Package ‘terraTCGAdat’

May 30, 2024

**Type** Package

**Title** OpenAccess TCGA Data on Terra as MultiAssayExperiment

**Version** 1.8.0

**Description** Leverage the existing open access TCGA data on Terra with well-established Bioconductor infrastructure. Make use of the Terra data model without learning its complexities. With a few functions, you can copy / download and generate a MultiAssayExperiment from the TCGA example workspaces provided by Terra.

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

**biocViews** Software, Infrastructure, DataImport

**Imports** BiocFileCache, dplyr, GenomicRanges, methods, RaggedExperiment, readr, S4Vectors, stats, tidyr, TCGAutils, utils

**Suggests** knitr, rmarkdown, BiocStyle, withr, testthat (>= 3.0.0)

**URL** https://github.com/waldronlab/terraTCGAdat

**BugReports** https://github.com/waldronlab/terraTCGAdat/issues

**VignetteBuilder** knitr

**License** Artistic-2.0

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**RoxygenNote** 7.2.3

**Config/testthat/edition** 3

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**Author** Marcel Ramos [aut, cre] (<https://orcid.org/0000-0002-3242-0582>)

**Maintainer** Marcel Ramos <marcel.ramos@roswellpark.org>
.getWorkspaceTable

Obtain the table of datasets from the Terra platform

Description

The datasets include all TCGA datasets that do not come from the Genomic Data Commons Data Repository because those data use a different data model.

Usage

.getWorkspaceTable(project = "^TCGA", cancerCode = ".*")

Arguments

  project character(1) A prefix for the regex search across all public projects on the terra platform (default: "^TCGA"). Usually, this does not change.
  cancerCode character(1) Corresponds to the TCGA cancer code (e.g., "ACC" for AdrenoCortical Carcinoma) of interest. The default value of (".*") provides all available cancer datasets.

Value

A ‘tibble’ ‘data.frame’ that match the project in put; by default, TCGA workspaces.
Obtain assay datasets from Terra

Usage

```r
getAssayData(
  assayName,
  sampleCode = "01",
  tablename = .DEFAULT_TABLENAME,
  workspace = terraTCGAworkspace(),
  namespace = .DEFAULT_NAMESPACE,
  metacols = .PARTICIPANT_METADATA_COLS,
  sampleIdx = TRUE
)
```

Arguments

- `assayName` character() The name of the assay dataset column from `getAssayTable` to import into the current workspace.
- `sampleCode` character(1) The sample code used to filtering samples e.g., "01" for Primary Solid Tumors, see `data("sampleTypes", package = "TCGAutils")` for reference
- `tablename` The Terra data model table from which to extract the clinical data (default: "sample")
- `workspace` character(1) The Terra Data Resources workspace from which to pull TCGA data (default: see `terraTCGAworkspace()`). This is set to a package-wide option.
- `namespace` character(1) The Terra Workspace Namespace that defaults to "broad-firecloud-tcga" and rarely needs to be changed.
- `metacols` The set of columns that comprise of the metadata columns. See the `.PARTICIPANT_METADATA_COLS` global variable
- `sampleIdx` numeric() index or TRUE. Specify an index for subsetting the assay data. This argument is mainly used for example and vignette purposes. To use all the data, use the default value (default: TRUE)

Value

Either a matrix or RaggedExperiment depending on the assay selected

See Also

`getAssayTable()`
Examples

```r
if (AnVIL::gcloud_exists())
  getAssayData(
    assayName = "protein_exp_mda_rppa_core_mdanderson_org_level_3__protein_normalization__data",
    sampleCode = c("01", "10"),
    workspace = "TCGA_ACC_OpenAccess_V1-0_DATA"
  )
```

---

**getAssayTable**

Obtain a reference table for assay data in the Terra data model

Description

The column names in the output can be used in the `getAssayData` function.

