Package 'seeker'

August 27, 2024

```
Type Package
Title Simplified Fetching and Processing of Microarray and RNA-Seq
            Data
Version 1.1.6
Description Wrapper around various existing tools and command-line interfaces,
            providing a standard interface, simple parallelization, and detailed logging.
            For microarray data, maps probe sets to standard gene IDs, building on
            'GEOquery' Davis and Meltzer (2007) <doi:10.1093/bioinformatics/btm254>,
            'ArrayExpress' Kauffmann et al. (2009) <doi:10.1093/bioinformatics/btp354>,
            Robust multi-array average 'RMA' Irizarry et al. (2003) <doi:10.1093/biostatistics/4.2.249>,
            and 'BrainArray' Dai et al. (2005) <doi:10.1093/nar/gni179>.
            For RNA-seq data, fetches metadata and raw reads from National Center for Biotechnology
            Information (NCBI) Sequence Read Archive (SRA), performs standard adapter and
            quality trimming using 'TrimGalore' Krueger <a href="https:">https:</a>
            //github.com/FelixKrueger/TrimGalore>,
            performs quality control checks using 'FastQC' Andrews <a href="https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/https://example.com/ht
            //github.com/s-andrews/FastQC>,
            quantifies transcript abundances using 'salmon' Pa-
            tro et al. (2017) <doi:10.1038/nmeth.4197> and potentially
            'refgenie' Stolarczyk et al. (2020) <doi:10.1093/gigascience/giz149>,
            aggregates the results using 'MultiQC' Ewels et al. (2016) <doi:10.1093/bioinformatics/btw354>,
            maps transcripts to genes using 'biomaRt' Durinkck et al. (2009) <doi:10.1038/nprot.2009.97>,
            and summarizes transcript-level quantifications for gene-level analyses using
            'tximport' Soneson et al. (2015) <doi:10.12688/f1000research.7563.2>.
URL https://seeker.hugheylab.org, https://github.com/hugheylab/seeker
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Depends R (>= 3.5)
Imports affy (>= 1.68.0), AnnotationDbi (>= 1.52.0), BiocManager (>=
            1.30.0), biomaRt (>= 2.36.1), checkmate (>= 2.1.0), curl (>=
            3.2), data.table (>= 1.11.8), foreach (>= 1.4.4), GEOquery (>=
            2.58.0), glue (>= 1.5.0), jsonlite (>= 1.7.2), methods, qs (>=
```

2 checkDefaultCommands

```
0.21.2), R.utils (>= 2.11.0), RCurl (>= 1.98), readr (>=
1.4.0), sessioninfo (>= 1.2.0), tximport (>= 1.8.0), withr (>=
2.4.2), yaml (>= 2.2.1)

Suggests ArrayExpress (>= 1.62.0), Biobase, doParallel (>= 1.0.17),
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Config/testthat/edition 3

VignetteBuilder knitr

NeedsCompilation no

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Repository CRAN

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```

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 ${\it checkDefaultCommand-} line\ interfaces$

Description

This function checks whether the command-line tools used by seeker are accessible in the expected places.

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Usage

```
checkDefaultCommands(keepIdx = FALSE)
```

Arguments

keepIdx Logical indicating whether to keep the idx column of the resulting data.table.

For internal use only.

Value

A data.table with columns for command, path, and version.

See Also

```
installSysDeps()
```

	D E .00
fastqc	Run FastQC

Description

This function calls fastqc using system2(). To run in parallel, register a parallel backend, e.g., using doParallel::registerDoParallel().

Usage

```
fastqc(filepaths, outputDir = "fastqc_output", cmd = "fastqc", args = NULL)
```

Arguments

filepaths	Paths to fastq files.	For single-end reads,	each element should be a single	
	filepath. For paired-	end reads, each element	t can be two filepaths separated by	

";".

outputDir Directory in which to store output. Will be created if it doesn't exist.

cmd Name or path of the command-line interface.

args Additional arguments to pass to the command-line interface.

Value

A vector of exit codes, invisibly.

