

# Package ‘TENET’

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**Title** R package for TENET (Tracing regulatory Element Networks using Epigenetic Traits) to identify key transcription factors

## Description

TENET identifies key transcription factors (TFs) and regulatory elements (REs) linked to a specific cell type by finding significantly correlated differences in gene expression and RE methylation between case and control input datasets, and identifying the top genes by number of significant RE DNA methylation site links. It also includes many additional tools to aid in visualization and analysis of the results, including plots displaying and comparing methylation and expression data and RE DNA methylation site link counts, survival analysis, TF motif searching in the vicinity of linked RE DNA methylation sites, custom TAD and peak overlap analysis, and UCSC Genome Browser track file generation. A utility function is also provided to download methylation, expression, and patient survival data from The Cancer Genome Atlas (TCGA) for use in TENET or other analyses.

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**Author** Rhie Lab at the University of Southern California [cre],  
Daniel Mullen [aut] (ORCID: <<https://orcid.org/0000-0002-7639-0549>>),  
Zexun Wu [aut] (ORCID: <<https://orcid.org/0000-0003-2566-1326>>),  
Ethan Nelson-Moore [aut] (ORCID:  
<<https://orcid.org/0009-0001-6903-9232>>),  
Suhn Rhie [aut] (ORCID: <<https://orcid.org/0000-0002-5522-5296>>)

**Maintainer** Rhie Lab at the University of Southern California <[rhie@ucla.edu](mailto:rhie@ucla.edu)>

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easyTENET	<i>Run the step 1 through step 6 functions with default arguments</i>
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**Description**

This function combines the step 1 through step 6 functions (step1MakeExternalDatasets, step2GetDifferentiallyMethylation, step3GetAnalysisZScores, step4SelectMostSignificantLinksPerDNAMethylationSite, step5OptimizeLinks, and step6DNAMethylationSitesPerGeneTabulation) into a single function. Arguments for this function generally reflect the arguments of the component functions without clearly defined defaults, with the exception of the step1MakeExternalDatasets function where all arguments have been included to support all of the options available to the user to define regions with relevant regulatory elements. All remaining arguments of the component functions are set to their default values.

**Usage**

```
easyTENET(  
  TENETMultiAssayExperiment,  
  extHM = NA,  
  extNDR = NA,  
  consensusEnhancer = TRUE,  
  consensusPromoter = FALSE,  
  consensusNDR = TRUE,  
  publicEnhancer = FALSE,  
  publicPromoter = FALSE,  
  publicNDR = FALSE,  
  cancerType = NA,  
  ENCODEPLS = FALSE,  
  ENCODEpELS = FALSE,  
  ENCODEdELS = FALSE,  
  assessPromoter = FALSE,  
  TSSDist = 1500,  
  minCaseCount,  
  coreCount = 1  
)
```

**Arguments**

TENETMultiAssayExperiment  
Specify a MultiAssayExperiment object containing expression and methylation SummarizedExperiment objects, such as one created by the TCGADownloader function. Coordinates for genes and DNA methylation sites should be included in the rowRanges of their respective SummarizedExperiment objects and should be annotated to the human hg38 genome. An argument of all functions except step1MakeExternalDatasets.

extHM	To use custom histone modification datasets, specify a path to a directory containing .bed, .narrowPeak, .broadPeak, and/or .gappedPeak files with these datasets. The files may optionally be compressed (.gz/.bz2/.xz). Otherwise, specify NA or do not specify this argument. An argument of the step1MakeExternalDatasets function.
extNDR	To use custom open chromatin or NDR datasets, specify a path to a directory containing .bed, .narrowPeak, .broadPeak, and/or .gappedPeak files with these datasets. The files may optionally be compressed (.gz/.bz2/.xz). Otherwise, specify NA or do not specify this argument. An argument of the step1MakeExternalDatasets function.
consensusEnhancer	Set to TRUE to use the consensus enhancer data included in TENET.AnnotationHub. Defaults to TRUE. An argument of the step1MakeExternalDatasets function.
consensusPromoter	Set to TRUE to use the consensus promoter data included in TENET.AnnotationHub. Defaults to FALSE. An argument of the step1MakeExternalDatasets function.
consensusNDR	Set to TRUE to use the consensus open chromatin data included in TENET.AnnotationHub. Defaults to TRUE. An argument of the step1MakeExternalDatasets function.
publicEnhancer	Set to TRUE to use the preprocessed publicly available enhancer (H3K27ac) datasets included in TENET.AnnotationHub. If set to TRUE, cancerType must be specified. Defaults to FALSE. An argument of the step1MakeExternalDatasets function.
publicPromoter	Set to TRUE to use the preprocessed publicly available promoter (H3K4me3) datasets included in TENET.AnnotationHub. If set to TRUE, cancerType must be specified. Defaults to FALSE. An argument of the step1MakeExternalDatasets function.
publicNDR	Set to TRUE to use the preprocessed publicly available open chromatin (ATAC-seq, DNase-seq) datasets included in TENET.AnnotationHub. If set to TRUE, cancerType must be specified. Defaults to FALSE. An argument of the step1MakeExternalDatasets function.
cancerType	If publicEnhancer, publicPromoter, and/or publicNDR is TRUE, specify a vector of cancer types from 'BLCA', 'BRCA', 'COAD', 'ESCA', 'HNSC', 'KIRP', 'LIHC', 'LUAD', 'LUSC', and 'THCA' to include the public data relevant to those cancer types. Defaults to NA. An argument of the step1MakeExternalDatasets function.
ENCODEPLS	Set to TRUE to use the ENCODE promoter-like elements dataset included in TENET.AnnotationHub. Defaults to FALSE. An argument of the step1MakeExternalDatasets function.
ENCODEpELS	Set to TRUE to use the ENCODE proximal enhancer-like elements dataset included in TENET.AnnotationHub. Defaults to FALSE. An argument of the step1MakeExternalDatasets function.
ENCODEdELS	Set to TRUE to use the ENCODE distal enhancer-like elements dataset included in TENET.AnnotationHub. Defaults to FALSE. An argument of the step1MakeExternalDatasets function.

assessPromoter	Set to TRUE to identify DNA methylation sites that mark promoter regions or FALSE to identify distal enhancer regions. Defaults to FALSE. An argument of the step2GetDifferentiallyMethylatedSites function.
TSSDist	Specify a positive integer distance in base pairs to any transcription start site within which DNA methylation sites are considered promoter DNA methylation sites. DNA methylation sites outside of the TSSDist from any transcription start site will be considered enhancer methylation sites. Defaults to 1500. An argument of the step2GetDifferentiallyMethylatedSites function.
minCaseCount	Specify a positive integer to be the minimum number of case samples to be considered for the hyper- or hypomethylated groups. Should be less than the total number of case samples. An argument of the step2GetDifferentiallyMethylatedSites function.
coreCount	Argument passed as the mc.cores argument to mclapply. See ?parallel::mclapply for more details. Defaults to 1. Used by the step3GetAnalysisZScores, step4SelectMostSignificantLinksPerDNAMethylationSite, and step5OptimizeLinks functions.

### Value

Returns the created MultiAssayExperiment object containing data from all step 1 through 6 functions.

### Examples

```
## This example creates a dataset of putative enhancer regulatory elements
## from consensus datasets and breast invasive carcinoma-relevant sources
## collected in the TENET.AnnotationHub package, then runs the step 2 through
## step 6 TENET functions analyzing RE DNA methylation sites in potential
## enhancer elements located over 1500 bp from transcription start sites
## listed for genes and transcripts in the GENCODE v36 human genome
## annotations, using a minimum case sample count of 5 and one CPU core
## to perform the analysis.

## Load the example TENET MultiAssayExperiment object from the
## TENET.ExperimentHub package
exampleTENETMultiAssayExperiment <-
  TENET.ExperimentHub::exampleTENETMultiAssayExperiment()

## Use the example TENET MultiAssayExperiment to run the step 1 through
## step 6 TENET functions
returnValue <- easyTENET(
  TENETMultiAssayExperiment = exampleTENETMultiAssayExperiment,
  extHM = NA,
  extNDR = NA,
  publicEnhancer = TRUE,
  publicNDR = TRUE,
  cancerType = "BRCA",
  ENCODEdELS = TRUE,
  minCaseCount = 5
)
```

```
## This example creates a dataset of putative promoter regulatory elements
## using bed-like files contained in the user's working directory, consensus
## NDR and promoter regions, and regions with promoter-like signatures from
## the ENCODE SCREEN project, but excluding cancer type-specific public
## datasets. This dataset is then used to analyze DNA methylation sites in
## promoter elements within 2000 bp of all transcription start sites
## provided in the MultiAssayExperiment only, identifying alterations found
## in at least 10 samples, and using 8 CPU cores to perform the analysis.

## Load the example TENET MultiAssayExperiment object from the
## TENET.ExperimentHub package
exampleTENETMultiAssayExperiment <-
  TENET.ExperimentHub::exampleTENETMultiAssayExperiment()

## Use the example TENET MultiAssayExperiment to run the step 1 through
## step 6 TENET functions
returnValue <- easyTENET(
  TENETMultiAssayExperiment = exampleTENETMultiAssayExperiment,
  extHM = ".",
  extNDR = ".",
  consensusEnhancer = FALSE,
  consensusPromoter = TRUE,
  ENCODEPLS = TRUE,
  assessPromoter = TRUE,
  TSSDist = 2000,
  minCaseCount = 10,
  coreCount = 8
)
```

---

humanTranscriptionFactorList

*Human transcription factor list*

---

## Description

A character vector of gene Ensembl IDs which were identified as human TFs by Lambert SA et al (PMID: 29425488). Candidate proteins were manually examined by a panel of experts based on available data. Proteins with experimentally demonstrated DNA binding specificity were considered TFs. Other proteins, such as co-factors and RNA binding proteins, were classified as non-TFs. **Citation:** Lambert SA, Jolma A, Campitelli LF, et al. The Human Transcription Factors. Cell. 2018 Feb 8;172(4):650-665. doi: 10.1016/j.cell.2018.01.029. Erratum in: Cell. 2018 Oct 4;175(2):598-599. PMID: 29425488.

## Usage

```
data("humanTranscriptionFactorList", package = "TENET")
```

**Format**

A character vector containing 1,639 Ensembl IDs of known human TFs.

**Source**

<http://humantfs.ccb.utoronto.ca/download.php>

**Examples**

```
data("humanTranscriptionFactorList", package = "TENET")
```

---

step1MakeExternalDatasets

*Create a GRanges object representing putative regulatory element regions, based on the data sources selected for inclusion, to be used in later TENET steps*

---

**Description**

This function creates a GRanges object containing regions representing putative regulatory elements, either enhancers or promoters, of interest to the user, based on the presence of specific histone marks and open chromatin/nucleosome-depleted regions. This function can take input from user-specified bed-like files (see <https://genome.ucsc.edu/FAQ/FAQformat.html#format1>) containing regions with histone modification (via the extHM argument) and/or open chromatin/nucleosome-depleted regions (via the extNDR argument), as well as preprocessed enhancer, promoter, and open chromatin datasets from many cell/tissue types included in the TENET.AnnotationHub repository. The resulting GRanges object will be returned. GRanges objects created by this function can be used by the step2GetDifferentiallyMethylatedSites function or other downstream functions. **Note:** Using datasets from TENET.AnnotationHub requires an internet connection, as those datasets are hosted in the Bioconductor AnnotationHub Data Lake.

**Usage**

```
step1MakeExternalDatasets(  
  extHM = NA,  
  extNDR = NA,  
  consensusEnhancer = TRUE,  
  consensusPromoter = FALSE,  
  consensusNDR = TRUE,  
  publicEnhancer = FALSE,  
  publicPromoter = FALSE,  
  publicNDR = FALSE,  
  cancerType = NA,  
  ENCODEPLS = FALSE,  
  ENCODEpELS = FALSE,  
  ENCODEdELS = FALSE  
)
```

**Arguments**

extHM	To use custom histone modification datasets, specify a path to a directory containing .bed, .narrowPeak, .broadPeak, and/or .gappedPeak files with these datasets. The files may optionally be compressed (.gz/.bz2/.xz). Otherwise, specify NA or do not specify this argument.
extNDR	To use custom open chromatin or NDR datasets, specify a path to a directory containing .bed, .narrowPeak, .broadPeak, and/or .gappedPeak files with these datasets. The files may optionally be compressed (.gz/.bz2/.xz). Otherwise, specify NA or do not specify this argument.
consensusEnhancer	Set to TRUE to use the consensus enhancer data included in TENET.AnnotationHub. Defaults to TRUE.
consensusPromoter	Set to TRUE to use the consensus promoter data included in TENET.AnnotationHub. Defaults to FALSE.
consensusNDR	Set to TRUE to use the consensus open chromatin data included in TENET.AnnotationHub. Defaults to TRUE.
publicEnhancer	Set to TRUE to use the preprocessed publicly available enhancer (H3K27ac) datasets included in TENET.AnnotationHub. If set to TRUE, cancerType must be specified. Defaults to FALSE.
publicPromoter	Set to TRUE to use the preprocessed publicly available promoter (H3K4me3) datasets included in TENET.AnnotationHub. If set to TRUE, cancerType must be specified. Defaults to FALSE.
publicNDR	Set to TRUE to use the preprocessed publicly available open chromatin (ATAC-seq, DNase-seq) datasets included in TENET.AnnotationHub. If set to TRUE, cancerType must be specified. Defaults to FALSE.
cancerType	If publicEnhancer, publicPromoter, and/or publicNDR is TRUE, specify a vector of cancer types from 'BLCA', 'BRCA', 'COAD', 'ESCA', 'HNSC', 'KIRP', 'LIHC', 'LUAD', 'LUSC', and 'THCA' to include the public data relevant to those cancer types. Defaults to NA.
ENCODEPLS	Set to TRUE to use the ENCODE promoter-like elements dataset included in TENET.AnnotationHub. Defaults to FALSE.
ENCODEpELS	Set to TRUE to use the ENCODE proximal enhancer-like elements dataset included in TENET.AnnotationHub. Defaults to FALSE.
ENCODEdELS	Set to TRUE to use the ENCODE distal enhancer-like elements dataset included in TENET.AnnotationHub. Defaults to FALSE.

**Value**

Returns the created regulatory element GRanges object.