Usage

```r
getAssayTable(
  tablename = .DEFAULT_TABLENAME,
  metacols = .PARTICIPANT_METADATA_COLS,
  workspace = terraTCGAworkspace(),
  namespace = .DEFAULT_NAMESPACE
)
```

Arguments

- `tablename`: The Terra data model table from which to extract the clinical data (default: "sample")
- `metacols`: The set of columns that comprise of the metadata columns. See the .PARTICIPANT_METADATA_COLS global variable
- `workspace`: character(1) The Terra Data Resources workspace from which to pull TCGA data (default: see terraTCGAworkspace()). This is set to a package-wide option.
- `namespace`: character(1) The Terra Workspace Namespace that defaults to "broad-firecloud-tcga" and rarely needs to be changed.

Value

A tibble of pointers to resources within the Terra data model

Examples

```r
if (AnVIL::gcloud_exists())
  getAssayTable(workspace = "TCGA_COAD_OpenAccess_V1-0_DATA")
```
**getClinical**

*Obtain clinical data*

**Description**

The participant table may contain curated demographic information e.g., sex, age, etc.

**Usage**

```r
getClinical(
  columnName, 
  participants = TRUE,
  tablename = .DEFAULT_TABLENAME,
  workspace = terraTCGAworkspace(),
  namespace = .DEFAULT_NAMESPACE,
  verbose = TRUE,
  metacols = .PARTICIPANT_METADATA_COLS,
  participantIds = NULL
)
```

**Arguments**

- **columnName**
  - The name of the column to extract files, see `getClinicalTable` table. If not provided, the first column in the table will be used to obtain the clinical information.

- **participants**
  - logical(1) Whether to merge the participant table from `avtable("participant")` to the clinical data

- **tablename**
  - The Terra data model table from which to extract the clinical data (default: "sample")

- **workspace**
  - character(1) The Terra Data Resources workspace from which to pull TCGA data (default: see `terraTCGAworkspace()`). This is set to a package-wide option.

- **namespace**
  - character(1) The Terra Workspace Namespace that defaults to "broad-firecloud-tcga" and rarely needs to be changed.

- **verbose**
  - logical(1) Whether to output additional information regarding the workspace and namespace (default: TRUE).

- **metacols**
  - The set of columns that comprise of the metadata columns. See the .PARTICIPANT_METADATA_COLS global variable

- **participantIds**
  - character() TCGA participant identifiers usually in the form of "TCGA-AB-1234". By default, all available participant identifiers will be used. (default: NULL)

**Value**

A `DataFrame` with clinical information from TCGA. The metadata i.e., `metadata(object)` includes the `columnName` used to obtain the data.
Examples

```r
if (AnVIL::gcloud_exists())
  getClinical(
    workspace = "TCGA_ACC_OpenAccess_V1-0_DATA",
    participantIds = c("TCGA-OR-A5J1", "TCGA-OR-A5J2",
                       "TCGA-OR-A5J3", "TCGA-OR-A5J4")
  )
```

### `getClinicalTable`

Obtain the reference table for clinical data

#### Description

The column names in the output table can be used in the `getClinical` function.

#### Usage

```r
getClinicalTable(
  tablename = .DEFAULT_TABLENAME,
  metacols = .PARTICIPANT_METADATA_COLS,
  workspace = terraTCGAworkspace(),
  namespace = .DEFAULT_NAMESPACE,
  verbose = TRUE
)
```

#### Arguments

- `tablename`: The Terra data model table from which to extract the clinical data (default: "sample")
- `metacols`: The set of columns that comprise of the metadata columns. See the `.PARTICIPANT_METADATA_COLS` global variable
- `workspace`: character(1) The Terra Data Resources workspace from which to pull TCGA data (default: see `terraTCGAworkspace()`). This is set to a package-wide option.
- `namespace`: character(1) The Terra Workspace Namespace that defaults to "broad-firecloud-tcga" and rarely needs to be changed.
- `verbose`: logical(1) Whether to output additional information regarding the workspace and namespace (default: TRUE).