```
seeker()
```

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fastqscreen

Run FastQ Screen

Description

This function calls fastq_screen using system2(). To run in parallel, register a parallel backend, e.g., using doParallel::registerDoParallel().

Usage

Arguments

filepaths Paths to fastq files. For single-end reads, each element should be a single

filepath. For paired-end reads, each element can be two filepaths separated by

";".

outputDir Directory in which to store output. Will be created if it doesn't exist.

cmd Name or path of the command-line interface.

args Additional arguments to pass to the command-line interface.

Value

A vector of exit codes, invisibly.

See Also

seeker()

fetch

Fetch files

Description

This function uses the NCBI SRA Toolkit via system2() to download files from SRA and convert them to fastq.gz. To process files in parallel, register a parallel backend, e.g., using doParallel::registerDoParallel(). Beware that intermediate files created by fasterq-dump are uncompressed and could require hundreds of gigabytes if files are processed in parallel.

fetch 5

Usage

```
fetch(
   accessions,
   outputDir,
   overwrite = FALSE,
   keepSra = FALSE,
   prefetchCmd = "prefetch",
   prefetchArgs = NULL,
   fasterqdumpCmd = "fasterq-dump",
   fasterqdumpArgs = NULL,
   pigzCmd = "pigz",
   pigzArgs = NULL
)
```

Arguments

	accessions	Character vector of SRA run accessions.
	outputDir	String indicating the local directory in which to save the files. Will be created if it doesn't exist.
	overwrite	Logical indicating whether to overwrite files that already exist in outputDir.
	keepSra	Logical indicating whether to keep the ".sra" files.
	prefetchCmd	String indicating command for prefetch, which downloads ".sra" files.
	prefetchArgs	Character vector indicating arguments to pass to prefetch.
	fasterqdumpCmd	String indicating command for fasterq-dump, which uses ".sra" files to create ".fastq" files.
fasterqdumpArgs		
		Character vector indicating arguments to pass to fasterq-dump.
	pigzCmd	String indicating command for pigz, which converts ".fastq" files to ".fastq.gz" files.
	pigzArgs	Character vector indicating arguments to pass to pigz.

Value

A list. As the function runs, it updates a tab-delimited log file in outputDir called "progress.tsv".

```
seeker(), fetchMetadata()
```

6 getPlatforms

fetchMetadata

Fetch metadata for a genomic study

Description

This function can use the API of the European Nucleotide Archive (recommended) or the Sequence Read Archive.

Usage

```
fetchMetadata(
  bioproject,
  host = c("ena", "sra"),
  fields = c("study_accession", "sample_accession", "secondary_sample_accession",
    "sample_alias", "sample_title", "experiment_accession", "run_accession", "fastq_md5",
    "fastq_ftp", "fastq_aspera"),
    file = NULL
)
```

Arguments

bioproject String indicating bioproject accession.

host String indicating from where to fetch the metadata.

fields Character vector indicating which fields to fetch, if host is "ena".

file String indicating output file path, if not NULL.

Value

A data.table.

See Also

```
seeker(), fetch()
```

getPlatforms

Get supported microarray platforms

Description

Get supported microarray platforms

Usage

```
getPlatforms(type = c("cdf", "mapping"))
```

getSalmonMetadata 7

Arguments

type

String indicating whether to get supported platforms for processing raw Affymetrix data using custom CDF or for mapping already processed data from probes to genes.

Value

A data.table.

getSalmonMetadata

Aggregrate metadata from salmon quantifications

Description

Aggregrate metadata from salmon quantifications

Usage

```
getSalmonMetadata(inputDir, outputDir = "data")
```

Arguments

inputDir

Directory that contains output from salmon.

outputDir

Directory in which to save the result, a file named "salmon_meta_info.csv". If

NULL, no file is saved.

Value

```
A data.table, invisibly.