**Examples**

```
## This example creates a dataset of putative enhancer regulatory elements
## from consensus datasets and breast invasive carcinoma-relevant sources
## collected in the TENET.AnnotationHub package.
```



```

returnGRanges <- step1MakeExternalDatasets(
  extHM = NA,
  extNDR = NA,
  publicEnhancer = TRUE,
  publicNDR = TRUE,
  cancerType = "BRCA",
  ENCODEdELS = TRUE
)

## This example creates a dataset of putative promoter regulatory elements
## using user provided bed-like files contained in the working
## directory, consensus NDR and promoter regions, and regions with
## promoter-like signatures from the ENCODE SCREEN project. This excludes any
## cancer type-specific public datasets.
returnGRanges <- step1MakeExternalDatasets(
  extHM = ".",
  extNDR = ".",
  consensusEnhancer = FALSE,
  consensusPromoter = TRUE,
  ENCODEPLS = TRUE
)

```

---

step2GetDifferentiallyMethylatedSites

*Identify differentially methylated RE DNA methylation sites*

---

## Description

This function identifies DNA methylation sites that mark putative regulatory elements (REs), including enhancer and promoter regions. These are sites that lie within regions with specific histone modifications and open chromatin regions, from a user-supplied GRanges object, such as one created by the `step1MakeExternalDatasets` function, and which are located at a user-specified distance relative to the transcription start sites (TSS) listed in either the `rowRanges` of the `element-Metadata` of the "expression" `SummarizedExperiment` in the `TENETMultiAssayExperiment` object, or the selected `geneAnnotationDataset` (which will be filtered to only genes and transcripts). After identifying DNA methylation sites representing the specified REs, the function classifies the RE DNA methylation sites as methylated, unmethylated, hypermethylated, or hypomethylated based on their differential methylation between the control and case samples supplied by the user, defined by cutoff values which are either automatically based on the mean methylation densities of the identified RE DNA methylation sites, or manually set by the user. **Note:** Using the algorithm to set cutoffs is recommended for use with DNA methylation array data, and may not work for whole-genome DNA methylation data.

## Usage

```

step2GetDifferentiallyMethylatedSites(
  TENETMultiAssayExperiment,
  regulatoryElementGRanges = NA,

```

```

geneAnnotationDataset = NA,
DNAMethylationArray = NA,
assessPromoter = FALSE,
TSSDist = 1500,
purityData = NA,
methCutoff = NA,
hypomethCutoff = NA,
hypermethCutoff = NA,
unmethCutoff = NA,
methUnmethProportionOffset = 0.2,
hypomethHypermethProportionOffset = 0.1,
minCaseCount,
cgDNAMethylationSitesOnly = TRUE
)

```

## Arguments

**TENETMultiAssayExperiment**

Specify a MultiAssayExperiment object containing expression and methylation SummarizedExperiment objects, such as one created by the TCGADownloader function. Coordinates for genes and DNA methylation sites should be included in the rowRanges of their respective SummarizedExperiment objects and should be annotated from the same genome build as the regions given in the regulatoryElementGRanges object.

**regulatoryElementGRanges**

Specify a GRanges object containing genomic regions representing regulatory elements of interest to the user. Coordinates for the regulatory element regions should be annotated to the same genome build as the gene and DNA methylation site coordinates given in the TENETMultiAssayExperiment object. If this argument is set to NA or not specified, this function will use all DNA methylation sites representing regulatory elements of interest as defined by the assessPromoter and TSSDist arguments. Defaults to NA.

**geneAnnotationDataset**

Specify a gene annotation dataset which is used to identify transcription start sites in order to find DNA methylation sites within regulatory elements of interest (promoters or enhancers) in conjunction with the settings of the assessPromoter and TSSDist arguments. The dataset will be filtered to only genes and transcripts. The argument must be either a GRanges object (such as one imported via `rtracklayer::import`) or a path to a GFF3 or GTF file. Both GENCODE and Ensembl annotations are supported. Other annotation datasets may work, but have not been tested. See the "Input data" section of the vignette for information on the required dataset format. Specify NA to use the start coordinates of all entries in the elementMetadata of the rowRanges of the "expression" SummarizedExperiment object within the TENETMultiAssayExperiment object, in which case no filtering will be done and all entries will be assumed to represent transcripts. Defaults to NA.

**DNAMethylationArray**

Specify the name of a DNA methylation probe array supported by the sesameData package (see `?sesameData::sesameData_getManifestGRanges`). If an

array is specified, RE DNA methylation sites and their locations in that array's manifest are cross-referenced with RE DNA methylation site IDs included in the rownames of the methylation dataset provided in the "methylation" SummarizedExperiment object within the TENETMultiAssayExperiment object, and only those overlapping will be considered for analysis. If set to NA, all RE DNA methylation sites with locations listed in the rowRanges of the "methylation" SummarizedExperiment object are used. Defaults to NA.

assessPromoter	Set to TRUE to identify DNA methylation sites that mark promoter regions or FALSE to identify distal enhancer regions. Defaults to FALSE.
TSSDist	Specify a positive integer distance in base pairs to any transcription start site (see geneAnnotationDataset) within which DNA methylation sites are considered promoter DNA methylation sites. DNA methylation sites outside of the TSSDist from any transcription start site will be considered enhancer methylation sites. Defaults to 1500.
purityData	Specify a SummarizedExperiment object which contains DNA methylation datasets collected from potential cell types which might affect the purity of the patient samples contained in the TENETMultiAssayExperiment. The coordinates for DNA methylation sites in this dataset should be included in the rowRanges of the purityData SummarizedExperiment object. Additionally, the DNA methylation site IDs in the purityData SummarizedExperiment object should overlap with DNA methylation sites present in the TENETMultiAssayExperiment and only those that do overlap will be considered for analysis. Defaults to NA.
methCutoff	Specify a number from 0 to 1 to be the beta-value cutoff for methylated RE DNA methylation sites. If unspecified or NA, an algorithm will be used to find the optimal cutoff value.
hypomethCutoff	Specify a number from 0 to 1 to be the beta-value cutoff for hypomethylated RE DNA methylation sites. Should be set lower than the methCutoff. If unspecified or NA, an algorithm will be used to find the optimal cutoff value.
hypermethCutoff	Specify a number from 0 to 1 to be the beta-value cutoff for hypermethylated RE DNA methylation sites. Should be set higher than the unmethCutoff. If unspecified or NA, an algorithm will be used to find the optimal cutoff value.
unmethCutoff	Specify a number from 0 to 1 to be the beta-value cutoff for unmethylated RE DNA methylation sites. If unspecified or NA, an algorithm will be used to find the optimal cutoff value.
methUnmethProportionOffset	Specify a number from 0 to 1 to be the proportion of the distance of the region between the first and last local maxima in the density plot of the mean methylation values of the RE DNA methylation sites in the control samples. This value is then added to the position of these local maxima to set the unmethylation and methylation cutoffs if they are not defined by the user. The value ideally should not exceed 0.5. Defaults to 0.2.
hypomethHypermethProportionOffset	Specify a number from 0 to 1 to be the proportion of the distance of the region between the first and last local maxima in the density plot of the mean methylation values of the RE DNA methylation sites in the case samples. This value is

then added to the calculated unmethylation and methylation cutoffs to then set the hypermethylation and hypomethylation cutoffs if they are not defined by the user. The value ideally should not exceed 0.5. Defaults to 0.1.

**minCaseCount** Specify a positive integer to be the minimum number of case samples to be considered for the hyper- or hypomethylated groups. Should be less than the total number of case samples.

**cgDNAMethylationSitesOnly**

Set to TRUE to include only RE DNA methylation sites with IDs that start with "cg". TRUE means that RE DNA methylation sites whose IDs do not start with "cg" will be removed from TENET analyses. Defaults to TRUE.

## Value

Returns the MultiAssayExperiment object given as the TENETMultiAssayExperiment argument with an additional list of data named "step2GetDifferentiallyMethylatedSites" in its metadata with the output data from this function. These data include the set of calculated cutoff values, the identities and counts of the classified RE DNA methylation sites, as well as plots of the mean methylation distributions of the identified regulatory element DNA methylation sites in the case and control samples and the set cutoff values. Of note for plots, if assessPromoter is TRUE, two distribution plots are saved, one using all promoter DNA methylation sites, and one using DNA methylation sites which are identified to overlap REs.

## Examples

```
## This example uses datasets provided in the TENET.ExperimentHub package to
## perform an example analysis, analyzing RE DNA methylation sites in
## potential enhancer elements, located over 1500 bp from transcription
## start sites listed for genes and transcripts in the GENCODE v36 human
## genome annotations, otherwise using default settings and a minimum case
## sample count of 5.
```

```
## Load the example TENET MultiAssayExperiment object, and the example
## GRanges object created by the TENET step 1 function, from the
## TENET.ExperimentHub package
exampleTENETMultiAssayExperiment <-
  TENET.ExperimentHub::exampleTENETMultiAssayExperiment()
exampleStep1MakeExternalDatasetsGRangesObject <-
  TENET.ExperimentHub::exampleTENETStep1MakeExternalDatasetsGRanges()
```

```
## Use the example datasets to identify differentially methylated
## RE DNA methylation sites
returnValue <- step2GetDifferentiallyMethylatedSites(
  TENETMultiAssayExperiment = exampleTENETMultiAssayExperiment,
  regulatoryElementGRanges =
    exampleTENETStep1MakeExternalDatasetsGRanges,
  minCaseCount = 5
)
```

```
## This example uses the same datasets, this time analyzing DNA methylation
## sites in promoter elements, considering all RE DNA methylation sites
## found within 2000 bp of all transcription start sites provided in the
```

```

## MultiAssayExperiment only. Additionally, the methylation cutoffs are
## manually set to 0.8, 0.7, 0.3, and 0.2 for the `methCutoff`,
## `hypomethCutoff`, `hypermethCutoff`, and `unmethCutoff` respectively. The
## `minCaseCount` is set to 10 samples and all RE DNA methylation sites
## regardless of ID will be considered.

## Load the example TENET MultiAssayExperiment object
## from the TENET.ExperimentHub package as well as the example GRanges object
## created by the TENET step 1 function from the TENET.ExperimentHub package
exampleTENETMultiAssayExperiment <-
  TENET.ExperimentHub::exampleTENETMultiAssayExperiment()
exampleStep1MakeExternalDatasetsGRangesObject <-
  TENET.ExperimentHub::exampleTENETStep1MakeExternalDatasetsGRanges()

## Use the example datasets to identify differentially methylated
## RE DNA methylation sites
returnValue <- step2GetDifferentiallyMethylatedSites(
  TENETMultiAssayExperiment = exampleTENETMultiAssayExperiment,
  regulatoryElementGRanges =
    exampleTENETStep1MakeExternalDatasetsGRanges,
  geneAnnotationDataset = NA,
  assessPromoter = TRUE,
  TSSDist = 2000,
  methCutoff = 0.8,
  hypomethCutoff = 0.7,
  hypermethCutoff = 0.3,
  unmethCutoff = 0.2,
  minCaseCount = 10,
  cgDNAMethylationSitesOnly = FALSE
)

```

---

**step3GetAnalysisZScores**

*Calculate Z-scores comparing the mean expression of each gene in the case samples that are hyper- or hypomethylated for each RE DNA methylation site chosen in step 2*

---

**Description**

This function takes the identified hyper- and hypomethylated RE DNA methylation sites from `step2GetDifferentiallyMethylatedSites` function and calculates Z-scores comparing the mean expression of each gene in the case samples that are hyper- or hypomethylated for each RE DNA methylation site, according to hyper- and hypomethylation cutoffs set during the previous function, to those that are not, across all hyper- or hypomethylated RE DNA methylation sites, calculating Z-scores for each unique RE DNA methylation site and gene combination, also known as a link.

**Usage**

```
step3GetAnalysisZScores(
  TENETMultiAssayExperiment,
  hypermethAnalysis = TRUE,
  hypomethAnalysis = TRUE,
  includeControl = FALSE,
  TFOnly = TRUE,
  zScoreCalculation = "oneSample",
  sparseResults = TRUE,
  pValue = 0.05,
  coreCount = 1
)
```

**Arguments**

TENETMultiAssayExperiment	Specify a MultiAssayExperiment object containing expression and methylation SummarizedExperiment objects, such as one created by the TCGADownloader function. This MultiAssayExperiment object should also contain the results from the step2GetDifferentiallyMethylatedSites function in the metadata of that object.
hypermethAnalysis	Set to TRUE to calculate Z-scores for hypermethylated RE DNA methylation sites. Defaults to TRUE.
hypomethAnalysis	Set to TRUE to calculate Z-scores for hypomethylated RE DNA methylation sites. Defaults to TRUE.
includeControl	Set to TRUE to include the control samples when identifying hyper/hypomethylated groups and calculating Z-scores. Defaults to FALSE.
TFOnly	Set to TRUE to only consider genes that are accepted transcription factors in "The Human Transcription Factors" by Lambert et al. 2018 when calculating Z-scores. Defaults to TRUE.
zScoreCalculation	Set to 'oneSample' to use a one-sample Z-score calculation or 'twoSample' to use a two sample Z-score calculation. Note that 'twoSample' tends to be much more lenient, and identifies many more significant RE DNA methylation site-gene links. Defaults to 'oneSample'.
sparseResults	Set to TRUE to save only the significant Z-scores for RE DNA methylation site-gene links. <b>Note:</b> If multiple testing correction will be performed in the subsequent step4SelectMostSignificantLinksPerDNAMethylationSite function, this argument should be set to FALSE. Defaults to TRUE.
pValue	Specify the p-value below which Z-scores will be considered significant during comparison of gene expression values between case samples that are hyper- or hypomethylated and those that are not. If sparseResults is set to TRUE, only significant Z-scores will be saved in the output MultiAssayExperiment object. Defaults to 0.05.
coreCount	Argument passed as the mc.cores argument to mclapply. See ?parallel::mclapply for more details. Defaults to 1.

**Value**

Returns the MultiAssayExperiment object given as the TENETMultiAssayExperiment argument with an additional list of data named "step3GetAnalysisZScores" in its metadata with the output of this function, which includes Z-scores for each TF or gene analyzed to each identified hyper- and/or hypomethylated RE DNA methylation site.

**Examples**

```
## This example uses the example MultiAssayExperiment provided in the
## TENET.ExperimentHub package to calculate one-sample Z-scores for links
## between both the hypermethylated and hypomethylated RE DNA methylation
## sites and the expression of transcription factor genes only, considering
## only the expression/methylation of case samples. Only significant
## Z-scores equivalent to  $p < 0.05$  will be saved to the
## TENETMultiAssayExperiment object. The analysis will be performed using
## one CPU core.

## Load the example TENET MultiAssayExperiment object
## from the TENET.ExperimentHub package
exampleTENETMultiAssayExperiment <-
  TENET.ExperimentHub::exampleTENETMultiAssayExperiment()

## Calculate Z-scores for hyper- and hypomethylated RE DNA methylation sites
returnValue <- step3GetAnalysisZScores(
  TENETMultiAssayExperiment = exampleTENETMultiAssayExperiment
)

## This example also uses the example MultiAssayExperiment, but calculates
## two-sample Z-scores for links between only hypomethylated RE DNA
## methylation sites and all genes, considering the expression/methylation
## of both case and control samples. For this analysis, all Z-scores are
## saved to the TENETMultiAssayExperiment object (though it should be noted
## this takes a large amount of memory). Only Z-scores with p-values less
## than 0.1 will be considered significant. This analysis will be performed
## using 8 CPU cores.

## Load the example TENET MultiAssayExperiment object
## from the TENET.ExperimentHub package
exampleTENETMultiAssayExperiment <-
  TENET.ExperimentHub::exampleTENETMultiAssayExperiment()

## Calculate Z-scores for hyper- and hypomethylated RE DNA methylation sites
returnValue <- step3GetAnalysisZScores(
  TENETMultiAssayExperiment = exampleTENETMultiAssayExperiment,
  hypermethAnalysis = FALSE,
  includeControl = TRUE,
  TFOnly = FALSE,
  zScoreCalculation = "twoSample",
  sparseResults = FALSE,
  pValue = 0.1,
  coreCount = 8
)
```

---

```
step4SelectMostSignificantLinksPerDNAMethylationSite
```

*Select the most significant RE DNA methylation site-gene links to each RE DNA methylation site*

---

## Description

This function takes the calculated Z-scores for the hyper- or hypomethylated G+ RE DNA methylation site-gene links and selects the most significant links to each regulatory element DNA methylation site, either up to a number specified by the user, or based on a significant p-value level set by the user after multiple testing correction is performed on the Z-scores output by the step3GetAnalysisZScores function per RE DNA methylation site in the RE DNA methylation site-gene pairs.