#### Value

A tibble of Google Storage resource locations e.g., gs://firecloud...
getTCGAdatalist

Examples

```r
if (AnVIL::gcloud_exists())
  getClinicalTable(
      workspace = "TCGA_AACC_OpenAccess_V1-0_DATA"
  )
```

---

getTCGAdatalist  
Import Terra TCGA data as a list

Description

Import Terra TCGA data as a list

Usage

```r
getTCGAdatalist(
  assayNames,  
sampleCode,  
workspace = terraTCGAWorkspace(),  
namespace = .DEFAULT_NAMESPACE,  
tablename = .DEFAULT_TABLENAME,  
sampleIdx = TRUE,  
verbose = TRUE
)
```

Arguments

- **assayNames**: character() A vector of assays selected from the colnames of `getAssayTable`
- **sampleCode**: character(1) The sample code used to filtering samples e.g., "01" for Primary Solid Tumors, see `data("sampleTypes", package = "TCGAutils")` for reference
- **workspace**: character(1) The Terra Data Resources workspace from which to pull TCGA data (default: see `terraTCGAWorkspace()`). This is set to a package-wide option.
- **namespace**: character(1) The Terra Workspace Namespace that defaults to "broad-firecloud-tcga" and rarely needs to be changed.
- **tablename**: The Terra data model table from which to extract the clinical data (default: "sample")
- **sampleIdx**: numeric() index or TRUE. Specify an index for subsetting the assay data. This argument is mainly used for example and vignette purposes. To use all the data, use the default value (default: TRUE)
- **verbose**: logical(1L) Whether to output additional details of the data facilitation.
sampleTypesTable

Value

A list of assay datasets

Examples

```r
if (AnVIL::gcloud_exists())
    getTCGAdatalist(
        assayNames = c("protein_exp__mda_rppa_core__mdanderson_org__Level_3__protein_normalization__data",
                        "snp__genome_wide_snp_6__broad_mit edu__Level_3__segmented_scna_minus_germline_cnv_hg18__seg"),
        sampleCode = c("01", "10"),
        workspace = "TCGA_COAD_OpenAccess_V1-0_DATA"
    )
```

sampleTypesTable  Get an overview of the samples available in the workspace

Description

The function provides an overview of samples from the avtables("sample") table for the current workspace. Along with the sample codes and frequencies, the output provides a description for each code and the short letter codes.

Usage

```r
sampleTypesTable(
    workspace = terraTCGAworkspace(),
    namespace = .DEFAULT_NAMESPACE,
    tablename = .DEFAULT_TABLENAME,
    verbose = TRUE
)
```

Arguments

- **workspace** character(1) The Terra Data Resources workspace from which to pull TCGA data (default: see terraTCGAworkspace()). This is set to a package-wide option.
- **namespace** character(1) The Terra Workspace Namespace that defaults to "broad-firecloud-tcga" and rarely needs to be changed.
- **tablename** The Terra data model table from which to extract the clinical data (default: "sample")
- **verbose** logical(1) Whether to output additional information regarding the workspace and namespace (default: TRUE).

Value

A tibble of sample codes and frequency along with their definition and short letter code
Examples

```r
if (AnVIL::gcloud_exists())
  sampleTypesTable(workspace = "TCGA_COAD_OpenAccess_V1-0_DATA")
```

### Description

Workspaces on Terra come pre-loaded with TCGA Data. The examples in the documentation correspond to the TCGA_COAD_OpenAccess_V1 workspace that can be found on app.terra.bio.