#' @seealso seeker(), salmon()
```

getTx2gene

Get mapping between transcripts and genes

Description

This function uses the biomaRt package.

Usage

```
getTx2gene(
  organism = "mmusculus",
  version = NULL,
  outputDir = "data",
  checkArgsOnly = FALSE
)
```

Arguments

organism String used to pass paste0(organism, "_gene_ensembl") as the dataset ar-

gument to biomaRt::useEnsembl(). To see available datasets, do mart = biomaRt::useEnsembl("general datasets)

version Passed to biomaRt::useEnsemb1(). NULL indicates the latest version. To see

available versions, do biomaRt::listEnsemblArchives().

outputDir Directory in which to save the result, a file named "tx2gene.csv.gz". If NULL, no

file is saved.

checkArgsOnly Logical indicating whether to only check function arguments. Used for testing.

Value

If checkArgsOnly is FALSE, a data.table based on the result from biomaRt::getBM(), with an attribute "version". Otherwise 0.

See Also

```
seeker(), tximport()
```

installCustomCdfPackages

Install custom CDF packages

Description

Install Brainarray custom CDFs for processing raw Affymetrix data. See http://brainarray.mbni.med.umich.edu/Brainarray/Database/CustomCDF/CDF_download.asp.

Usage

```
installCustomCdfPackages(pkgs, ver = 25, dryRun = FALSE)
```

Arguments

pkgs Character vector of package names, e.g., "hgu133ahsentrezgcdf".

ver Integer version number (25 as of 5 Jan 2021).

dryRun Logical indicating whether to actually install the packages.

Value

A character vector of URLs, invisibly.

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installSysDeps

Install seeker's system dependencies

Description

This function installs and configures the various programs required for seeker to fetch and process RNA-seq data.

Usage

```
installSysDeps(
    sraToolkitDir,
    minicondaDir,
    refgenieDir,
    rprofileDir,
    minicondaEnv = "seeker",
    refgenieGenomes = NULL,
    fastqscreenDir = NULL
)
```

Arguments

sraToolkitDir String indicating directory in which to install the SRA Toolkit. Recommended to use "~", the home directory. If NULL, the Toolkit will not be installed. minicondaDir String indicating directory in which to install Miniconda. Recommended to use "~", the home directory. If NULL, Miniconda will not be installed. refgenieDir String indicating directory in which to store the directory of genome assets from refgenie, which will be named "refgenie_genomes". Recommended to use "~", the home directory. Only used if minicondaDir is not NULL. rprofileDir String indicating directory in which to create or modify .Rprofile, which is run by R on startup. Common options are "~" or ".". minicondaEnv String indicating name of the Miniconda environment in which to install various conda packages (fastq-screen, fastqc, multiqc, pigz, refgenie, salmon, and trimgalore). refgenieGenomes Character vector indicating genome assets, such as transcriptome indexes for salmon(), to pull from refgenomes using refgenie. If NULL, no assets are fetched. String indicating directory in which to download the genomes for fastqscreen(). fastqscreenDir

This takes a long time. If NULL, genomes are not downloaded.

Value

NULL, invisibly

```
seeker()
```

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multiqc

Run MultiQC

Description

This function calls multiqc using system2().

Usage

```
multiqc(
  parentDir = ".",
  outputDir = "multiqc_output",
  cmd = "multiqc",
  args = NULL
)
```

Arguments

parentDir Directory that contains output to be aggregated.

outputDir Directory in which to store output. Will be created if it doesn't exist.

cmd Name or path of the command-line interface.

args Additional arguments to pass to the command-line interface.

Value

An exit code, invisibly.

See Also

seeker()

salmon

Run Salmon

Description

This function calls salmon using system2(). To run in parallel, register a parallel backend, e.g., using doParallel::registerDoParallel().

Usage

```
salmon(
  filepaths,
  samples,
  indexDir,
  outputDir = "salmon_output",
  cmd = "salmon",
  args = c("-1 A -q --seqBias --gcBias --no-version-check -p",
    foreach::getDoParWorkers()),
  compress = TRUE
)
```

Arguments

filepaths Paths to fastq files. For single-end reads, each element should be a single

filepath. For paired-end reads, each element should be two filepaths separated

by ";".

samples Corresponding sample names for fastq files.

indexDir Directory that contains salmon index.

outputDir Directory in which to store output. Will be created if it doesn't exist.

cmd Name or path of the command-line interface.

args Additional arguments to pass to the command-line interface.

compress Logical indicating whether to gzip the quantification file (quant.sf) from salmon.

Does not affect downstream analysis.

Value

A vector of exit codes, invisibly.

See Also