## Usage

```
step4SelectMostSignificantLinksPerDNAMethylationSite(
  TENETMultiAssayExperiment,
  hypermethGplusAnalysis = TRUE,
  hypomethGplusAnalysis = TRUE,
  linksPerREDNAMethylationSiteMaximum = 25,
  multipleTestingPValue = 0.05,
  coreCount = 1
)
```

## Arguments

TENETMultiAssayExperiment

Specify a MultiAssayExperiment object containing expression and methylation SummarizedExperiment objects, such as one created by the TCGADownloader function. This MultiAssayExperiment object should also contain the results from the step2GetDifferentiallyMethylatedSites and step3GetAnalysisZScores functions in its metadata.

hypermethGplusAnalysis

Set to TRUE to analyze hypermethylated G+ RE DNA methylation site-gene links. Requires the hypermethAnalysis parameter to have been set to TRUE in step 3.

hypomethGplusAnalysis

Set to TRUE to analyze hypomethylated G+ RE DNA methylation site-gene links. Requires the hypomethAnalysis parameter to have been set to TRUE in step 3.

linksPerREDNAMethylationSiteMaximum

This parameter must either be set to an integer n greater than 0, in which case only the n most significant RE DNA methylation site-gene link pairs from step 3 will be selected per RE DNA methylation site, or a character string describing a multiple testing correction method supported by p.adjust (see ?stats::p.adjust)



to perform multiple testing correction on the Z-scores from step 3, using the `multipleTestingPValue` argument to set a significant p-value cutoff. **Note:** If multiple testing correction is performed, `sparseResults` should have been set to `FALSE` in the `step3GetAnalysisZScores` function. Defaults to 25 (maximum links per unique RE DNA methylation site).

`multipleTestingPValue`

Cutoff for multiple testing corrected p-values. This argument is only used if the `linksPerREDNAMethylationSiteMaximum` argument is set to a multiple testing correction method. Defaults to 0.05.

`coreCount`

Argument passed as the `mc.cores` argument to `mclapply`. See `?parallel::mclapply` for more details. Defaults to 1.

## Value

Returns the `MultiAssayExperiment` object given as the `TENETMultiAssayExperiment` argument with an additional list of data named "step4SelectMostSignificantLinksPerDNAMethylationSite" in its metadata with the output of this function, which includes the most significant selected gene links to the hyper- or hypomethylated RE DNA methylation sites.

## Examples

```
## This example uses the example MultiAssayExperiment provided in the
## TENET.ExperimentHub package to identify the 25 most significant links
## between both hyper- and hypomethylated enhancer DNA methylation sites and
## all genes, using one CPU core to perform the analysis.

## Load the example TENET MultiAssayExperiment object
## from the TENET.ExperimentHub package
exampleTENETMultiAssayExperiment <-
  TENET.ExperimentHub::exampleTENETMultiAssayExperiment()

## Perform the analysis
returnValue <- step4SelectMostSignificantLinksPerDNAMethylationSite(
  TENETMultiAssayExperiment = exampleTENETMultiAssayExperiment
)

## This example also uses the example MultiAssayExperiment but identifies
## the most significant links between only hypomethylated enhancer DNA
## methylation sites and all genes by performing Bonferroni multiple testing
## correction using a significant p-value of 0.10, using 8 CPU cores to
## perform the analysis. Note: running this code with the
## exampleTENETMultiAssayExperiment will produce a warning message as
## sparseResults was set to TRUE when the example dataset was generated.
## However, this function will still run and is valid as an example.

## Load the example TENET MultiAssayExperiment object
## from the TENET.ExperimentHub package
exampleTENETMultiAssayExperiment <-
  TENET.ExperimentHub::exampleTENETMultiAssayExperiment()

## Perform the analysis
```

```

returnValue <- step4SelectMostSignificantLinksPerDNAMethylationSite(
  TENETMultiAssayExperiment = exampleTENETMultiAssayExperiment,
  hypermethGplusAnalysis = FALSE,
  linksPerREDNAMethylationSiteMaximum = "bonferroni",
  multipleTestingPValue = 0.1,
  coreCount = 8
)

```

---

step5OptimizeLinks	<i>Find final RE DNA methylation site-gene links using various optimization metrics</i>
--------------------	---

---

## Description

This function takes the most significant hyper- or hypomethylated G+ RE DNA methylation site-gene links selected in step 4, and selects optimized links based on the relative expression of the given gene in hyper- or hypomethylated case samples compared to control samples, using an unpaired two-sided Wilcoxon rank-sum test to check that the hyper- or hypomethylated samples for that given RE DNA methylation site-gene link also show appropriately higher/lower expression of the linked gene in a number of case samples greater than or equal to the minCaseCount number specified in the step2GetDifferentiallyMethylatedSites function that have maximum/minimum methylation above/below the hyperStringency/hypoStringency cutoff values selected.

## Usage

```

step5OptimizeLinks(
  TENETMultiAssayExperiment,
  hypermethGplusAnalysis = TRUE,
  hypomethGplusAnalysis = TRUE,
  expressionPvalCutoff = 0.05,
  hyperStringency = NA,
  hypoStringency = NA,
  coreCount = 1
)

```

## Arguments

TENETMultiAssayExperiment	Specify a MultiAssayExperiment object containing expression and methylation SummarizedExperiment objects, such as one created by the TCGADownloader function. This MultiAssayExperiment object should also contain the results from the step2GetDifferentiallyMethylatedSites, step3GetAnalysisZScores, and step4SelectMostSignificantLinksPerDNAMethylationSite functions in its metadata.
hypermethGplusAnalysis	Set to TRUE to optimize hypermethylated G+ links. Requires hypermethAnalysis from step 4 to have been set to TRUE.

hypomethGplusAnalysis	Set to TRUE to optimize hypomethylated G+ links. Requires hypomethAnalysis from step 4 to have been set to TRUE.
expressionPvalCutoff	Cutoff for Benjamini-Hochberg corrected Wilcoxon p-values used during comparison of gene expression values between hyper/hypomethylated case and control samples. Defaults to 0.05.
hyperStringency	Specify a number from 0 to 1 to be the beta-value cutoff to optimize for hypermethylated links with methylation values above the cutoff if a more/less selective cutoff is desired. Defaults to the hypermethCutoff value specified in step2GetDifferentiallyMethylatedSites.
hypoStringency	Specify a number from 0 to 1 to be the beta-value cutoff to optimize for hypomethylated links with methylation values below the cutoff if a more/less selective cutoff is desired. Defaults to the hypomethCutoff value specified in step2GetDifferentiallyMethylatedSites.
coreCount	Argument passed as the mc.cores argument to mcmapply. See ?parallel::mcmapply for more details. Defaults to 1.

## Value

Returns the MultiAssayExperiment object given as the TENETMultiAssayExperiment argument with an additional list of data named "step5OptimizeLinks" in its metadata with the output of this function, which includes the gene and RE DNA methylation site IDs of each optimized gene-RE DNA methylation site link, the Z-scores and corresponding p-values calculated in steps 3-4 for those links, and the various optimization metrics performed by this step.

## Examples

```
## This example uses the example MultiAssayExperiment provided in the
## TENET.ExperimentHub package to optimize links between the hypermethylated
## and hypomethylated RE DNA methylation sites and the expression of
## transcription factor genes. For this analysis, the default p-value cutoff
## of 0.05 is used, with `hyperStringency` and `hypoStringency` values set
## to the originally calculated hyper- and hypomethylation cutoffs. The
## analysis uses one CPU core.

## Load the example TENET MultiAssayExperiment object
## from the TENET.ExperimentHub package
exampleTENETMultiAssayExperiment <-
  TENET.ExperimentHub::exampleTENETMultiAssayExperiment()

## Perform the link optimization
returnValue <- step5OptimizeLinks(
  TENETMultiAssayExperiment = exampleTENETMultiAssayExperiment
)

## This example also uses the example MultiAssayExperiment provided in the
## TENET.ExperimentHub package, but it only runs on hypomethylated RE DNA
## methylation sites and uses a p-value cutoff of 0.01. The `hypoStringency`
```

```

## value is set to 0.3 specifically rather than using the originally set
## hypomethylation cutoff. Eight CPU cores are used to perform the analysis.

## Load the example TENET MultiAssayExperiment object
## from the TENET.ExperimentHub package
exampleTENETMultiAssayExperiment <-
  TENET.ExperimentHub::exampleTENETMultiAssayExperiment()

## Perform the link optimization
returnValue <- step5OptimizeLinks(
  TENETMultiAssayExperiment = exampleTENETMultiAssayExperiment,
  hypermethGplusAnalysis = FALSE,
  expressionPvalCutoff = 0.01,
  hypoStringency = 0.3,
  coreCount = 8
)

```

---

```
step6DNAMethylationSitesPerGeneTabulation
```

*Tabulate the total number of RE DNA methylation sites linked to each of the genes*

---

## Description

This function takes the final optimized regulatory element RE DNA methylation site-gene links identified by the `step5OptimizeLinks` function and tabulates the number of these links per gene. This tabulation is done separately for both of the hyper- or hypomethylated G+ analysis quadrants, as selected by the user.

## Usage

```

step6DNAMethylationSitesPerGeneTabulation(
  TENETMultiAssayExperiment,
  geneAnnotationDataset = NA,
  hypermethGplusAnalysis = TRUE,
  hypomethGplusAnalysis = TRUE
)

```

## Arguments

TENETMultiAssayExperiment

Specify a MultiAssayExperiment object containing expression and methylation SummarizedExperiment objects, such as one created by the `TCGADownloader` function. This MultiAssayExperiment object should also contain the results from the `step5OptimizeLinks` function in its metadata.

**geneAnnotationDataset**

Specify a gene annotation dataset which is used to identify names for genes by their Ensembl IDs. The argument must be either a GRanges object (such as one imported via `rtracklayer::import`) or a path to a GFF3 or GTF file. Both GENCODE and Ensembl annotations are supported. Other annotation datasets may work, but have not been tested. See the "Input data" section of the vignette for information on the required dataset format. Specify NA to use the names for genes listed in the "geneName" column of the elementMetadata of the rowRanges of the "expression" SummarizedExperiment object within the TENETMultiAssayExperiment object. Defaults to NA.

**hypermethGplusAnalysis**

Set to TRUE to calculate total links by gene for hypermethylated RE DNA methylation sites with G+ links. Defaults to TRUE.

**hypomethGplusAnalysis**

Set to TRUE to calculate total links by gene for hypomethylated RE DNA methylation sites with G+ links. Defaults to TRUE.

**Value**

Returns the MultiAssayExperiment object given as the TENETMultiAssayExperiment argument with an additional list of data named "step6DNAMethylationSitesPerGeneTabulation" in its meta-data with the output of this function, which includes data frames containing significant hyper- and/or hypomethylated G+ link counts per gene after all TENET steps through step5OptimizeLinks have been run.

**Examples**

```
## This example uses the example MultiAssayExperiment provided in the
## TENET.ExperimentHub package to tabulate both hyper- and hypomethylated G+
## RE DNA methylation site-gene links, using genes with names provided in the
## provided MultiAssayExperiment object.
```

```
## Load the example TENET MultiAssayExperiment object
## from the TENET.ExperimentHub package
exampleTENETMultiAssayExperiment <-
  TENET.ExperimentHub::exampleTENETMultiAssayExperiment()
```

```
## Calculate linked RE DNA methylation sites per gene
returnValue <- step6DNAMethylationSitesPerGeneTabulation(
  TENETMultiAssayExperiment = exampleTENETMultiAssayExperiment
)
```

```
## This example also uses the example MultiAssayExperiment provided
## in the TENET.ExperimentHub package, but it only runs on hypomethylated
## RE DNA methylation sites.
```

```
## Load the example TENET MultiAssayExperiment object
## from the TENET.ExperimentHub package
exampleTENETMultiAssayExperiment <-
  TENET.ExperimentHub::exampleTENETMultiAssayExperiment()
```

```
## Calculate linked RE DNA methylation sites per gene
returnValue <- step6DNAMethylationSitesPerGeneTabulation(
  TENETMultiAssayExperiment = exampleTENETMultiAssayExperiment,
  hypermethGplusAnalysis = FALSE
)
```

---

```
step7ExpressionVsDNAMethylationScatterplots
```

*Create scatterplots displaying the expression of the top genes and the methylation levels of each of their linked RE DNA methylation sites, along with copy number variation, somatic mutation, and purity data, if provided by the user*

---

## Description

This function takes the top genes/transcription factors (TFs) by number of linked RE DNA methylation sites identified by the `step6DNAMethylationSitesPerGeneTabulation` function up to the number specified by the user, or all genes linked to selected RE DNA methylation sites in a list specified by the user, and generates scatterplots displaying the expression level of each of these genes in the x-axis and the methylation level of each RE DNA methylation site linked to them for both of the hyper- or hypomethylated G+ analysis quadrants, as selected by the user. The scatterplots incorporate copy number variation (CNV), somatic mutation (SM), and purity information from each sample, if provided by the user.