### Usage

```r
terraTCGAdata(
  clinicalName,
  assays,
  participants = TRUE,
  sampleCode = NULL,
  split = FALSE,
  workspace = terraTCGAworkspace(),
  namespace = .DEFAULT_NAMESPACE,
  tablename = .DEFAULT_TABLENAME,
  verbose = TRUE,
  sampleIdx = TRUE
)
```

### Arguments

- **clinicalName**: character(1) The column name taken from `getClinicalTable()` and downloaded to be included as the `colData`.
- **assays**: character() A character vector of assay names taken from `getAssayTable()`.
- **participants**: logical(1) Whether to merge the participant table from `avtable("participant")` to the clinical data.
- **sampleCode**: character() A character vector of sample codes from `sampleTypesTable()`. By default, (NULL) all samples are downloaded and kept in the data.
- **split**: logical(1L) Whether or not to split the `MultiAssayExperiment` by sample types using `splitAssays` helper function (default FALSE).
- **workspace**: character(1) The Terra Data Resources workspace from which to pull TCGA data (default: see `terraTCGAworkspace()`). This is set to a package-wide option.
- **namespace**: character(1) The Terra Workspace Namespace that defaults to "broad-firecloud-tcga" and rarely needs to be changed.
### TerraTCGAworkspace

**Description**

Terra allows access to about 71 open access TCGA datasets. A dataset workspace can be set using the `terraTCGAworkspace` function with a `projectName` input. Use the `selectTCGAworkspace` function to select a TCGA data workspace from an interactive table.

**Usage**

```r
terraTCGAworkspace(projectName = getOption("terraTCGAdata.workspace", NULL))

selectTCGAworkspace(
    projectName = getOption("terraTCGAdata.workspace", NULL),
    verbose = FALSE,
    ...
)
```

#### Parameters

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>tablename</code></td>
<td>The Terra data model table from which to extract the clinical data (default: &quot;sample&quot;)</td>
</tr>
<tr>
<td><code>verbose</code></td>
<td>logical(1) Whether to output additional information regarding the workspace and namespace (default: TRUE).</td>
</tr>
<tr>
<td><code>sampleIdx</code></td>
<td>numeric() index or TRUE. Specify an index for subsetting the assay data. This argument is mainly used for example and vignette purposes. To use all the data, use the default value (default: TRUE)</td>
</tr>
</tbody>
</table>

**Value**

A `MultiAssayExperiment` object with n number of assays corresponding to the `assays` argument.

**Examples**

```r
if (AnVIL::gcloud_exists())
    terraTCGAdata(
        clinicalName = "clin_bio_nationwidechildrens_org_Level_1_biospecimen_clin",
        assays = c("protein_exp_mda_rppa_core_manderson_org_Level_3_protein_normalization_data",
                    "rnaseqv2_illumina_hiseq_rnaseqv2_unc_edu_Level_3_RSEM_genes_normalized_data"),
        workspace = "TCGA_COAD_OpenAccess_V1-0_DATA",
        sampleCode = NULL,
        sampleIdx = 1:4,
        split = FALSE
    )
```
terraTCGAWorkspace

Arguments

projectName character(1) A project code usually in the form of TCGA_CODE_OpenAccess_V1-0_DATA. See selectTCGAWorkspace to interactively select from a table of project codes.

verbose logical(1) Whether to provide more informative messages when the "terraTCGAdata.workspace" option is set.

... further arguments passed down to lower level functions, not intended for the end user.

Details

Note that GDC workspaces are not supported and are excluded from the search results. GDC workspaces use a Terra workflow to download TCGA data rather than providing Google Bucket storage locations for easy data retrieval. To reset the terraTCGAWorkspace, use terraTCGAWorkspace(NULL) and you will be prompted to select from a list of TCGA workspaces. You may also check the current active workspace by running terraTCGAWorkspace() without any inputs.

Value

A Terra TCGA Workspace name

Functions

- selectTCGAWorkspace(): Function to interactively select from the available TCGA data workspaces in Terra. The 'projectName' argument and 'terraTCGAdata.workspace' option must be 'NULL' to enable the interactive gadget.

Examples

```r
if (AnVIL::gcloud_exists() && interactive()) {
  selectTCGAWorkspace()
  terraTCGAWorkspace()
}
```
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