```
seeker(), getSalmonMetadata()
```

seeker

Process RNA-seq data end to end

Description

This function selectively performs various steps to process RNA-seq data. See also the vignettes: browseVignettes('seeker').

Usage

```
seeker(params, parentDir = ".", dryRun = FALSE)
```

Arguments

params

Named list of parameters with components:

- study: String used to name the output directory within parentDir.
- metadata: Named list with components:
 - run: Logical indicating whether to fetch metadata. See fetchMetadata().
 If TRUE, saves a file parentDir/study/metadata.csv. If FALSE, expects that file to already exist. The unmodified fetched or found metadata is saved to a file parentDir/study/metadata_original.csv. Following components are only checked if run is TRUE.
 - bioproject: String indicating the study's bioproject accession.
 - include: Optional named list for specifying which rows of metadata to include for further processing, with components:
 - * colname: String indicating column in metadata
 - * values: Vector indicating values within colname
 - exclude: Optional named list for specifying which rows of metadata to exclude from further processing (superseding include), with components:
 - * colname: String indicating column in metadata
 - * values: Vector indicating values within colname
- fetch: Named list with components:
 - run: Logical indicating whether to fetch files from SRA. See fetch(). If TRUE, saves files to parentDir/study/fetch_output. Whether TRUE or FALSE, expects metadata to have a column "run_accession", and updates metadata with column "fastq_fetched" containing paths to files in parentDir/study/fetch_output. Following components are only checked if run is TRUE.
 - keep: Logical indicating whether to keep fastq.gz files when all processing steps have completed. NULL indicates TRUE.
 - overwrite: Logical indicating whether to overwrite files that already exist. NULL indicates to use the default in fetch().
 - keepSra: Logical indicating whether to keep the ".sra" files. NULL indicates to use the default in fetch().
 - prefetchCmd: String indicating command for prefetch, which down-loads ".sra" files. NULL indicates to use the default in fetch().
 - prefetchArgs: Character vector indicating arguments to pass to prefetch.
 NULL indicates to use the default in fetch().
 - fasterqdumpCmd: String indicating command for fasterq-dump, which uses ".sra" files to create ".fastq" files. NULL indicates to use the default in fetch().
 - prefetchArgs: Character vector indicating arguments to pass to fasterqdump. NULL indicates to use the default in fetch().
 - pigzCmd: String indicating command for pigz, which converts ".fastq" files to ".fastq.gz" files. NULL indicates to use the default in fetch().
 - pigzArgs: Character vector indicating arguments to pass to pigz. NULL indicates to use the default in fetch().

- trimgalore: Named list with components:
 - run: Logical indicating whether to perform quality/adapter trimming of reads. See trimgalore(). If TRUE, expects metadata to have a column "fastq_fetched" containing paths to fastq files in parentDir/study/fetch_output, saves trimmed files to parentDir/study/trimgalore_output, and updates metadata with column "fastq_trimmed". If FALSE, expects and does nothing. Following components are only checked if run is TRUE.
 - keep: Logical indicating whether to keep trimmed fastq files when all processing steps have completed. NULL indicates TRUE.
 - cmd: Name or path of the command-line interface. NULL indicates to use the default in trimgalore().
 - args: Additional arguments to pass to the command-line interface.
 NULL indicates to use the default in trimgalore().
 - pigzCmd: String indicating command for pigz, which converts ".fastq" files to ".fastq.gz" files. NULL indicates to use the default in trimgalore().
- fastqc: Named list with components:
 - run: Logical indicating whether to perform QC on reads. See fastqc(). If TRUE and trimgalore\$run is TRUE, expects metadata to have a column "fastq_trimmed" containing paths to fastq files in parentDir/study/trimgalore_output. If TRUE and trimgalore\$run is FALSE, expects metadata to have a column "fastq_fetched" containing paths to fastq files in parentDir/study/fetch_output. If TRUE, saves results to parentDir/study/fastqc_output. If FALSE, expects and does nothing. Following components are only checked if run is TRUE.
