set with set_cbioportal_db(<your_db>)

Value

A data frame of Fusions

Examples

```
get_fusions_by_study Get Fusions By Study
```

Description

Get Fusions By Study

Usage

```
get_fusions_by_study(
  study_id = NULL,
  molecular_profile_id = NULL,
  base_url = NULL
```

```
get_genes

get_structural_variants_by_study(
    study_id = NULL,
    molecular_profile_id = NULL,
    base_url = NULL
)
```

Arguments

study_id A study ID to query mutations. If NULL, guesses study ID based on molecular_profile_id.

molecular_profile_id a molecular profile to query mutations. If NULL, guesses molecular_profile_id

based on study ID.

 $base_url \qquad \qquad The \ database \ URL \ to \ query \ If \ NULL \ will \ default \ to \ URL \ set \ with \ set_cbioportal_db(<your_db>)$

Value

A dataframe of fusions

Examples

get_genes

Get A List of Genes for a Specified Database

Description

Get A List of Genes for a Specified Database

Usage

```
get_genes(base_url = NULL)
```

Arguments

base_url

The database URL to query If NULL will default to URL set with set_cbioportal_db(<your_db>)

Value

A dataframe of gene ids, hugo symbols, and gene types

Examples

```
## Not run:
get_genes(base_url = 'www.cbioportal.org/api')
## End(Not run)
```

```
get_genetics_by_sample
```

Get All Genomic Information By Sample IDs

Description

Get All Genomic Information By Sample IDs

Usage

```
get_genetics_by_sample(
    sample_id = NULL,
    study_id = NULL,
    sample_study_pairs = NULL,
    genes = NULL,
    panel = NULL,
    add_hugo = TRUE,
    base_url = NULL,
    return_segments = FALSE
)
```

Arguments

sample_id a vector of sample IDs (character)

study_id

A string indicating the study ID from which to pull data. If no study ID, will guess the study ID based on your URL and inform. Only 1 study ID can be passed. If mutations/cna from more than 1 study needed, see sample_study_pairs

sample_study_pairs

A dataframe with columns: sample_id, study_id and molecular_profile_id (optional). Variations in capitalization of column names are accepted. This can be used in place of sample_id, study_id, molecular_profile_id arguments above if you need to pull samples from several different studies at once. If passed this will take overwrite sample_id, study_id, molecular_profile_id if also passed.

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genes A vector of Entrez ids or Hugo symbols. If Hugo symbols are supplied, they will

be converted to entrez ids using the get_entrez_id() function. If panel and genes are both supplied, genes from both arguments will be returned. If both are NULL (default), it will return gene results for all available genomic data for

that sample.

panel One or more panel IDs to query (e.g. 'IMPACT468'). If panel and genes are

both supplied, genes from both arguments will be returned. If both are NULL (default), it will return gene results for all available genomic data for that sample.

add_hugo Logical indicating whether HugoGeneSymbol should be added to your resulting

data frame, if not already present in raw API results. Argument is TRUE by default. If FALSE, results will be returned as is (i.e. any existing Hugo Symbol

columns in raw results will not be removed).

base_url The database URL to query If NULL will default to URL set with set_cbioportal_db(<your_db>)

return_segments

Default is FALSE where copy number segmentation data won't be returned in addition to the mutation, cna and structural variant data. TRUE will return any

available segmentation data with results.

Value

A list of mutations, cna and structural variants (including fusions), if available. Will also return copy number segmentation data if return_segments = TRUE.

