

Package ‘HumanTranscriptomeCompendium’

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Title Tools to work with a Compendium of 181000 human transcriptome sequencing studies

Description Provide tools for working with a compendium of human transcriptome sequences (originally htxcomp).

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Author Sean Davis, Vince Carey

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Imports shiny, ssrch, S4Vectors, SummarizedExperiment, utils

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Maintainer VJ Carey <stvjc@channing.harvard.edu>

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addRD	<i>add gene-level rowData derived from transcript level rowRanges</i>
-------	---

Description

add gene-level rowData derived from transcript level rowRanges

Usage

addRD(x)

Arguments

x result of htx_load()

Value

RangedSummarizedExperiment with enhanced rowData

Examples

```
# this function operates on a SummarizedExperiment that has
# transcript-level rowRanges but gene-level quantifications
addRD
```

ca43k	<i>app to survey 43000 cancer transcriptomes</i>
-------	--

Description

app to survey 43000 cancer transcriptomes

Usage

```
ca43k()
```

Value

a SummarizedExperiment

Note

Copies source code and metadata to a temporary folder and executes shiny::runApp there; sets working directory to folder where ca43k was called when app is exited. Also will return either NULL or a SummarizedExperiment at conclusion.

Examples

```
ca43k
```

get_ds4841	<i>return instance of ssrch::DocSet with metadata on 4841 human transcriptome studies in NCBI SRA</i>
------------	---

Description

return instance of ssrch::DocSet with metadata on 4841 human transcriptome studies in NCBI SRA

Usage

```
get_ds4841(  
  cache = BiocFileCache::BiocFileCache(),  
  csv_zip_path = path_doc4842()  
)
```

Arguments

cache	instance of 'BiocFileCache', defaults to 'BiocFileCache::BiocFileCache()'
csv_zip_path	a path leading to the zip file of CSV for metadata in the DocSet instance

Value

instance of DocSet as defined in ssrch package

Note

will bind the correct value of 'zipf' in 'environment(ds4841@doc_retriever)', which depends on details of installation

Examples

```
get_ds4841()
```

htx_app

explore SRA metadata

Description

explore SRA metadata

Usage

```
htx_app()
```

Value

a SummarizedExperiment can be requested through an event

Note

This function deals with extraction of compendium elements. The overall scope is determined by HumanTranscriptomeCompendium::studTable which is the list of all studies with taxon 9606, strategy RNA-seq, source transcriptomic. Some studies will not have experiments in the compendium, and if such are selected, a warning will be generated in the session.

Examples

```
if (interactive()) htx_app()
```

htx_load	<i>load a SummarizedExperiment shell for the Human Transcriptome Compendium</i>
----------	---

Description

load a SummarizedExperiment shell for the Human Transcriptome Compendium

Usage

```
htx_load(  
  remotePath = "https://biocfound-bigrnatx.s3.us-west-2.amazonaws.com/rangedHtxGeneSE.rds",  
  cache = BiocFileCache::BiocFileCache(),  
  genesOnly = TRUE  
)
```

Arguments

remotePath	path to an RDS representation of the DelayedArray-based SummarizedExperiment
cache	a BiocFileCache instance, defaulting to value of BiocFileCache()
genesOnly	logical(1) if TRUE return reference to SummarizedExperiment with gene-level quantifications; in this case the remotePath value is set to 'https://s3.amazonaws.com/bcfound-bigrna/rangedHtxGeneSE.rds'.

Value

a RangedSummarizedExperiment instance

Examples

```
htx_load
```

htx_query_by_study_accession	<i>retrieve 'restfulSE' SummarizedExperiment instance for selected studies in htx compendium</i>
------------------------------	--

Description

retrieve 'restfulSE' SummarizedExperiment instance for selected studies in htx compendium

Usage

```
htx_query_by_study_accession(studies, htxSE, ...)
```

Arguments

<code>studies</code>	character vector of study accessions
<code>htxSE</code>	SummarizedExperiment instance, typically the result of <code>htx_load()</code> , which we don't want to repeat needlessly
<code>...</code>	passed to 'htx_load', ignored if 'se' is nonmissing

Value

SummarizedExperiment instance

Note

This function was designed to perform a single query on a 'fresh' compendium image from 'htx_load()'. However, one could consider iterating the process to build up metadata on multiple series of studies. This is not likely to succeed without careful manipulation of the `colData` of the input SummarizedExperiment. A message will be written if the input SummarizedExperiment appears to be other than a 'fresh' 'htx_load' result.

Examples

```
htx_query_by_study_accession("ERP011411")
```

<code>htx_query_by_text</code>	<i>subset compendium through keyword lookup</i>
--------------------------------	---

Description

subset compendium through keyword lookup

Usage

```
htx_query_by_text(query, ..., tryGrep = TRUE, ignore.case = TRUE)
```

Arguments

<code>query</code>	character(1) to be found in <code>ls(ssrch::kw2docs(get_ds4841()))</code>
<code>...</code>	passed to 'htx_query_by_study_accession'
<code>tryGrep</code>	logical(1) if TRUE, 'query' does not match any keyword directly, it will be treated as a regular expression and the vector of keywords will be grepped for pattern 'query'; defaults to TRUE
<code>ignore.case</code>	logical(1) used when <code>tryGrep</code> is TRUE, defaults to TRUE

Value

SummarizedExperiment instance

Note

The DocSet instance returned by `'get_ds4841()'` is used. Lookups are case-sensitive. Look carefully at note for `'htx_query_by_study_accession'` to understand logic of incrementing metadata on a given input SummarizedExperiment.