 - keep: Logical indicating whether to keep fastqc files when all processing steps have completed. NULL indicates TRUE.
 - cmd: Name or path of the command-line interface. NULL indicates to use the default in fastqc().
 - args: Additional arguments to pass to the command-line interface.
 NULL indicates to use the default in fastqc().
- salmon: Named list with components:
 - run: Logical indicating whether to quantify transcript abundances. See salmon(). If TRUE and trimgalore\$run is TRUE, expects metadata to have a column "fastq_trimmed" containing paths to fastq files in parentDir/study/trimgalore_output. If TRUE and trimgalore\$run is FALSE, expects metadata to have a column "fastq_fetched" containing paths to fastq files in parentDir/study/fetch_output. If TRUE, saves results to parentDir/study/salmon_output and parentDir/study/salmon_meta_info.csv. If FALSE, expects and does nothing. Following components are only checked if run is TRUE.
 - indexDir: Directory that contains salmon index.
 - sampleColname: String indicating column in metadata containing sample ids. NULL indicates "sample_accession", which should work for data from SRA and ENA.
 - keep: Logical indicating whether to keep quantification results when all processing steps have completed. NULL indicates TRUE.

- cmd: Name or path of the command-line interface. NULL indicates to use the default in salmon().
- args: Additional arguments to pass to the command-line interface.
 NULL indicates to use the default in salmon().
- multiqc: Named list with components:
 - run: Logical indicating whether to aggregate results of various processing steps. See multiqc(). If TRUE, saves results to parentDir/study/multiqc_output. If FALSE, expects and does nothing. Following components are only checked if run is TRUE.
 - cmd: Name or path of the command-line interface. NULL indicates to use the default in multiqc().
 - args: Additional arguments to pass to the command-line interface.
 NULL indicates to use the default in multiqc().
- tximport: Named list with components:
 - run: Logical indicating whether to summarize transcript- or gene-level estimates for downstream analysis. See tximport(). If TRUE, expects metadata to have a column sampleColname of sample ids, and expects a directory parentDir/study/salmon_output containing directories of quantification results, and saves results to parentDir/study/tximport_output.qs. If FALSE, expects and does nothing. Following components are only checked if run is TRUE.
 - tx2gene: Optional named list with components:
 - * organism: String indicating organism and thereby ensembl gene dataset. See getTx2gene().
 - * version: Optional number indicating ensembl version. NULL indicates the latest version. See getTx2gene().
 - * filename: Optional string indicating name of pre-existing text file in parentDir/params\$study containing mapping between transcripts (first column) and genes (second column), with column names in the first row. If filename is specified, organism and version must not be specified.

If not NULL, saves a file parentDir/study/tx2gene.csv.gz.

- countsFromAbundance: String indicating whether or how to estimate counts using estimated abundances. See tximport::tximport().
- ignoreTxVersion: Logical indicating whether to the version suffix on transcript ids. NULL indicates to use TRUE. See tximport::tximport().

params can be derived from a yaml file, see vignette("introduction", package = "seeker"). The yaml representation of params will be saved to parentDir/params\$study/params.yml

parentDir

Directory in which to store the output, which will be a directory named according to params\$study.

dryRun

Logical indicating whether to check the validity of inputs without actually fetching or processing any data.

Value

Path to the output directory parentDir/params\$study, invisibly.

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See Also