## Usage

```
step7ExpressionVsDNAMethylationScatterplots(
  TENETMultiAssayExperiment,
  geneAnnotationDataset = NA,
  hypermethGplusAnalysis = TRUE,
  hypomethGplusAnalysis = TRUE,
  topGeneNumber = 10,
  DNAMethylationSites = NA,
  simpleOrComplex = "simple",
  CNVData = NA,
  SMDData = NA,
  purityData = NA,
  coreCount = 1
)
```

## Arguments

TENETMultiAssayExperiment

Specify a MultiAssayExperiment object containing expression and methylation SummarizedExperiment objects, such as one created by the `TCGADownloader` function. This MultiAssayExperiment object should also contain the results from the `step5OptimizeLinks` and `step6DNAMethylationSitesPerGeneTabulation` functions in its metadata.

geneAnnotationDataset	Specify a gene annotation dataset which is used to identify names for genes by their Ensembl IDs. The argument must be either a GRanges object (such as one imported via <code>rtracklayer::import</code> ) or a path to a GFF3 or GTF file. Both GENCODE and Ensembl annotations are supported. Other annotation datasets may work, but have not been tested. See the "Input data" section of the vignette for information on the required dataset format. Specify NA to use the names for genes listed in the "geneName" column of the elementMetadata of the rowRanges of the "expression" SummarizedExperiment object within the TENETMultiAssayExperiment object. Defaults to NA.
hypermethGplusAnalysis	Set to TRUE to create scatterplots for genes with hypermethylated RE DNA methylation sites with G+ links and each of their linked RE DNA methylation sites. Defaults to TRUE.
hypomethGplusAnalysis	Set to TRUE to create scatterplots for genes with hypomethylated RE DNA methylation sites with G+ links and each of their linked RE DNA methylation sites. Defaults to TRUE.
topGeneNumber	Specify the number of top genes/TFs, based on the most linked RE DNA methylation sites of a given analysis type, for which to create scatterplots. Defaults to 10.
DNAMethylationSites	Supply a vector of RE DNA methylation site IDs for which scatterplots of those RE DNA methylation sites with expression of any linked genes/TFs of the specified analysis types will be generated.
simpleOrComplex	Set to 'simple' to create scatterplots without using copy number variation, somatic mutation, or purity data from samples. Otherwise set to 'complex' to use such data in the scatterplots. If set to 'complex', data on the samples' copy number variation, somatic mutation data, and purity data will need to be provided to the subsequent CNVData, SMDData, and purityData arguments respectively. Defaults to 'simple'.
CNVData	Specify the CNV status for each of the top genes and each sample in the TENETMultiAssayExperiment. CNV status should be given in the form of an integer value with a minimum of -2, representing the change in copy number for each of the top genes, by the 'hypermethGplusAnalysis', 'hypomethGplusAnalysis', and 'topGeneNumber' settings, with -2 representing a loss of both copies, -1 a single copy loss, 0 for no copy number change, and positive values indicating a gain of that many copies, although copy number gains of 2 or more will be grouped together. These data can be given in the form of a data frame/matrix, or a path to a file that contains the vital status data. If a data frame or matrix is given, then the rownames of the supplied data frame or matrix must include the sample names as they appear in the colData of the TENETMultiAssayExperiment object. If a single string is provided, then it is assumed to be a path to a file containing tab-delimited CNV data. It is expected that the names of the samples, again corresponding with the sample names in the colData, are given in the first column of the file, which will be loaded as row names. It is also assumed the first row of the loaded file contains column headers. In both

cases, it is assumed a given data frame/matrix, or the one loaded from a given file path, will contain CNV data for each of the top genes, as determined by the user settings for the 'hypermethGplusAnalysis' and 'hypomethGplusAnalysis' arguments, as well as the 'topGeneNumber', with the gene IDs for those genes followed by "\_CNV" in the column names. If this variable is set to NA, then the CNV data will be assumed to already be contained in the colData of the TENETMultiAssayExperiment under columns similarly named as the gene ID followed by "\_CNV" for each of the top genes, as determined by the user settings for the 'hypermethGplusAnalysis' and 'hypomethGplusAnalysis' arguments, as well as the 'topGeneNumber'. Note: if a given gene is missing this information, then the plot will be generated without considering the CNV status of that gene. Defaults to NA, and is only considered if 'simpleOrComplex' is set to "complex".

SMDData	Specify the SM status for each of the top genes and each sample in the TENETMultiAssayExperiment. SM status should be given in the form as either 0 or 1, or "no mutation" or "mutation", indicating whether each sample harbors a somatic mutation for each of the top genes as determined by 'hypermethGplusAnalysis', 'hypomethGplusAnalysis', and 'topGeneNumber' settings. This argument can be given in the form of a data frame/matrix, or a path to a file that contains the SM data, or it can be set to NA. See the documentation for 'CNV-Data' for more information. Note: if a given gene is missing this information, then the plot will be generated without considering the SM status of that gene. Defaults to NA, and is only considered if 'simpleOrComplex' is set to "complex".
purityData	Specify the cellularity/purity data for samples in the TENETMultiAssayExperiment. Purity values should range from 0 to 1. This data can be given in a variety of forms, including a vector, data frame/matrix, or a path to a file that contains the purity data. If a vector is given, the names of the vector elements must correspond to the names of the samples in the rownames of the colData of the TENETMultiAssayExperiment object. If no names are provided for the vector, then the number of elements in the vector must equal the number of samples in the colData, and are assumed to align with the samples as they are ordered in the colData. If a data frame or matrix is given, then the rownames of the supplied data frame or matrix must include the sample names as they appear in the colData of the TENETMultiAssayExperiment object, and the first column of the data frame or matrix will be assumed to include the purity data. If a single string is provided, then it is assumed to be a path to a tab-delimited file containing purity data in the second column, and the names of the samples, again corresponding with the sample names in the colData, in the first column, which will be loaded as the row names. The first row of the file must contain column names. If this variable is set to NA, then the purity data will be assumed to already be contained in the colData of the TENETMultiAssayExperiment under a column titled "purity". Defaults to NA, and is only considered if 'simpleOrComplex' is set to "complex".
coreCount	Argument passed as the mc.cores argument to mcmapply. See ?parallel::mcmapply for more details. Defaults to 1.



**Value**

Returns the MultiAssayExperiment object given as the TENETMultiAssayExperiment argument with an additional list of information named 'step7ExpressionVsDNAMethylationScatterplots' in its metadata with the output of this function. This list is subdivided into hypermethGplus or hypomethGplus results as selected by the user, which are further subdivided into lists with data for the top overall genes, and for top TF genes only. An additional list named 'selectedDNAMethylationSites' is also generated if the user has specified RE DNA methylation sites of interest to the RE DNA methylation sites argument of this function. Within each of these lists, a final list is generated for each of the top genes/TFs, or specified RE DNA methylation sites, which contains the scatterplots for each RE DNA methylation site linked to the top genes/TFs, or for each gene linked to the user specified REs if DNA methylation sites are specified, for each analysis type. In each scatterplot, the expression of the gene is plotted on the X-axis, and the methylation of the linked RE DNA methylation site is plotted on the Y-axis. If the user has opted to create complex plots, and the CNV and SM data are available for the plotted gene, the CNV and SM status for each case sample will be reflected in each point's symbol used (with SM status taking precedence over CNV), while the purity of each sample will be reflected in the size of the point.

**Examples**

```
## This example uses the example MultiAssayExperiment provided in the
## TENET.ExperimentHub package to create scatterplots for the top 10
## genes/TFs, by number of linked hyper- or hypomethylated RE DNA
## methylation sites, showing expression of these genes and the DNA
## methylation level of their linked RE DNA methylation sites. Gene names
## will be retrieved from the rowRanges of the 'expression'
## SummarizedExperiment object in the example MultiAssayExperiment. Only
## simple scatterplots will be created. The analysis will be performed using
## one CPU core.

## Load the example TENET MultiAssayExperiment object
## from the TENET.ExperimentHub package
exampleTENETMultiAssayExperiment <-
  TENET.ExperimentHub::exampleTENETMultiAssayExperiment()

## Use the example dataset to create the scatterplots
returnValue <- step7ExpressionVsDNAMethylationScatterplots(
  TENETMultiAssayExperiment = exampleTENETMultiAssayExperiment
)

## This example uses the example MultiAssayExperiment provided in the
## TENET.ExperimentHub package to create scatterplots for only the top 5
## genes/TFs, by number of linked hypomethylated RE DNA methylation sites,
## as well as for a given vector with some example RE DNA methylation sites
## of interest. Gene names will be retrieved from the rowRanges of the
## 'expression' SummarizedExperiment object in the example
## MultiAssayExperiment. For each plot, complex scatterplots displaying each
## sample's CNV and SM status for the gene in the plot, as well as purity,
## where available. The gene CNV and SM status, as well as purity from each
## patient sample in the analyses, will be taken from specific columns
## present in the exampleTENETClinicalDataFrame object. The analysis will be
## performed using 8 CPU cores.
```

```
## Load the example TENET MultiAssayExperiment object
## from the TENET.ExperimentHub package
exampleTENETMultiAssayExperiment <-
  TENET.ExperimentHub::exampleTENETMultiAssayExperiment()

## Again, this loads the data frame with example clinical data for patients
## in the TENET MultiAssayExperiment object from the TENET.ExperimentHub
## package
exampleTENETClinicalDataFrame <-
  TENET.ExperimentHub::exampleTENETClinicalDataFrame()

## Use the example dataset to create the scatterplots
returnValue <- step7ExpressionVsDNAMethylationScatterplots(
  TENETMultiAssayExperiment = exampleTENETMultiAssayExperiment,
  hypermethGplusAnalysis = FALSE,
  topGeneNumber = 5,
  DNAMethylationSites = c("cg03095778", "cg24011501", "cg12989041"),
  simpleOrComplex = "complex",
  CNVData = exampleTENETClinicalDataFrame[seq(4, 42, by = 2)],
  SMDData = exampleTENETClinicalDataFrame[seq(5, 43, by = 2)],
  purityData = exampleTENETClinicalDataFrame[3],
  coreCount = 8
)
```

---

```
step7LinkedDNAMethylationSiteCountHistograms
```

*Create histograms displaying the number of genes or transcription factors linked to a given number of RE DNA methylation sites*

---

## Description

This function generates histograms displaying the number of all genes, or transcription factor (TF) genes only, with links to a given number of regulatory element DNA methylation sites from both of the hyper- or hypomethylated G+ analysis quadrants, as selected by the user.

## Usage

```
step7LinkedDNAMethylationSiteCountHistograms(
  TENETMultiAssayExperiment,
  hypermethGplusAnalysis = TRUE,
  hypomethGplusAnalysis = TRUE
)
```

## Arguments

**TENETMultiAssayExperiment**  
Specify a MultiAssayExperiment object containing expression and methylation SummarizedExperiment objects, such as one created by the TCGADownloader

function. This MultiAssayExperiment object should also contain the results from the step6DNAMethylationSitesPerGeneTabulation function in its metadata.

hypermethGplusAnalysis

Set to TRUE to create histograms of genes/TFs linked to hypermethylated RE DNA methylation sites with G+ links. Defaults to TRUE.

hypomethGplusAnalysis

Set to TRUE to create histograms of genes/TFs linked to hypomethylated RE DNA methylation sites with G+ links. Defaults to TRUE.

## Value

Returns the MultiAssayExperiment object given as the TENETMultiAssayExperiment argument with an additional list of information named 'step7LinkedDNAMethylationSiteCountHistograms' in its metadata with the output of this function. This list is subdivided into hypermethGplus or hypomethGplus results as selected by the user. Each of these contains a histogram displaying the number of all genes, or TFs only, linked to a given number of RE DNA methylation sites in each of the selected analysis quadrants.

## Examples

```
## This example uses the example MultiAssayExperiment provided in the
## TENET.ExperimentHub package to create histograms displaying counts of
## genes/TFs by the number of linked hyper- or hypomethylated G+ RE DNA
## methylation sites.

## Load the example TENET MultiAssayExperiment object
## from the TENET.ExperimentHub package
exampleTENETMultiAssayExperiment <-
  TENET.ExperimentHub::exampleTENETMultiAssayExperiment()

## Use the example dataset to create the RE DNA methylation site count
## histograms
returnValue <- step7LinkedDNAMethylationSiteCountHistograms(
  TENETMultiAssayExperiment = exampleTENETMultiAssayExperiment
)

## This example also uses the example MultiAssayExperiment provided in the
## TENET.ExperimentHub package, but only creates a histogram of counts of
## genes/TFs by the number of linked hypomethylated G+ RE DNA methylation
## sites.

## Load the example TENET MultiAssayExperiment object
## from the TENET.ExperimentHub package
exampleTENETMultiAssayExperiment <-
  TENET.ExperimentHub::exampleTENETMultiAssayExperiment()

## Use the example dataset to create the RE DNA methylation site count
## histograms
returnValue <- step7LinkedDNAMethylationSiteCountHistograms(
  TENETMultiAssayExperiment = exampleTENETMultiAssayExperiment,
```

```

    hypermethGplusAnalysis = FALSE
  )

```

---

## step7LinkedDNAMethylationSitesMotifSearching

*Perform motif searching for transcription factor motifs in the vicinity of RE DNA methylation sites linked to the specified transcription factors*

---

### Description

This function takes a named list of transcription factors (TFs) and their binding motifs, and for each of the TFs, it identifies if the specified motif for that TF is found within a user-specified distance to each RE DNA methylation site linked to that TF from both of the hyper- or hypomethylated G+ analysis quadrants, as selected by the user.

### Usage

```

step7LinkedDNAMethylationSitesMotifSearching(
  TENETMultiAssayExperiment,
  TFMotifList,
  geneAnnotationDataset = NA,
  DNAMethylationArray = NA,
  hypermethGplusAnalysis = TRUE,
  hypomethGplusAnalysis = TRUE,
  distanceFromREDNAMethylationSites = 100,
  matchPWMMinScore = "75%",
  coreCount = 1
)

```

### Arguments

- |                           |   |
|---------------------------|---|
| TENETMultiAssayExperiment | Specify a MultiAssayExperiment object containing expression and methylation SummarizedExperiment objects, such as one created by the TCGADownloader function. This MultiAssayExperiment object should also contain the results from the step50optimizeLinks function in its metadata.   |
| TFMotifList               | Specify a named list mapping transcription factor gene names and/or IDs to their respective motif position weight matrix (PWM). The PWMs should be in the form of a 4xN matrix. If a PWM is not available, the user can select one by querying the MotifDb package database. See the example below for more information. <b>Note:</b> The name of each PWM in the list must match the gene name or Ensembl ID of a gene in the TENETMultiAssayExperiment with RE DNA methylation sites linked to it for the specified analysis types. |

geneAnnotationDataset	Specify a gene annotation dataset which is used to identify names for genes by their Ensembl IDs. The argument must be either a GRanges object (such as one imported via <code>rtracklayer::import</code> ) or a path to a GFF3 or GTF file. Both GENCODE and Ensembl annotations are supported. Other annotation datasets may work, but have not been tested. See the "Input data" section of the vignette for information on the required dataset format. Specify NA to use the names for genes listed in the "geneName" column of the elementMetadata of the rowRanges of the "expression" SummarizedExperiment object within the TENETMultiAssayExperiment object. Defaults to NA.
DNAMethylationArray	Specify the name of a DNA methylation probe array supported by the sesameData package (see <code>?sesameData::sesameData_getManifestGRanges</code> ). If an array is specified, RE DNA methylation sites and their locations in that array's manifest are cross-referenced with RE DNA methylation site IDs included in the rownames of the methylation dataset provided in the "methylation" SummarizedExperiment object within the TENETMultiAssayExperiment object, and only those overlapping will be considered for analysis. If set to NA, all RE DNA methylation sites with locations listed in the rowRanges of the "methylation" SummarizedExperiment object are used. Defaults to NA.
hypermethGplusAnalysis	Set to TRUE to do motif searching in the vicinity of hypermethylated RE DNA methylation sites with G+ links to the TF of interest. Defaults to TRUE.
hypomethGplusAnalysis	Set to TRUE to do motif searching in the vicinity of hypomethylated RE DNA methylation sites with G+ links to the TF of interest. Defaults to TRUE.
distanceFromREDNAMethylationSites	Specify a positive integer in base pairs to be the distance from the linked RE DNA methylation sites within which motif searching will be performed. Defaults to 100.
matchPWMMinScore	Specify the min.score argument passed to the matchPWM function for motif searching. See <code>?Biostrings::matchPWM</code> for more details. Defaults to "75%".
coreCount	Argument passed as the mc.cores argument to mclapply. See <code>?parallel::mclapply</code> for more details. Defaults to 1.