Examples

```
## Not run:
get_genetics_by_sample(sample_id = c("TCGA-OR-A5J2-01","TCGA-OR-A5J6-01"),
    study_id = "acc_tcga",
    return_segments = TRUE)
## End(Not run)
```

get_genetics_by_study Get All Genomic Information By Study

Description

Get All Genomic Information By Study

Usage

```
get_genetics_by_study(
   study_id = NULL,
   add_hugo = TRUE,
   base_url = NULL,
   return_segments = FALSE
)
```

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Arguments

study_id A study ID to query mutations. If NULL, guesses study ID based on molecu-

lar_profile_id.

add_hugo Logical indicating whether HugoGeneSymbol should be added to your resulting

data frame, if not already present in raw API results. Argument is TRUE by default. If FALSE, results will be returned as is (i.e. any existing Hugo Symbol

columns in raw results will not be removed).

base_url The database URL to query If NULL will default to URL set with set_cbioportal_db(<your_db>)

return_segments

Default is FALSE where copy number segmentation data won't be returned in addition to the mutation, cna and structural variant data. TRUE will return any

available segmentation data with results.

Value

A list of mutations, cna and structural variants (including fusions), if available. Will also return copy number segmentation data if return_segments = TRUE.

Examples

```
## Not run:
get_genetics_by_study(study_id = "prad_msk_2019")
## End(Not run)
```

get_gene_panel

Retrieve Genes Included For a Specified Panel ID

Description

Retrieve Genes Included For a Specified Panel ID

Usage

```
get_gene_panel(panel_id = NULL, base_url = NULL)
```

Arguments

panel_id name of panel. See available_gene_panels() to get panel ID

base_url The database URL to query. If NULL will default to URL set with set_cbioportal_db(<your_db>)

Value

A dataframe of genes in a specified panel

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Examples

```
## Not run:
get_gene_panel(panel_id = "IMPACT468", base_url = 'www.cbioportal.org/api')
## End(Not run)
```

get_hugo_symbol

Get Hugo Symbol for a given set of Entrez IDs

Description

Get Hugo Symbol for a given set of Entrez IDs

Usage

```
get_hugo_symbol(entrez_id = NULL, base_url = NULL)
```

Arguments

entrez_id a character or numeric vector of Entrez gene IDs

base_url The database URL to query

Value

A dataframe with Entrez Gene IDs and Hugo Symbols

Examples

```
## Not run:
get_hugo_symbol(entrez_id = 2261, base_url = 'www.cbioportal.org/api')
get_hugo_symbol(entrez_id = c(2261, 7157) , base_url = 'www.cbioportal.org/api')
## End(Not run)
```

```
get_mutations_by_sample
```

Get Mutations By Sample ID

Description

Get Mutations By Sample ID

Usage

```
get_mutations_by_sample(
   sample_id = NULL,
   study_id = NULL,
   molecular_profile_id = NULL,
   sample_study_pairs = NULL,
   genes = NULL,
   panel = NULL,
   add_hugo = TRUE,
   base_url = NULL
)
```

Arguments

sample_id

a vector of sample IDs (character)

study_id

A string indicating the study ID from which to pull data. If no study ID, will guess the study ID based on your URL and inform. Only 1 study ID can be passed. If mutations/cna from more than 1 study needed, see sample_study_pairs

molecular_profile_id

A string indicating the molecular profile ID from which to pull data. If ID supplied, will guess the molecular profile ID based on the study ID. Only 1 molecular profile ID can be passed. If mutations from more than 1 study needed, see sample_study_pairs

sample_study_pairs

A dataframe with columns: sample_id, study_id and molecular_profile_id (optional). Variations in capitalization of column names are accepted. This can be used in place of sample_id, study_id, molecular_profile_id arguments above if you need to pull samples from several different studies at once. If passed this will take overwrite sample_id, study_id, molecular_profile_id if also passed.

genes

A vector of Entrez ids or Hugo symbols. If Hugo symbols are supplied, they will be converted to entrez ids using the get_entrez_id() function. If panel and genes are both supplied, genes from both arguments will be returned. If both are NULL (default), it will return gene results for all available genomic data for that sample.

panel

One or more panel IDs to query (e.g. 'IMPACT468'). If panel and genes are both supplied, genes from both arguments will be returned. If both are NULL (default), it will return gene results for all available genomic data for that sample.

add_hugo

base_url

Logical indicating whether HugoGeneSymbol should be added to your resulting data frame, if not already present in raw API results. Argument is TRUE by default. If FALSE, results will be returned as is (i.e. any existing Hugo Symbol columns in raw results will not be removed).