```
fetchMetadata(), fetch(), trimgalore(), fastqc(), salmon(), multiqc(), tximport(), installSysDeps(),
seekerArray()
```

Examples

```
## Not run:
doParallel::registerDoParallel()
params = yaml::read_yaml('my_params.yaml')
seeker(params)
## End(Not run)
```

seekerArray

Process microarray data end to end

Description

This function fetches data and metadata from NCBI GEO and ArrayExpress, processes raw Affymetrix data using RMA and custom CDFs from Brainarray, and maps probes to genes. See also the vignettes: browseVignettes('seeker').

Usage

```
seekerArray(
   study,
   geneIdType,
   platform = NULL,
   parentDir = ".",
   metadataOnly = FALSE
)
```

Arguments

study String indicating the study accession and used to name the output directory

within parentDir. Must start with "GSE", "E-", or "LOCAL". If starts with "GSE", data are fetched using GEOquery::getGEO(). If starts with "E-", data are fetched using ArrayExpress::getAE(). If starts with "LOCAL", data in the form of cel(.gz) files must in the directory parentDir/study/raw, and parentDir/study must contain a file "sample_metadata.csv" that has a column sample_id con-

taining the names of the cel(.gz) files without the file extension.

geneIdType String indicating whether to map probes to gene IDs from Ensembl ("ensembl")

or Entrez ("entrez").

platform String indicating the GEO-based platform accession for the raw data. See https:

//www.ncbi.nlm.nih.gov/geo/browse/?view=platforms. Only necessary if study starts with "LOCAL", or starts with "GSE" and the study uses multiple

platforms.

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parentDir Directory in which to store the output, which will be a directory named accord-

ing to study.

metadataOnly Logical indicating whether to only process the sample metadata, and skip pro-

cessing the expression data.

Details

The standard output:

naive_expression_set.qs: Initial ExpresssionSet generated by GEOquery::getGEO or ArrayExpress::ae2bioc().
 Should generally not be used if sample_metadata.csv and gene_expression_matrix.qs are available.

- sample_metadata.csv: Table of sample metadata. Column sample_id matches colnames of the gene expression matrix.
- gene_expression_matrix.qs: Rows correspond to genes, columns to samples. Expression values are log2-transformed.
- custom_cdf_name.txt: Name of custom CDF package used by affy::justRMA() to process and normalize raw Affymetrix data and map probes to genes.
- feature_metadata.qs: GPL object, if gene expression matrix was generated from processed data
- probe_gene_mapping.csv.gz: Table of probes and genes, if gene expression matrix was generated from processed data.
- "raw" directory: Contains raw Affymetrix files.
- params.yml: Parameters used to process the dataset.
- session.log: R session information.

The output may include other files from NCBI GEO or ArrayExpress. Files with extension "qs" can be read into R using qs::qread().

Value

Path to the output directory parentDir/study, invisibly.

See Also

```
seeker()
```

Examples

```
## Not run:
seekerArray('GSE25585', 'entrez')
## End(Not run)
```

trimgalore 17

|--|

Description

This function calls trim_galore using system2(), and is only designed to handle standard adapter/quality trimming. To run in parallel, register a parallel backend, e.g., using doParallel::registerDoParallel().

Usage

```
trimgalore(
  filepaths,
  outputDir = "trimgalore_output",
  cmd = "trim_galore",
  args = NULL,
  pigzCmd = "pigz"
)
```

Arguments

filepaths	Paths to fastq files. For single-end reads, each element should be a single filepath. For paired-end reads, each element should be two filepaths separated by ";".
outputDir	Directory in which to store output. Will be created if it doesn't exist.
cmd	Name or path of the command-line interface.
args	Additional arguments to pass to the command-line interface. Output files will always be compressed. Arguments "-gzip", "-cores", "-j", and "-basename" are not allowed. Arguments "-o" and "-paired" should not be specified here.
pigzCmd	String for pigz command, which will gzip the output files.

Value

A vector of exit codes, invisibly.

```
seeker()
```

18 tximport

tximport

Run tximport on RNA-seq quantifications

Description

This function uses the tximport package.

Usage

```
tximport(
  inputDir,
  tx2gene,
  samples = NULL,
  outputDir = "data",
  type = c("salmon", "kallisto"),
  countsFromAbundance = "lengthScaledTPM",
  ignoreTxVersion = TRUE,
  ...
)
```

Arguments

```
Directory that contains the quantification directories.
inputDir
tx2gene
                 NULL or data.frame of mapping between transcripts and genes, as returned by
                  getTx2gene(), passed to tximport::tximport().
samples
                  Names of quantification directories to include. NULL indicates all.
outputDir
                 Directory in which to save the result, a file named "tximport_output.qs", using
                  qs::qsave(). If NULL, no file is saved.
type
                  Passed to tximport::tximport().
countsFromAbundance
                  Passed to tximport::tximport().
ignoreTxVersion
                 Passed to tximport::tximport().
                 Additional arguments passed to tximport::tximport().
```

Value

```
A list, as returned by tximport::tximport(), invisibly.
```

```
seeker(), getTx2gene()
```

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