## Value

Returns the MultiAssayExperiment object given as the TENETMultiAssayExperiment argument with an additional list of information named 'step7LinkedDNAMethylationSitesMotifSearching' in its metadata with the output of this function. This list is subdivided into hypermethGplus or hypomethGplus results as selected by the user. Each contains a list for each specified TF gene with a seqLogo of the gene's motif PWM, a table of the PWM, and two additional tables with information on motif occurrences found in the vicinity of linked RE DNA methylation sites. The first table contains information about each motif found in the vicinity of the linked RE DNA methylation sites for the TF gene at the specified matchPWMMinScore. The second lists each RE DNA methylation site linked to the TF gene for the analysis types specified, and the number of motif occurrences found in the vicinity of each of those RE DNA methylation sites.

**Examples**

```
## These examples require the MotifDb package
if (requireNamespace("MotifDb", quietly = TRUE)) {
  ## Show available motifs for example TF FOXA1
  names(MotifDb::query(MotifDb::MotifDb, "FOXA1"))

  ## The seqLogos for all input motifs will also be included in the output
  ## of this function. Alternatively, individual motifs can be visualized
  ## with the seqLogo function from the seqLogo package.
  seqLogo::seqLogo(MotifDb::query(MotifDb::MotifDb, "FOXA1")[[3]])

  ## Once PWMs have been selected for use, a list containing them must be
  ## created
  exampleTFMotifList <- list(
    "FOXA1" = MotifDb::query(MotifDb::MotifDb, "FOXA1")[[3]],
    "ESR1" = MotifDb::query(MotifDb::MotifDb, "ESR1")[[4]]
  )

  ## This example uses the example MultiAssayExperiment provided in the
  ## TENET.ExperimentHub package to perform motif overlapping for all
  ## hyper- and hypomethylated RE DNA methylation sites linked to the
  ## FOXA1 and ESR1 genes using the motifs for each gene specified in the
  ## exampleTFMotifList. Gene names and locations, and the locations of RE
  ## DNA methylation sites, will be retrieved from the rowRanges of the
  ## 'expression' and 'methylation' SummarizedExperiment objects in the
  ## example MultiAssayExperiment. Regions within 100 bp of linked RE DNA
  ## methylation sites will be checked for motifs, and a similarity
  ## threshold of 75% will be used to identify motifs. The analysis will
  ## be performed using one CPU core.

  ## Load the example TENET MultiAssayExperiment object
  ## from the TENET.ExperimentHub package
  exampleTENETMultiAssayExperiment <-
    TENET.ExperimentHub::exampleTENETMultiAssayExperiment()

  ## Use the example dataset to do the motif searching
  returnValue <- step7LinkedDNAMethylationSitesMotifSearching(
    TENETMultiAssayExperiment = exampleTENETMultiAssayExperiment,
    TFMotifList = exampleTFMotifList
  )

  ## This example uses the example MultiAssayExperiment provided in the
  ## TENET.ExperimentHub package to perform motif overlapping for only
  ## hypomethylated RE DNA methylation sites linked to the FOXA1 and ESR1
  ## genes using the motifs for each gene specified in the
  ## exampleTFMotifList. Gene names and locations, and the locations of RE
  ## DNA methylation sites, will be retrieved from the rowRanges of the
  ## 'expression' and 'methylation' SummarizedExperiment objects in the
  ## example MultiAssayExperiment. Regions within 50 bp of linked RE DNA
  ## methylation sites will be checked for motifs, and a similarity
  ## threshold of 80% will be used to identify motifs. The analysis will
  ## be performed using 8 CPU cores.
```

```

## Load the example TENET MultiAssayExperiment object
## from the TENET.ExperimentHub package
exampleTENETMultiAssayExperiment <-
  TENET.ExperimentHub::exampleTENETMultiAssayExperiment()

## Use the example dataset to do the motif searching
returnValue <- step7LinkedDNAMethylationSitesMotifSearching(
  TENETMultiAssayExperiment = exampleTENETMultiAssayExperiment,
  TFMotifList = exampleTFMotifList,
  hypermethGplusAnalysis = FALSE,
  distanceFromREDNAMethylationSites = 50,
  matchPWMMinScore = "80%",
  coreCount = 8
)
} else {
  message(
    "These examples require the MotifDb package. Run ",
    'BiocManager::install("MotifDb") and rerun these examples.'
  )
}

```

---

step7SelectedDNAMethylationSitesCaseVsControlBoxplots

*Generate boxplots comparing the methylation level of the specified RE  
DNA methylation sites in case and control samples*

---

## Description

This function takes a vector of RE DNA methylation sites specified by the user and generates boxplots displaying the methylation level of each of these DNA methylation sites in the case compared to control samples, along with the results of a Student's t-test comparing the methylation level between these two groups.

## Usage

```

step7SelectedDNAMethylationSitesCaseVsControlBoxplots(
  TENETMultiAssayExperiment,
  DNAMethylationSites,
  coreCount = 1
)

```

## Arguments

TENETMultiAssayExperiment  
Specify a MultiAssayExperiment object containing a methylation Summarized-Experiment object, such as one created by the TCGADownloader function.

**DNAMethylationSites**

Supply a vector of RE DNA methylation site IDs for which boxplots with the methylation of those RE DNA methylation sites will be created.

**coreCount**

Argument passed as the mc.cores argument to mcmapply. See ?parallel::mcmapply for more details. Defaults to 1.

**Value**

Returns the MultiAssayExperiment object given as the TENETMultiAssayExperiment argument with an additional list of data named 'step7SelectedDNAMethylationSitesCaseVsControlBoxplots' in its metadata with the output of this function, which contains boxplots showing the methylation of the RE DNA methylation sites of interest in the case and control samples, with Student's t-test p-values and the ID of the RE DNA methylation site in the title.

**Examples**

```
## This example uses the example MultiAssayExperiment provided in the
## TENET.ExperimentHub package to generate boxplots for several selected
## RE DNA methylation sites, using one CPU core.

## Load the example TENET MultiAssayExperiment object
## from the TENET.ExperimentHub package
exampleTENETMultiAssayExperiment <-
  TENET.ExperimentHub::exampleTENETMultiAssayExperiment()

## Use the example dataset to create RE DNA methylation site case vs. control
## boxplots
returnValue <- step7SelectedDNAMethylationSitesCaseVsControlBoxplots(
  TENETMultiAssayExperiment = exampleTENETMultiAssayExperiment,
  DNAMethylationSites = c("cg03095778", "cg24011501", "cg12989041"),
  coreCount = 1
)
```

---

step7StatesForLinks	<i>Identify which of the case samples harbor each of the identified regulatory element DNA methylation site-gene links</i>
---------------------	--

---

**Description**

This function generates data frames with information for each of the case samples and each RE DNA methylation site-gene link from both of the hyper- or hypomethylated G+ analysis quadrants, as selected by the user, on if a given sample is said to "harbor" each link, depending on if the methylation of the given sample for the RE DNA methylation site in the link is above or below the hyper- or hypomethylation cutoff defined in the step2GetDifferentiallyMethylatedSites function and the expression of the gene in the link is significantly greater than, or less than, the mean expression in the control samples, as determined by a Bonferroni-corrected 1-sided t-test with a p-value threshold of 0.05.



**Usage**

```
step7StatesForLinks(
  TENETMultiAssayExperiment,
  hypermethGplusAnalysis = TRUE,
  hypomethGplusAnalysis = TRUE,
  coreCount = 1
)
```

**Arguments**

TENETMultiAssayExperiment	Specify a MultiAssayExperiment object containing expression and methylation SummarizedExperiment objects, such as one created by the TCGADownloader function. This MultiAssayExperiment object should also contain the results from the step5OptimizeLinks functions in its metadata.
hypermethGplusAnalysis	Set to TRUE to analyze which case samples harbor each hypermethylated G+RE DNA methylation site-gene link. Defaults to TRUE.
hypomethGplusAnalysis	Set to TRUE to analyze which case samples harbor each hypomethylated G+RE DNA methylation site-gene link. Defaults to TRUE.
coreCount	Argument passed as the mc.cores argument to mcmapply. See ?parallel::mcmapply for more details. Defaults to 1.

**Value**

Returns the MultiAssayExperiment object given as the TENETMultiAssayExperiment argument with an additional list of information named 'step7StatesForLinks' in its metadata with the output of this function. This list is subdivided into hypermethGplus or hypomethGplus results as selected by the user, which contain data frames for each of the specified analysis types with case samples in the columns and each RE DNA methylation site-gene link for that analysis type in the rows, with a 1 indicating the sample is positive for that link and a 0 indicating it is not. NA values are shown for samples that lack methylation data for the site or expression data for the gene.

**Examples**

```
## This example uses the example MultiAssayExperiment provided in the
## TENET.ExperimentHub package to create data frames for link states between
## genes and both hyper- and hypomethylated G+ RE DNA methylation sites. The
## analysis will be performed using one CPU core.
```

```
## Load the example TENET MultiAssayExperiment object
## from the TENET.ExperimentHub package
exampleTENETMultiAssayExperiment <-
  TENET.ExperimentHub::exampleTENETMultiAssayExperiment()
```

```
## Use the example dataset to do the states for links analysis
returnValue <- step7StatesForLinks(
  TENETMultiAssayExperiment = exampleTENETMultiAssayExperiment
```

```

)

## This example uses the example MultiAssayExperiment provided in the
## TENET.ExperimentHub package to create data frames for link states between
## genes and only hypomethylated G+ RE DNA methylation sites. The analysis
## will be performed using 8 CPU cores.

## Load the example TENET MultiAssayExperiment object
## from the TENET.ExperimentHub package
exampleTENETMultiAssayExperiment <-
  TENET.ExperimentHub::exampleTENETMultiAssayExperiment()

## Use the example dataset to do the states for links analysis
returnValue <- step7StatesForLinks(
  TENETMultiAssayExperiment = exampleTENETMultiAssayExperiment,
  hypermethGplusAnalysis = FALSE,
  coreCount = 8
)

```

---

step7TopGenesCaseVsControlExpressionBoxplots

*Create boxplots comparing the expression level of the top genes/transcription factors in case and control samples*

---

## Description

This function takes the top genes/transcription factors (TFs) for each analysis type by number of linked RE DNA methylation sites identified by the `step6DNAMethylationSitesPerGeneTabulation` function, up to the number specified by the user, and generates boxplots displaying the expression level of each of these genes in the case compared to control samples, along with the results of a Student's t-test comparing the expression level between these two groups.

## Usage

```

step7TopGenesCaseVsControlExpressionBoxplots(
  TENETMultiAssayExperiment,
  geneAnnotationDataset = NA,
  hypermethGplusAnalysis = TRUE,
  hypomethGplusAnalysis = TRUE,
  topGeneNumber = 10,
  coreCount = 1
)

```

## Arguments

**TENETMultiAssayExperiment**  
Specify a MultiAssayExperiment object containing expression and methylation SummarizedExperiment objects, such as one created by the TCGADownloader

function. This MultiAssayExperiment object should also contain the results from the step5OptimizeLinks and step6DNAMethylationSitesPerGeneTabulation functions in its metadata.

#### geneAnnotationDataset

Specify a gene annotation dataset which is used to identify names for genes by their Ensembl IDs. The argument must be either a GRanges object (such as one imported via `rtracklayer::import`) or a path to a GFF3 or GTF file. Both GENCODE and Ensembl annotations are supported. Other annotation datasets may work, but have not been tested. See the "Input data" section of the vignette for information on the required dataset format. Specify NA to use the names for genes listed in the "geneName" column of the elementMetadata of the rowRanges of the "expression" SummarizedExperiment object within the TENETMultiAssayExperiment object. Defaults to NA.

#### hypermethGplusAnalysis

Set to TRUE to create expression boxplots for the top genes/TFs with the most hypermethylated RE DNA methylation sites with G+ links. Defaults to TRUE.

#### hypomethGplusAnalysis

Set to TRUE to create expression boxplots for the top genes/TFs with the most hypomethylated RE DNA methylation sites with G+ links. Defaults to TRUE.

**topGeneNumber** Specify the number of top genes/TFs, based on the most linked RE DNA methylation sites of a given analysis type, for which to generate expression boxplots. Defaults to 10.

**coreCount** Argument passed as the `mc.cores` argument to `mcmapply`. See `?parallel::mcmapply` for more details. Defaults to 1.

### Value

Returns the MultiAssayExperiment object given as the TENETMultiAssayExperiment argument with an additional list of information named 'step7TopGenesCaseVsControlExpressionBoxplots' in its metadata with the output of this function. This list is subdivided into hypermethGplus or hypomethGplus results as selected by the user, which are further subdivided into lists with plots for the top overall genes, and for top TF genes only. These contain boxplots showing the expression of the genes of interest in the case and control samples, with Student's t-test p-values and the name and ID of the gene in the title.

### Examples

```
## This example uses the example MultiAssayExperiment provided in the
## TENET.ExperimentHub package to create expression boxplots in case vs.
## control samples for the top 10 genes/TFs, by number of linked hyper- or
## hypomethylated RE DNA methylation sites. Gene names will be retrieved
## from the rowRanges of the 'expression' SummarizedExperiment object in the
## example MultiAssayExperiment. The analysis will be performed using one
## CPU core.
```

```
## Load the example TENET MultiAssayExperiment object
## from the TENET.ExperimentHub package
exampleTENETMultiAssayExperiment <-
  TENET.ExperimentHub::exampleTENETMultiAssayExperiment()
```

```
## Use the example dataset to create expression case vs. control boxplots
returnValue <- step7TopGenesCaseVsControlExpressionBoxplots(
  TENETMultiAssayExperiment = exampleTENETMultiAssayExperiment
)

## This example uses the example MultiAssayExperiment provided in the
## TENET.ExperimentHub package to create expression boxplots in case vs.
## control samples for the top 5 genes/TFs, by number of linked
## hypomethylated RE DNA methylation sites only. Gene names will be
## retrieved from the rowRanges of the 'expression' SummarizedExperiment
## object in the example MultiAssayExperiment. The analysis will be
## performed using 8 CPU cores.

## Load the example TENET MultiAssayExperiment object
## from the TENET.ExperimentHub package
exampleTENETMultiAssayExperiment <-
  TENET.ExperimentHub::exampleTENETMultiAssayExperiment()

## Use the example dataset to create expression case vs. control boxplots
returnValue <- step7TopGenesCaseVsControlExpressionBoxplots(
  TENETMultiAssayExperiment = exampleTENETMultiAssayExperiment,
  hypermethGplusAnalysis = FALSE,
  topGeneNumber = 5,
  coreCount = 8
)
```

---

## step7TopGenesCircosPlots

*Generate Circos plots displaying the links between top identified genes and each of the RE DNA methylation sites linked to them*

---

### Description

This function takes the top genes/TFs by number of linked regulatory element DNA methylation sites identified by the `step6DNAMethylationSitesPerGeneTabulation` function, up to the number specified by the user, and generates Circos plots for each gene showing the genomic links between each gene and each RE DNA methylation site linked to the gene for the analysis types specified.