Examples

```
htx_query_by_text("HNRNPC")
```

```
HumanTranscriptomeCompendium.colnames
```

character vector of available samples in HDF cloud assay representation

Description

character vector of available samples in HDF cloud assay representation

Usage

```
HumanTranscriptomeCompendium.colnames
```

Format

character vector

Source

compendium processing

Examples

```
head(HumanTranscriptomeCompendium::HumanTranscriptomeCompendium.colnames)
```

```
load_bigrnaFiles      obtain listing of contents of BigRNA compendium (salmon runs)
```

Description

obtain listing of contents of BigRNA compendium (salmon runs)

Usage

```
load_bigrnaFiles(cache = BiocFileCache::BiocFileCache())
```

Arguments

cache instance of 'BiocFileCache', defaults to 'BiocFileCache::BiocFileCache()'

Value

a named vector

Examples

```
if (interactive()) head(load_bigrnaFiles())
```

load_experTable	<i>obtain listing of experiments and submission date/time in compendium</i>
-----------------	---

Description

obtain listing of experiments and submission date/time in compendium

Usage

```
load_experTable(cache = BiocFileCache::BiocFileCache())
```

Arguments

cache instance of 'BiocFileCache', defaults to 'BiocFileCache::BiocFileCache()'

Value

a data.frame

Examples

```
if (interactive()) head(load_experTable())
```

load_studTable	<i>obtain listing of all studies in compendium</i>
----------------	--

Description

obtain listing of all studies in compendium

Usage

```
load_studTable(cache = BiocFileCache::BiocFileCache())
```

Arguments

cache instance of ‘BiocFileCache’, defaults to ‘BiocFileCache::BiocFileCache()’

Value

a data.frame

Examples

```
if (interactive()) head(load_studTable())
```

path_doc4842	<i>return path to metadata csvs in zip file</i>
--------------	---

Description

return path to metadata csvs in zip file

Usage

```
path_doc4842(cache = BiocFileCache::BiocFileCache())
```

Arguments

cache instance of ‘BiocFileCache’, defaults to ‘BiocFileCache::BiocFileCache()’

Value

path to zipfile

Note

CSVs were retrieved using methods provided at <https://api-omicidx.cancerdatasoci.org/sra/1.0/ui/> and zipped together. Function will lodge zipfile in ‘cache’ if not present.

Examples

```
path_doc4842()
```

procExpToGene	<i>acquire a single sample from bigRNA compendium specified by accession and develop gene-level quantifications using tximport</i>
---------------	--

Description

acquire a single sample from bigRNA compendium specified by accession and develop gene-level quantifications using tximport

Usage

```
procExpToGene(
  acc,
  tx2gene = tx2gene_gencode27(),
  urlprefix = "http://bigrna-test.cancerdatasci.org/data/?accession=",
  manifestdata = HumanTranscriptomeCompendium::load_bigrnaFiles(),
  regexp = "quant.sf.bz2|json"
)
```

Arguments

acc	character(1) sample-level accession as defined in SRA
tx2gene	a data.frame instance mapping transcript identifiers used in the compendium to gene identifiers. See note.
urlprefix	character(1) where the salmon run outputs are lodged, with acc a subfolder defined through the manifestData parameter.
manifestdata	a character vector defining folders (under results/human/27/ with salmon outputs).
regexp	a character(1) regular expression for filtering filename elements in manifestdata to define which salmon output components in the bigrna compendium are retrieved.

Value

the result of a tximport run

Note

The tx2gene_gencode function supplied with this package uses the tximportData package contents to create the data.frame for use as tx2gene. The system2 function is used to generate folders to be used by tximport.

Examples

```
# this example involves nontrivial internet communications
args(procExpToGene)

td = tempdir()
od = getwd()
setwd(td)
nn = procExpToGene("ERX1097381")
str(nn)
setwd(od)
```

tx2gene_gencode27	<i>generate a data.frame mapping gencode 27 ensembl transcript identifiers to ensembl gene identifiers</i>
-------------------	--

Description

generate a data.frame mapping gencode 27 ensembl transcript identifiers to ensembl gene identifiers

Usage

```
tx2gene_gencode27()
```

Value

a data.frame with 200401 rows mapping transcript identifiers in column 1 to 58288 gene symbols in column 2.

Note

Uses CSV in tximportData to acquire the information.

Examples

```
head(tx2gene_gencode27())
```

uniqueAcc_120518	<i>experiment accessions available in compendium as of may 12 2018</i>
------------------	--

Description

experiment accessions available in compendium as of may 12 2018

Usage

```
uniqueAcc_120518
```

Format

```
data.frame
```

Source

SRAdBv2 may 12 2018

Examples

```
head(HumanTranscriptomeCompendium::uniqueAcc_120518)
```

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