The database URL to query If NULL will default to URL set with set_cbioportal_db(<your_db>)

Value

A data frame of mutations (maf file format)

Examples

```
## Not run:
get_mutations_by_sample(sample_id = c("TCGA-OR-A5J2-01","TCGA-OR-A5J6-01"),
study_id = "acc_tcga",
base_url = "public")
## End(Not run)
```

get_mutations_by_study

Get Mutations By Study ID

Description

Get Mutations By Study ID

Usage

```
get_mutations_by_study(
  study_id = NULL,
  molecular_profile_id = NULL,
  add_hugo = TRUE,
  base_url = NULL
)
```

Arguments

study_id A study ID to query mutations. If NULL, guesses study ID based on molecu-

lar_profile_id.

molecular_profile_id

a molecular profile to query mutations. If NULL, guesses molecular_profile_id

based on study ID.

add_hugo Logical indicating whether HugoGeneSymbol should be added to your resulting

data frame, if not already present in raw API results. Argument is TRUE by default. If FALSE, results will be returned as is (i.e. any existing Hugo Symbol

columns in raw results will not be removed).

base_url The database URL to query If NULL will default to URL set with set_cbioportal_db(<your_db>)

Value

A dataframe of mutations (maf file format)

Examples

```
## Not run:
get_mutations_by_study(study_id = "prad_msk_2019")
get_mutations_by_study(molecular_profile_id = "prad_msk_2019_mutations")
## End(Not run)
```

get_panel_by_sample

Get Gene Panel by study ID and sample ID

Description

Get Gene Panel by study ID and sample ID

Usage

```
get_panel_by_sample(
  study_id = NULL,
  sample_id = NULL,
  sample_study_pairs = NULL,
  base_url = NULL
)
```

Arguments

study_id

A string indicating the study ID from which to pull data. If no study ID, will guess the study ID based on your URL and inform. Only 1 study ID can be passed. If mutations/cna from more than 1 study needed, see sample_study_pairs

sample_id

a vector of sample IDs (character)

sample_study_pairs

A dataframe with columns: sample_id, study_id and molecular_profile_id (optional). Variations in capitalization of column names are accepted. This can be used in place of sample_id, study_id, molecular_profile_id arguments above if you need to pull samples from several different studies at once. If passed this will take overwrite sample_id, study_id, molecular_profile_id if also passed.

base_url

The database URL to query If NULL will default to URL set with set_cbioportal_db(<your_db>)

Value

a dataframe of a specific clinical attribute

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Examples

```
## Not run:
get_panel_by_sample(study_id = "blca_plasmacytoid_mskcc_2016",
    sample_id = "DS-sig-010-P2",
    base_url = 'www.cbioportal.org/api')

## End(Not run)

get_samples_by_patient
```

Get sample IDs for a given set of patient IDs

Description

Get sample IDs for a given set of patient IDs

Usage

```
get_samples_by_patient(patient_id = NULL, study_id = NULL, base_url = NULL)
```

Arguments

patient_id	A character string of sample IDs to query
study_id	A character string indicating which study ID should be searched. Only 1 study allowed. If NULL, we will guess a default study ID based on your database URL.
base_url	The database URL to query If NULL will default to URL set with set chioportal db(<vour db="">)</vour>

Value

A dataframe of patient IDs and corresponding sample IDs. If patient has multiple samples, there will be multiple rows per patient.