### Usage

```
step7TopGenesCircosPlots(
  TENETMultiAssayExperiment,
  DNAMethylationArray = NA,
  geneAnnotationDataset = NA,
  hypermethGplusAnalysis = TRUE,
  hypomethGplusAnalysis = TRUE,
```

```

    topGeneNumber = 10,
    coreCount = 1
)

```

## Arguments

### TENETMultiAssayExperiment

Specify a MultiAssayExperiment object containing expression and methylation SummarizedExperiment objects, such as one created by the TCGADownloader function. This MultiAssayExperiment object should also contain the results from the step5OptimizeLinks and step6DNAMethylationSitesPerGeneTabulation functions in its metadata.

### DNAMethylationArray

Specify the name of a DNA methylation probe array supported by the sesame-Data package (see ?sesameData::sesameData\_getManifestGRanges). If an array is specified, RE DNA methylation sites and their locations in that array's manifest are cross-referenced with RE DNA methylation site IDs included in the rownames of the methylation dataset provided in the "methylation" SummarizedExperiment object within the TENETMultiAssayExperiment object, and only those overlapping will be considered for analysis. If set to NA, all RE DNA methylation sites with locations listed in the rowRanges of the "methylation" SummarizedExperiment object are used. Defaults to NA.

### geneAnnotationDataset

Specify a gene annotation dataset which is used to identify names for genes by their Ensembl IDs. The argument must be either a GRanges object (such as one imported via rtracklayer::import) or a path to a GFF3 or GTF file. Both GENCODE and Ensembl annotations are supported. Other annotation datasets may work, but have not been tested. See the "Input data" section of the vignette for information on the required dataset format. Specify NA to use the names for genes listed in the "geneName" column of the elementMetadata of the rowRanges of the "expression" SummarizedExperiment object within the TENETMultiAssayExperiment object. Defaults to NA.

### hypermethGplusAnalysis

Set to TRUE to create Circos plots displaying links between the top genes/TFs by most hypermethylated RE DNA methylation sites with G+ links and their linked RE DNA methylation sites of that type. Defaults to TRUE.

### hypomethGplusAnalysis

Set to TRUE to create Circos plots displaying links between the top genes/TFs by most hypomethylated RE DNA methylation sites with G+ links and their linked RE DNA methylation sites of that type. Defaults to TRUE.

**topGeneNumber** Specify the number of top genes/TFs, based on the most linked RE DNA methylation sites of a given analysis type, for which to generate Circos plots showing the links between the genes and each of their linked RE DNA methylation sites. Defaults to 10.

**coreCount** Argument passed as the mc.cores argument to mclapply. See ?parallel::mclapply for more details. Defaults to 1.

**Value**

Returns the MultiAssayExperiment object given as the TENETMultiAssayExperiment argument with an additional list of information named 'step7TopGenesCircosPlots' in its metadata with the output of this function. This list is subdivided into hypermethGplus or hypomethGplus results as selected by the user, which are further subdivided into lists with data for the top overall genes and for top TF genes only. Each of these lists contain Circos plots for each of the top genes/TFs visualizing the links between the genes and their linked RE DNA methylation sites for the selected analysis types.

**Examples**

```
## This example uses the example MultiAssayExperiment provided in the
## TENET.ExperimentHub package to create Circos plots for the top 10
## genes/TFs, by number of linked hyper- or hypomethylated RE DNA
## methylation sites. Gene names and locations and RE DNA methylation site
## locations will be retrieved from the rowRanges of the 'expression' and
## 'methylation' SummarizedExperiment objects in the example
## MultiAssayExperiment. The analysis will be performed using one CPU core.

## Load the example TENET MultiAssayExperiment object
## from the TENET.ExperimentHub package
exampleTENETMultiAssayExperiment <-
  TENET.ExperimentHub::exampleTENETMultiAssayExperiment()

## Use the example dataset to create Circos plots
returnValue <- step7TopGenesCircosPlots(
  TENETMultiAssayExperiment = exampleTENETMultiAssayExperiment
)

## This example uses the example MultiAssayExperiment provided in the
## TENET.ExperimentHub package to create Circos plots for the top 5
## genes/TFs, by number of linked hypomethylated RE DNA methylation sites
## only. Gene names and locations will be retrieved from the rowRanges of
## the 'expression' and 'methylation' SummarizedExperiment objects in the
## example MultiAssayExperiment. RE DNA methylation site IDs and locations
## will be retrieved from the HM450 array via the sesameData package. Eight
## CPU cores will be used to create plots.

## Load the example TENET MultiAssayExperiment object
## from the TENET.ExperimentHub package
exampleTENETMultiAssayExperiment <-
  TENET.ExperimentHub::exampleTENETMultiAssayExperiment()

## Use the example dataset to create Circos plots
returnValue <- step7TopGenesCircosPlots(
  TENETMultiAssayExperiment = exampleTENETMultiAssayExperiment,
  DNAMethylationArray = "HM450",
  hypermethGplusAnalysis = FALSE,
  topGeneNumber = 5,
  coreCount = 8
)
```

---

**step7TopGenesDNAMethylationHeatmaps**

*Generate heatmaps displaying the methylation level of all RE DNA methylation sites linked to the top genes/transcription factors, along with the expression of those genes in the column headers, in the case samples within the supplied MultiAssayExperiment object*

---

**Description**

This function takes the top genes/transcription factors (TFs) for each analysis type by number of linked RE DNA methylation sites identified by the `step6DNAMethylationSitesPerGeneTabulation` function, up to the number specified by the user, and generates heatmaps displaying the methylation level of the unique RE DNA methylation sites linked to any of those genes, along with the expression of those genes in the case samples only.

**Usage**

```
step7TopGenesDNAMethylationHeatmaps(  
  TENETMultiAssayExperiment,  
  geneAnnotationDataset = NA,  
  hypermethGplusAnalysis = TRUE,  
  hypomethGplusAnalysis = TRUE,  
  topGeneNumber = 10  
)
```

**Arguments**

`TENETMultiAssayExperiment`

Specify a `MultiAssayExperiment` object containing expression and methylation `SummarizedExperiment` objects, such as one created by the `TCGADownloader` function. This `MultiAssayExperiment` object should also contain the results from the `step2GetDifferentiallyMethylatedSites`, `step5OptimizeLinks`, and `step6DNAMethylationSitesPerGeneTabulation` functions in its metadata.

`geneAnnotationDataset`

Specify a gene annotation dataset which is used to identify names for genes by their Ensembl IDs. The argument must be either a `GRanges` object (such as one imported via `rtracklayer::import`) or a path to a GFF3 or GTF file. Both GENCODE and Ensembl annotations are supported. Other annotation datasets may work, but have not been tested. See the "Input data" section of the vignette for information on the required dataset format. Specify `NA` to use the names for genes listed in the "geneName" column of the `elementMetadata` of the `rowRanges` of the "expression" `SummarizedExperiment` object within the `TENETMultiAssayExperiment` object. Defaults to `NA`.

**hypermethGplusAnalysis**

Set to TRUE to create heatmaps showing DNA methylation levels of RE DNA methylation sites linked to the top genes/TFs with the most hypermethylated RE DNA methylation sites with G+ links. Defaults to TRUE.

**hypomethGplusAnalysis**

Set to TRUE to create heatmaps showing DNA methylation levels of RE DNA methylation sites linked to the top genes/TFs with the most hypomethylated RE DNA methylation sites with G+ links. Defaults to TRUE.

**topGeneNumber**

Specify the number of top genes/TFs, based on the most linked RE DNA methylation sites of a given analysis type, for which to generate heatmaps with their linked RE DNA methylation sites' methylation levels. Defaults to 10.

**Value**

Returns the MultiAssayExperiment object given as the TENETMultiAssayExperiment argument with an additional list of information named 'step7TopGenesDNAMethylationHeatmaps' in its metadata with the output of this function. This list is subdivided into hypermethGplus or hypomethGplus results as selected by the user, which are further subdivided into lists with data for the top overall genes and for top TF genes only. Each of these contains a single heatmap, with the expression of the top genes/TFs in the column headers and the methylation of their unique linked RE DNA methylation sites in the body of the heatmaps.

**Examples**

```
## This example uses the example MultiAssayExperiment provided in the
## TENET.ExperimentHub package to create heatmaps for the top 10 genes/TFs,
## by number of linked hyper- or hypomethylated RE DNA methylation sites, as
## well as the unique RE DNA methylation sites linked to those 10 genes.
## Gene names will be retrieved from the rowRanges of the 'expression'
## SummarizedExperiment object in the example MultiAssayExperiment.
```

```
## Load the example TENET MultiAssayExperiment object
## from the TENET.ExperimentHub package
exampleTENETMultiAssayExperiment <-
  TENET.ExperimentHub::exampleTENETMultiAssayExperiment()
```

```
## Use the example dataset to create methylation heatmaps
returnValue <- step7TopGenesDNAMethylationHeatmaps(
  TENETMultiAssayExperiment = exampleTENETMultiAssayExperiment
)
```

```
## This example uses the example MultiAssayExperiment provided in the
## TENET.ExperimentHub package to create heatmaps for only the top 5
## genes/TFs, by number of linked hypomethylated RE DNA methylation sites
## only, as well as the unique RE DNA methylation sites linked to those 5
## genes/TFs. Gene names will be retrieved from the rowRanges of the
## 'expression' SummarizedExperiment object in the example
## MultiAssayExperiment.
```

```
## Load the example TENET MultiAssayExperiment object
## from the TENET.ExperimentHub package
```



```

exampleTENETMultiAssayExperiment <-
  TENET.ExperimentHub::exampleTENETMultiAssayExperiment()

## Use the example dataset to create methylation heatmaps
returnValue <- step7TopGenesDNAMethylationHeatmaps(
  TENETMultiAssayExperiment = exampleTENETMultiAssayExperiment,
  hypermethGplusAnalysis = FALSE,
  topGeneNumber = 5
)

```

---

step7TopGenesExpressionCorrelationHeatmaps

*Generate mirrored heatmaps displaying the correlation of the expression values of the top genes/TFs*

---

## Description

This function takes the top genes/TFs for each analysis type by number of linked RE DNA methylation sites identified by the step6DNAMethylationSitesPerGeneTabulation function, up to the number specified by the user, and generates heatmaps showing the correlation R-values between the expression of the top genes/TFs, as well as data frames with the R-values.

## Usage

```

step7TopGenesExpressionCorrelationHeatmaps(
  TENETMultiAssayExperiment,
  geneAnnotationDataset = NA,
  hypermethGplusAnalysis = TRUE,
  hypomethGplusAnalysis = TRUE,
  topGeneNumber = 10
)

```

## Arguments

TENETMultiAssayExperiment

Specify a MultiAssayExperiment object containing expression and methylation SummarizedExperiment objects, such as one created by the TCGADownloader function. This MultiAssayExperiment object should also contain the results from the step6DNAMethylationSitesPerGeneTabulation function in its metadata.

geneAnnotationDataset

Specify a gene annotation dataset which is used to identify names for genes by their Ensembl IDs. The argument must be either a GRanges object (such as one imported via rtracklayer::import) or a path to a GFF3 or GTF file. Both GENCODE and Ensembl annotations are supported. Other annotation datasets may work, but have not been tested. See the "Input data" section of the vignette for information on the required dataset format. Specify NA to use the

names for genes listed in the "geneName" column of the elementMetadata of the rowRanges of the "expression" SummarizedExperiment object within the TENETMultiAssayExperiment object. Defaults to NA.

**hypermethGplusAnalysis** Set to TRUE to create heatmaps and tables showing expression correlation values for the top genes/TFs with the most hypermethylated RE DNA methylation sites with G+ links. Defaults to TRUE.

**hypomethGplusAnalysis** Set to TRUE to create heatmaps and tables showing expression correlation values for the top genes/TFs with the most

**topGeneNumber** Specify the number of top genes/TFs, based on the most linked RE DNA methylation sites of a given analysis type, for which to generate expression correlation heatmaps and tables. Defaults to 10.

### Value

Returns the MultiAssayExperiment object given as the TENETMultiAssayExperiment argument with an additional list of information named 'step7TopGenesExpressionCorrelationHeatmaps' in its metadata with the output of this function. This list is subdivided into hypermethGplus or hypomethGplus results as selected by the user, which are further subdivided into lists with data for the top overall genes and for top TF genes only. These contain the mirrored heatmaps displaying the expression correlation values for the expression of top genes/TFs as well as data frames with names and correlation values for each of those genes/TFs.

### Examples

```
## This example uses the example MultiAssayExperiment provided in the
## TENET.ExperimentHub package to create correlation heatmaps, and
## corresponding data frames, for the top 10 genes/TFs by number of linked
## hyper- or hypomethylated RE DNA methylation sites. Gene names will be
## retrieved from the rowRanges of the 'expression' SummarizedExperiment
## object in the example MultiAssayExperiment.

## Load the example TENET MultiAssayExperiment object
## from the TENET.ExperimentHub package
exampleTENETMultiAssayExperiment <-
  TENET.ExperimentHub::exampleTENETMultiAssayExperiment()

## Use the example dataset to create expression correlation heatmaps
returnValue <- step7TopGenesExpressionCorrelationHeatmaps(
  TENETMultiAssayExperiment = exampleTENETMultiAssayExperiment
)

## This example uses the example MultiAssayExperiment provided in the
## TENET.ExperimentHub package to create correlation heatmaps, and
## corresponding data frames, for the top 5 genes/TFs by number of linked
## hypomethylated RE DNA methylation sites only. Gene names will be retrieved
## from the rowRanges of the 'expression' SummarizedExperiment object in the
## example MultiAssayExperiment.

## Load the example TENET MultiAssayExperiment object
```

```
## from the TENET.ExperimentHub package
exampleTENETMultiAssayExperiment <-
  TENET.ExperimentHub::exampleTENETMultiAssayExperiment()

## Use the example dataset to create expression correlation heatmaps
returnValue <- step7TopGenesExpressionCorrelationHeatmaps(
  TENETMultiAssayExperiment = exampleTENETMultiAssayExperiment,
  hypermethGplusAnalysis = FALSE,
  topGeneNumber = 5
)
```

---

```
step7TopGenesOverlappingLinkedDNAMethylationSitesHeatmaps
```

*Generate binary heatmaps displaying which of the top genes/transcription factors share links with each of the unique regulatory element DNA methylation sites linked to at least one top gene/TF*

---

## Description

This function takes the top genes/TFs for each analysis type by number of linked RE DNA methylation sites identified by the step6DNAMethylationSitesPerGeneTabulation function, up to the number specified by the user, and identifies the unique RE DNA methylation sites linked to them, then generates two-color binary heatmaps displaying which of the top genes/TFs the RE DNA methylation sites are linked to, as well as data frames with that information.

## Usage

```
step7TopGenesOverlappingLinkedDNAMethylationSitesHeatmaps(
  TENETMultiAssayExperiment,
  geneAnnotationDataset = NA,
  hypermethGplusAnalysis = TRUE,
  hypomethGplusAnalysis = TRUE,
  topGeneNumber = 10
)
```

## Arguments

TENETMultiAssayExperiment

Specify a MultiAssayExperiment object containing expression and methylation SummarizedExperiment objects, such as one created by the TCGADownloader function. This MultiAssayExperiment object should also contain the results from the step5OptimizeLinks and step6DNAMethylationSitesPerGeneTabulation functions in its metadata.

geneAnnotationDataset

Specify a gene annotation dataset which is used to identify names for genes by their Ensembl IDs. The argument must be either a GRanges object (such as one

imported via `rtracklayer::import`) or a path to a GFF3 or GTF file. Both GENCODE and Ensembl annotations are supported. Other annotation datasets may work, but have not been tested. See the "Input data" section of the vignette for information on the required dataset format. Specify NA to use the names for genes listed in the "geneName" column of the `elementMetadata` of the `rowRanges` of the "expression" `SummarizedExperiment` object within the `TENETMultiAssayExperiment` object. Defaults to NA.

#### `hypermethGplusAnalysis`

Set to TRUE to create heatmaps and tables showing the linked RE DNA methylation sites for the top genes/TFs with the most hypermethylated RE DNA methylation sites with G+ links. Defaults to TRUE.

#### `hypomethGplusAnalysis`

Set to TRUE to create heatmaps and tables showing the linked RE DNA methylation sites for the top genes/TFs with the most hypomethylated RE DNA methylation sites with G+ links. Defaults to TRUE.

`topGeneNumber` Specify the number of top genes/TFs, based on the most linked RE DNA methylation sites of a given analysis type, for which to generate linked RE DNA methylation site heatmaps and tables. Defaults to 10.

### Value

Returns the `MultiAssayExperiment` object given as the `TENETMultiAssayExperiment` argument with an additional list of information named `'step7TopGenesOverlappingLinkedDNAMethylationSitesHeatmaps'` in its metadata with the output of this function. This list is subdivided into `hypermethGplus` or `hypomethGplus` results as selected by the user, which are further subdivided into lists with data for the top overall genes and for top TF genes only. These contain the binary heatmaps displaying the unique RE DNA methylation sites linked to the top genes/TFs in the columns and each of the top genes/TFs in the rows, with black indicating the given RE DNA methylation site is linked to the given gene/TF. Data frames are also included with this same data, with 1s indicating an RE DNA methylation site is linked to the gene/TF.

### Examples

```
## This example uses the example MultiAssayExperiment provided in the
## TENET.ExperimentHub package to create overlap heatmaps, and corresponding
## data frames, for the top 10 genes/TFs by number of linked hyper- or
## hypomethylated RE DNA methylation sites. Gene names will be retrieved
## from the rowRanges of the 'expression' SummarizedExperiment object in the
## example MultiAssayExperiment.

## Load the example TENET MultiAssayExperiment object
## from the TENET.ExperimentHub package
exampleTENETMultiAssayExperiment <-
  TENET.ExperimentHub::exampleTENETMultiAssayExperiment()

## Use the example dataset to create overlap heatmaps
returnValue <- step7TopGenesOverlappingLinkedDNAMethylationSitesHeatmaps(
  TENETMultiAssayExperiment = exampleTENETMultiAssayExperiment
)
```

```
## This example uses the example MultiAssayExperiment provided in the
## TENET.ExperimentHub package to create overlap heatmaps, and corresponding
## data frames, for the top 5 genes/TFs by number of linked hypomethylated
## RE DNA methylation sites only. Gene names will be retrieved from the
## rowRanges of the 'expression' SummarizedExperiment object in the example
## MultiAssayExperiment.

## Load the example TENET MultiAssayExperiment object
## from the TENET.ExperimentHub package
exampleTENETMultiAssayExperiment <-
  TENET.ExperimentHub::exampleTENETMultiAssayExperiment()

## Use the example dataset to create overlap heatmaps
returnValue <- step7TopGenesOverlappingLinkedDNAMethylationSitesHeatmaps(
  TENETMultiAssayExperiment = exampleTENETMultiAssayExperiment,
  hypermethGplusAnalysis = FALSE,
  topGeneNumber = 5
)
```

---

`step7TopGenesSurvival` *Perform Kaplan-Meier and Cox regression analyses to assess the association of top gene expression and linked RE DNA methylation site methylation with patient survival*

---

## Description

This function takes the top genes/transcription factors (TFs) by number of linked RE DNA methylation sites identified by the `step6DNAMethylationSitesPerGeneTabulation` function up to the number specified by the user and generates survival plots and tables with statistics from survival analyses assessing the survival association of the expression level of each gene as well as the methylation level of each RE DNA methylation site linked to them, using percentile cutoffs as specified by the user for Kaplan-Meier analyses.

## Usage

```
step7TopGenesSurvival(
  TENETMultiAssayExperiment,
  geneAnnotationDataset = NA,
  hypermethGplusAnalysis = TRUE,
  hypomethGplusAnalysis = TRUE,
  topGeneNumber = 10,
  vitalStatusData = NA,
  survivalTimeData = NA,
  highProportion = 0.5,
  lowProportion = 0.5,
  generatePlots = TRUE,
  coreCount = 1
)
```

## Arguments

### TENETMultiAssayExperiment

Specify a MultiAssayExperiment object containing expression and methylation SummarizedExperiment objects, such as one created by the TCGADownloader function. This MultiAssayExperiment object should also contain the results from the step2GetDifferentiallyMethylatedSites, step5OptimizeLinks, and step6DNAMethylationSitesPerGeneTabulation functions in its metadata. Additionally, the colData of the MultiAssay object must contain a 'vital\_status' and 'time' column, containing data on the patients' survival status and time to event/censorship, respectively.

### geneAnnotationDataset

Specify a gene annotation dataset which is used to identify names for genes by their Ensembl IDs. The argument must be either a GRanges object (such as one imported via `rtracklayer::import`) or a path to a GFF3 or GTF file. Both GENCODE and Ensembl annotations are supported. Other annotation datasets may work, but have not been tested. See the "Input data" section of the vignette for information on the required dataset format. Specify NA to use the names for genes listed in the "geneName" column of the elementMetadata of the rowRanges of the "expression" SummarizedExperiment object within the TENETMultiAssayExperiment object. Defaults to NA.

### hypermethGplusAnalysis

Set to TRUE to perform survival analyses on top genes/TFs by most hypermethylated RE DNA methylation sites with G+ links, as well as their linked RE DNA methylation sites.

### hypomethGplusAnalysis

Set to TRUE to perform survival analyses on top genes/TFs by most hypomethylated RE DNA methylation sites with G+ links, as well as their linked RE DNA methylation sites.

### topGeneNumber

Specify the number of top genes/TFs, based on the most linked RE DNA methylation sites of a given analysis type, for which to perform survival analyses. Defaults to 10.

### vitalStatusData

Specify the vital status data for samples in the TENETMultiAssayExperiment. Vital status should be given in the form of either "alive" or "dead" (case-insensitive), or 1 or 2, indicating that the sample was collected from a patient who was alive/censored or dead/reached the outcome of interest, respectively. These data can be given in a variety of forms, including a vector, data frame/ matrix, or a path to a file that contains the vital status data. If a vector is given, the names of the vector elements must correspond to the names of the samples in the rownames of the colData of the TENETMultiAssayExperiment object. If no names are provided for the vector, then the number of elements in the vector must equal the number of samples in the colData, and are assumed to align with the samples as they are ordered in the colData. If a data frame or matrix is given, its rownames must include the sample names as they appear in the colData of the TENETMultiAssayExperiment object, and its first column must include the vital status data. If a single string is provided, then it is assumed to be a path to a tab-delimited file containing vital status data in the second column, and the

names of the samples, again corresponding with the sample names in the colData, in the first column, which will be loaded as the row names. The first row of the file must contain column names. If this variable is set to NA, then the vital status data will be assumed to already be contained in the colData of the TENETMultiAssayExperiment under a column titled "vital\_status". Defaults to NA.

#### survivalTimeData

Specify the survival time data for samples in the TENETMultiAssayExperiment. Survival time should be given in the form of a numeric variable. These data can be given in a variety of forms, including a vector, data frame/matrix, or a path to a file that contains the survival time data. See the documentation for the vitalStatusData argument for more information. If this variable is set to NA, then the survival time data will be assumed to already be contained in the colData of the TENETMultiAssayExperiment under a column titled "time". Defaults to NA.

**highProportion** Set a number ranging from 0 to 1, indicating the proportion of all samples to include in the high expression/methylation group for Kaplan-Meier survival analyses. Defaults to 0.5.

**lowProportion** Set a number ranging from 0 to 1, indicating the proportion of all samples to include in the low expression/methylation group for Kaplan-Meier survival analyses. The total value of the highProportion and lowProportion arguments should not exceed 1. **Note:** If both this value and the highProportion value are set to 0.5, samples at exactly the 50th percentile will be assigned to the "Low" group. Defaults to 0.5.

**generatePlots** Set to TRUE to create and save plots displaying the Kaplan-Meier survival results for the genes/TFs of interest, as well as the RE DNA methylation sites linked to them. Defaults to TRUE.

**coreCount** Argument passed as the mc.cores argument to mclapply. See ?parallel::mclapply for more details. Defaults to 1.

### Value

Returns the MultiAssayExperiment object given as the TENETMultiAssayExperiment argument with an additional list of information named 'step7TopGenesSurvival' in its metadata with the output of this function. This list is subdivided into hypermethGplus or hypomethGplus results as selected by the user, which are further subdivided into lists with data for the top overall genes, and for top TF genes only. Each contains a list with data frames containing survival statistics for the top genes/TFs as well as their linked RE DNA methylation sites, from both Kaplan-Meier and Cox regression analyses. Additionally, they will also include a list of Kaplan-Meier plots if generatePlots is TRUE.

### Examples

```
## This example uses the example MultiAssayExperiment provided in the
## TENET.ExperimentHub package to perform Kaplan-Meier and Cox regression
## survival analyses for the top 10 genes/TFs, by number of linked hyper- or
## hypomethylated RE DNA methylation sites, as well as for all unique RE DNA
## methylation sites linked to those
```

```
## 10 genes/TFs. The vital status and survival time of patients will be
## taken from the "vital_status" and "time" columns present in the colData of
## the example MultiAssayExperiment. Gene names will be retrieved from the
## rowRanges of the 'expression' SummarizedExperiment object in the example
## MultiAssayExperiment. For Kaplan-Meier analyses, the patient samples with
## complete clinical information in the highest half of
## expression/methylation will be compared to the patient samples with
## complete clinical information in the lowest half. Kaplan-Meier plots will
## be saved for the genes and RE DNA methylation sites, and the analysis will
## be performed using one CPU core.
```

```
## Load the example TENET MultiAssayExperiment object
## from the TENET.ExperimentHub package
exampleTENETMultiAssayExperiment <-
  TENET.ExperimentHub::exampleTENETMultiAssayExperiment()
```

```
## Use the example dataset to do the survival analysis
returnValue <- step7TopGenesSurvival(
  TENETMultiAssayExperiment = exampleTENETMultiAssayExperiment
)
```

```
## This example uses the example MultiAssayExperiment provided in the
## TENET.ExperimentHub package to perform Kaplan-Meier and Cox regression
## survival analyses for only the top 5 genes/TFs, by number of linked
## hypomethylated RE DNA methylation sites, as well as for all unique
## RE DNA methylation sites linked to those 5 genes/TFs only. The vital
## status and survival time of patients will be
## taken from specific columns in a separate data frame with example patient
## data from the TENET.ExperimentHub package. Gene names will be retrieved
## from the rowRanges of the 'expression' SummarizedExperiment object in the
## example MultiAssayExperiment. For Kaplan-Meier analyses, the patient
## samples with complete clinical information in the highest quartile of
## expression/methylation will be compared to the patient samples with
## complete clinical information in the lowest quartile. Kaplan-Meier plots
## will not be saved for the genes and RE DNA methylation sites, and the
## analysis will be performed using 8 CPU cores.
```

```
## Load the example TENET MultiAssayExperiment object
## from the TENET.ExperimentHub package
exampleTENETMultiAssayExperiment <-
  TENET.ExperimentHub::exampleTENETMultiAssayExperiment()
```

```
## Again, this loads the data frame with example clinical data for patients
## in the TENET MultiAssayExperiment object from the TENET.ExperimentHub
## package
exampleTENETClinicalDataFrame <-
  TENET.ExperimentHub::exampleTENETClinicalDataFrame()
```

```
## Use the example datasets to do the survival analysis
returnValue <- step7TopGenesSurvival(
  TENETMultiAssayExperiment = exampleTENETMultiAssayExperiment,
  hypermethGplusAnalysis = FALSE,
  topGeneNumber = 5,
```



```

    vitalStatusData = exampleTENETClinicalDataFrame["vital_status"],
    survivalTimeData = exampleTENETClinicalDataFrame["time"],
    highProportion = 0.25,
    lowProportion = 0.25,
    generatePlots = FALSE,
    coreCount = 8
  )

```

---

### step7TopGenesTADTables

*Create tables using user-supplied topologically associating domain (TAD) information which identify the topologically associating domains containing each RE DNA methylation site linked to the top genes/transcription factors, as well as other genes in the same topologically associating domain as potential downstream targets*

---

### Description

This function takes the top genes/transcription factors (TFs) by number of linked RE DNA methylation sites identified by the step6DNAMethylationSitesPerGeneTabulation function up to the number specified by the user and generates tables with information for each of the RE DNA methylation sites linked to them for both of the hyper- or hypomethylated G+ analysis quadrants, as selected by the user. These tables note which of the top genes/TFs each RE DNA methylation site is linked to, as well as the total number and names of genes which happen to lie within the same topologically associating domain (TAD) of each RE DNA methylation site in each of the user-supplied TAD files.

### Usage

```

step7TopGenesTADTables(
  TENETMultiAssayExperiment,
  TADFiles,
  geneAnnotationDataset = NA,
  DNAMethylationArray = NA,
  hypermethGplusAnalysis = TRUE,
  hypomethGplusAnalysis = TRUE,
  topGeneNumber = 10,
  coreCount = 1
)

```

### Arguments

TENETMultiAssayExperiment

Specify a MultiAssayExperiment object containing expression and methylation SummarizedExperiment objects, such as one created by the TCGADownloader function. This MultiAssayExperiment object should also contain the results from the step5OptimizeLinks and step6DNAMethylationSitesPerGeneTabulation functions in its metadata.