Examples

```
## Not run:
get_samples_by_patient(patient_id = c("P-0000034", "P-0000036"))
## End(Not run)
```

```
get_segments_by_sample
```

Get Copy Number Segmentation Data By Sample ID

Description

Get Copy Number Segmentation Data By Sample ID

Usage

```
get_segments_by_sample(
  sample_id = NULL,
  study_id = NULL,
  sample_study_pairs = NULL,
  base_url = NULL
)
```

Arguments

sample_id

a vector of sample IDs (character)

study_id

A string indicating the study ID from which to pull data. If no study ID, will guess the study ID based on your URL and inform. Only 1 study ID can be passed. If mutations/cna from more than 1 study needed, see sample_study_pairs

sample_study_pairs

A dataframe with columns: sample_id, study_id and molecular_profile_id (optional). Variations in capitalization of column names are accepted. This can be used in place of sample_id, study_id, molecular_profile_id arguments above if you need to pull samples from several different studies at once. If passed this will take overwrite sample_id, study_id, molecular_profile_id if also passed.

base_url

The database URL to query If NULL will default to URL set with set_cbioportal_db(<your_db>)

Value

A dataframe of CNA segments

Examples

get_segments_by_study

get_segments_by_study Get Copy Number Segmentation Data By Study

Description

Get Copy Number Segmentation Data By Study

Usage

```
get_segments_by_study(study_id = NULL, add_hugo = TRUE, base_url = NULL)
```

Arguments

study_id A study ID to query mutations. If NULL, guesses study ID based on molecu-

lar_profile_id.

add_hugo Logical indicating whether HugoGeneSymbol should be added to your resulting

data frame, if not already present in raw API results. Argument is TRUE by default. If FALSE, results will be returned as is (i.e. any existing Hugo Symbol

columns in raw results will not be removed).

base_url The database URL to query If NULL will default to URL set with set_cbioportal_db(<your_db>)

Value

A dataframe of CNA segments

Examples

```
## Not run:
get_segments_by_study(study_id = "prad_msk_2019")
get_segments_by_study(molecular_profile_id = "prad_msk_2019_cna")
## End(Not run)
```

get_study_info Get Metadata on All Available Studies in Database or a Specified Study

Description

Get Metadata on All Available Studies in Database or a Specified Study

Usage

```
get_study_info(study_id = NULL, base_url = NULL)
```

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Arguments

study_id one or more study IDs (see available_studies() to lookup IDs)

base_url The database URL to query. If NULL will default to URL set with set_cbioportal_db(<your_db>)

Value

A dataframe of study metadata

Examples

```
## Not run:
set_cbioportal_db("public")
get_study_info("acc_tcga")
## End(Not run)
```

impact_gene_info

IMPACT Gene Meta Data

Description

Dataframe labeling all genes included in IMPACT panels along with their corresponding platform ID and Entrez ID.

Usage

```
impact_gene_info
```

Format

A data frame with 470 genes

hugo_symbol Factor w/ 574 levels, Column containing all HUGO symbols genes included in IM-PACT

entrez_id Integer, contains all Entrez IDs for genes included in IMPACT

platform_341 Character, indicates whether each gene was included in IMPACT platform 341. Options are included and not included

platform_410 Character, indicates whether each gene was included in IMPACT platform 410. Options are included and not included

platform_468 Character, indicates whether each gene was included in IMPACT platform 468. Options are included and not included

alias A nested dataframe of aliases for each gene and corresponding entrez gene ids for aliases if they exist

Source

```
http://www.cbioportal.org/
```

set_cbioportal_db 29

set_cbioportal_db

Connect to cBioPortal DB

Description

This function sets a base cBioPortal URL

Usage

```
set_cbioportal_db(db = NULL)
```

Arguments

db

The database URL to use as base URL for calls, or "public" for https://www.cbioportal.org/

Value

No return value, called for side effects. Will display an alert notifying if the user has successfully authenticated to cBioPortal.

Author(s)

Karissa Whiting, Daniel D. Sjoberg

Examples

```
## Not run:
set_cbioportal_db(db = "public")
## End(Not run)
```

 $test_cbioportal_db$

Test the Database Connection Anytime During your R Session

Description

Helps troubleshoot API issues during an R session

Usage

```
test_cbioportal_db()
```

Value

No return value, called for side effects. Will display an alert notifying if the user has successfully authenticated to cBioPortal

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Author(s)

Karissa Whiting, Daniel D. Sjoberg

Examples

```
## Not run:
set_cbioportal_db("public")
test_cbioportal_db()
## End(Not run)
```

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