TADFiles	Specify a data frame, matrix, or GRanges object with information on the TAD compartments of interest, organized in a bed-like manner (see <a href="https://genome.ucsc.edu/FAQ/FAQformat.html#format1">https://genome.ucsc.edu/FAQ/FAQformat.html#format1</a> ), or a path to a directory that contains bed-like files that contain such TAD information. If a path is provided, multiple bed-formatted TAD files can be included in the specified directory. The files may optionally be compressed (.gz/.bz2/.xz). <b>Note:</b> Data frames and matrices will be assumed to use 1-indexed coordinates; <code>rtracklayer::import.bed</code> converts coordinates to 1-indexed upon import.
geneAnnotationDataset	Specify a gene annotation dataset which is used to identify names for genes by their Ensembl IDs. The argument must be either a GRanges object (such as one imported via <code>rtracklayer::import</code> ) or a path to a GFF3 or GTF file. Both GENCODE and Ensembl annotations are supported. Other annotation datasets may work, but have not been tested. See the "Input data" section of the vignette for information on the required dataset format. Specify NA to use the names for genes listed in the "geneName" column of the elementMetadata of the rowRanges of the "expression" SummarizedExperiment object within the TENETMultiAssayExperiment object. Defaults to NA.
DNAMethylationArray	Specify the name of a DNA methylation probe array supported by the sesameData package (see <code>?sesameData::sesameData_getManifestGRanges</code> ). If an array is specified, RE DNA methylation sites and their locations in that array's manifest are cross-referenced with RE DNA methylation site IDs included in the rownames of the methylation dataset provided in the "methylation" SummarizedExperiment object within the TENETMultiAssayExperiment object, and only those overlapping will be considered for analysis. If set to NA, all RE DNA methylation sites with locations listed in the rowRanges of the "methylation" SummarizedExperiment object are used. Defaults to NA.
hypermethGplusAnalysis	Set to TRUE to create TAD tables for the RE DNA methylation sites linked to the top genes/TFs by most hypermethylated RE DNA methylation sites with G+ links.
hypomethGplusAnalysis	Set to TRUE to create TAD tables for the RE DNA methylation sites linked to the top genes/TFs by most hypomethylated RE DNA methylation sites with G+ links.
topGeneNumber	Specify the number of top genes/TFs, based on the most linked RE DNA methylation sites of a given analysis type, for which to generate TAD tables for the RE DNA methylation sites linked to those genes. Defaults to 10.
coreCount	Argument passed as the <code>mc.cores</code> argument to <code>mcmapply</code> . See <code>?parallel::mcmapply</code> for more details. Defaults to 1.

## Value

Returns the MultiAssayExperiment object given as the TENETMultiAssayExperiment argument with an additional list of information named 'step7TopGenesTADTables' in its metadata with the output of this function. This list is subdivided into hypermethGplus or hypomethGplus results as selected by the user, which are further subdivided into data frames with data for the unique RE

DNA methylation sites linked to the top overall genes, and for top TF genes only. This includes the top genes/TFs each RE DNA methylation site is linked to, and, for each TAD file, if an RE DNA methylation site was found in a TAD in that file, as well as the gene count and identities of other genes found in the same TAD as each RE DNA methylation site.

## Examples

```
## This example uses the example MultiAssayExperiment provided in the
## TENET.ExperimentHub package to do overlapping for all unique RE DNA
## methylation sites linked to the top 10 genes, by number of linked hyper-
## or hypomethylated RE DNA methylation sites, using a GRanges object
## containing topologically associating domain (TAD) data from the
## TENET.ExperimentHub package. Gene names and locations, and the locations
## of RE DNA methylation sites, will be retrieved from the rowRanges of the
## 'expression' and 'methylation' SummarizedExperiment objects in the
## example MultiAssayExperiment. The analysis will be performed using one
## CPU core.

## Load the example TENET MultiAssayExperiment object
## from the TENET.ExperimentHub package
exampleTENETMultiAssayExperiment <-
  TENET.ExperimentHub::exampleTENETMultiAssayExperiment()

## Load the example TAD GRanges object from the TENET.ExperimentHub package
exampleTENETTADRegions <- TENET.ExperimentHub::exampleTENETTADRegions()

## Use the example datasets to do the TAD overlapping
returnValue <- step7TopGenesTADTables(
  TENETMultiAssayExperiment = exampleTENETMultiAssayExperiment,
  TADFiles = exampleTENETTADRegions
)

## This example uses the example MultiAssayExperiment provided in the
## TENET.ExperimentHub package to do overlapping for all unique RE DNA
## methylation sites linked to the top 5 genes only, by number of linked
## hypomethylated RE DNA methylation sites only, with bed-like files
## containing topologically associating domain (TAD) data located in the
## user's R working directory. This analysis will be done using gene names
## and their locations which are included in the "geneName" column of the
## elementMetadata of the rowRanges of the "expression" SummarizedExperiment
## object within the TENETMultiAssayExperiment object, and RE DNA
## methylation sites and their locations which are included in the HM450
## array retrieved via the sesameData package. The analysis will be
## performed using 8 CPU cores.

## Load the example TENET MultiAssayExperiment object
## from the TENET.ExperimentHub package
exampleTENETMultiAssayExperiment <-
  TENET.ExperimentHub::exampleTENETMultiAssayExperiment()

## Use the example dataset to do the TAD overlapping
returnValue <- step7TopGenesTADTables(
```

```

TENETMultiAssayExperiment = exampleTENETMultiAssayExperiment,
TADFiles = ".",
DNAMethylationArray = "HM450",
hypermethGplusAnalysis = FALSE,
topGeneNumber = 5,
coreCount = 8
)

```

---

### step7TopGenesUCSCBedFiles

*Create bed-formatted interact files which can be loaded on the UCSC Genome Browser to display links between top genes and transcription factors and their linked RE DNA methylation sites*

---

## Description

This function takes the top genes/transcription factors (TFs) by number of linked RE DNA methylation sites identified by the `step6DNAMethylationSitesPerGeneTabulation` function, up to the number specified by the user, and generates bed-formatted interact files (see <https://genome.ucsc.edu/goldenPath/help/interact.html>) that can be uploaded to the UCSC Genome Browser (<https://genome.ucsc.edu>) to visualize the links between each of the top specified genes/TFs and the RE DNA methylation sites linked to them for both of the hyper- or hypomethylated G+ analysis quadrants, as selected by the user.

## Usage

```

step7TopGenesUCSCBedFiles(
  TENETMultiAssayExperiment,
  outputDirectory,
  geneAnnotationDataset = NA,
  DNAMethylationArray = NA,
  hypermethGplusAnalysis = TRUE,
  hypomethGplusAnalysis = TRUE,
  topGeneNumber = 10
)

```

## Arguments

**TENETMultiAssayExperiment**  
Specify a `MultiAssayExperiment` object containing expression and methylation `SummarizedExperiment` objects, such as one created by the `TCGADownloader` function. This `MultiAssayExperiment` object should also contain the results from the `step5OptimizeLinks` and `step6DNAMethylationSitesPerGeneTabulation` functions in its metadata.

**outputDirectory**  
Specify the path to the output directory in which to save the .bed files created by this function. The directory will be created if it does not exist.

**geneAnnotationDataset**

Specify a gene annotation dataset which is used to identify names for genes by their Ensembl IDs. The argument must be either a GRanges object (such as one imported via `rtracklayer::import`) or a path to a GFF3 or GTF file. Both GENCODE and Ensembl annotations are supported. Other annotation datasets may work, but have not been tested. See the "Input data" section of the vignette for information on the required dataset format. Specify NA to use the names for genes listed in the "geneName" column of the elementMetadata of the rowRanges of the "expression" SummarizedExperiment object within the TENETMultiAssayExperiment object. Defaults to NA.

**DNAMethylationArray**

Specify the name of a DNA methylation probe array supported by the sesameData package (see `?sesameData::sesameData_getManifestGRanges`). If an array is specified, RE DNA methylation sites and their locations in that array's manifest are cross-referenced with RE DNA methylation site IDs included in the rownames of the methylation dataset provided in the "methylation" SummarizedExperiment object within the TENETMultiAssayExperiment object, and only those overlapping will be considered for analysis. If set to NA, all RE DNA methylation sites with locations listed in the rowRanges of the "methylation" SummarizedExperiment object are used. Defaults to NA.

**hypermethGplusAnalysis**

Set to TRUE to create interact files showing links between the top genes/TFs by most RE hypermethylated RE DNA methylation sites with G+ links, and these linked RE DNA methylation sites. Defaults to TRUE.

**hypomethGplusAnalysis**

Set to TRUE to create interact files showing links between the top genes/TFs by most hypomethylated RE DNA methylation sites with G+ links, and these linked RE DNA methylation sites. Defaults to TRUE.

**topGeneNumber**

Specify the number of top genes/TFs, based on the most linked RE DNA methylation sites of a given analysis type, for which to generate interact files showing the links between those genes and each of their linked RE DNA methylation sites. Defaults to 10.

**Value**

Outputs .bed formatted interact files to upload to the UCSC Genome Browser to the specified output directory. These files display the interactions between the top genes/TFs and their linked RE DNA methylation sites for the given analysis types. Returns a list of lists named after each selected analysis type, each containing the file paths to the created .bed files for top genes and top TFs for that analysis type.

**Examples**

```
## This example uses the example MultiAssayExperiment provided in the
## TENET.ExperimentHub package to create UCSC Genome Browser interact files
## for the top 10 genes/TFs by number of linked hyper- or hypomethylated RE
## DNA methylation sites. The interact files for the top genes/TFs will be
## saved in the user's working directory. Gene names and locations, and the
## locations of RE DNA methylation sites, will be retrieved from the
```

```

## rowRanges of the 'expression' and 'methylation' SummarizedExperiment
## objects in the example MultiAssayExperiment.

## Load the example TENET MultiAssayExperiment object
## from the TENET.ExperimentHub package
exampleTENETMultiAssayExperiment <-
  TENET.ExperimentHub::exampleTENETMultiAssayExperiment()

## Use the example dataset to create and save the UCSC Genome Browser
## interact files
filePaths <- step7TopGenesUCSCBedFiles(
  TENETMultiAssayExperiment = exampleTENETMultiAssayExperiment,
  outputDirectory = "."
)

## Get the path to the bed file for the top TFs by number of
## hypomethylated G+ RE DNA methylation sites
filePaths$hypoGplus$topTFs

## This example uses the example MultiAssayExperiment provided in the
## TENET.ExperimentHub package to create UCSC Genome Browser interact files
## for the top 5 genes/TFs by number of linked hypomethylated RE DNA
## methylation sites only. The interact files for the top genes/TFs will be
## saved in the user's working directory. Gene names and locations will be
## retrieved from the rowRanges of the 'expression' and 'methylation'
## SummarizedExperiment objects in the example MultiAssayExperiment. RE DNA
## methylation site IDs and locations will be retrieved from the HM450 array
## via the sesameData package.

## Load the example TENET MultiAssayExperiment object
## from the TENET.ExperimentHub package
exampleTENETMultiAssayExperiment <-
  TENET.ExperimentHub::exampleTENETMultiAssayExperiment()

## Use the example dataset to create and save the UCSC Genome Browser
## interact files
filePaths <- step7TopGenesUCSCBedFiles(
  TENETMultiAssayExperiment = exampleTENETMultiAssayExperiment,
  outputDirectory = ".",
  DNAMethylationArray = "HM450",
  hypermethGplusAnalysis = FALSE,
  topGeneNumber = 5
)

## Get the path to the bed file for the top TFs by number of
## hypomethylated G+ RE DNA methylation sites.
## Note: Since we performed analyses only using TFs in the step 3 function,
## the top genes are all TFs, so topTFs will be NA here, and topGenes
## should be used instead.
filePaths$hypoGplus$topGenes

```

---

step7TopGenesUserPeakOverlap

*Identify if RE DNA methylation sites linked to top genes/transcription factors are located within a specific distance of specified genomic regions*

---

## Description

This function takes the top genes/transcription factors (TFs) by number of linked RE DNA methylation sites identified by the step6DNAMethylationSitesPerGeneTabulation function, up to the number specified by the user, and identifies if the RE DNA methylation sites linked to those genes/TFs from the hyper- or hypomethylated G+ analysis quadrants are found in the vicinity of genomic regions (peaks) of interest, supplied by the user in the form of .bed, .narrowPeak, .broadPeak, and/or gappedPeak files in a specified directory or given as a data frame or GRanges object.

## Usage

```
step7TopGenesUserPeakOverlap(
  TENETMultiAssayExperiment,
  peakData,
  geneAnnotationDataset = NA,
  DNAMethylationArray = NA,
  hypermethGplusAnalysis = TRUE,
  hypomethGplusAnalysis = TRUE,
  topGeneNumber = 10,
  distanceFromREDNAMethylationSites = 100,
  coreCount = 1
)
```

## Arguments

TENETMultiAssayExperiment	Specify a MultiAssayExperiment object containing expression and methylation SummarizedExperiment objects, such as one created by the TCGADownloader function. This MultiAssayExperiment object should also contain the results from the step5OptimizeLinks and step6DNAMethylationSitesPerGeneTabulation functions in its metadata.
peakData	Specify a data frame, matrix, or GRanges object with genomic regions (peaks) of interest, organized in a bed-like manner (see <a href="https://genome.ucsc.edu/FAQ/FAQformat.html#format1">https://genome.ucsc.edu/FAQ/FAQformat.html#format1</a> ), or a path to a directory containing .bed, .narrowPeak, .broadPeak, and/or .gappedPeak files with peaks of interest. The files may optionally be compressed (.gz/.bz2/.xz).
geneAnnotationDataset	Specify a gene annotation dataset which is used to identify names for genes by their Ensembl IDs. The argument must be either a GRanges object (such as one imported via rtracklayer::import) or a path to a GFF3 or GTF file. Both

GENCODE and Ensembl annotations are supported. Other annotation datasets may work, but have not been tested. See the "Input data" section of the vignette for information on the required dataset format. Specify NA to use the names for genes listed in the "geneName" column of the elementMetadata of the rowRanges of the "expression" SummarizedExperiment object within the TENETMultiAssayExperiment object. Defaults to NA.

#### DNAMethylationArray

Specify the name of a DNA methylation probe array supported by the sesameData package (see `?sesameData::sesameData_getManifestGRanges`). If an array is specified, RE DNA methylation sites and their locations in that array's manifest are cross-referenced with RE DNA methylation site IDs included in the rownames of the methylation dataset provided in the "methylation" SummarizedExperiment object within the TENETMultiAssayExperiment object, and only those overlapping will be considered for analysis. If set to NA, all RE DNA methylation sites with locations listed in the rowRanges of the "methylation" SummarizedExperiment object are used. Defaults to NA.

#### hypermethGplusAnalysis

Set to TRUE to create data frames with the peak overlap information for the unique hypermethylated RE DNA methylation sites linked to the top genes/TFs by most hypermethylated RE DNA methylation sites with G+ links. Defaults to TRUE.

#### hypomethGplusAnalysis

Set to TRUE to create data frames with the peak overlap information for the unique hypomethylated RE DNA methylation sites linked to the top genes/TFs by most hypomethylated RE DNA methylation sites with G+ links. Defaults to TRUE.

**topGeneNumber** Specify the number of top genes/TFs, based on the most linked RE DNA methylation sites of a given analysis type, for which to generate data showing overlap with user peak files for the RE DNA methylation sites linked to those genes. Defaults to 10.

#### distanceFromREDNAMethylationSites

Specify either 0 or a positive integer in base pairs to be the distance from the linked RE DNA methylation sites within which an RE DNA methylation site will be considered to overlap a peak. Defaults to 100.

**coreCount** Argument passed as the `mc.cores` argument to `mclapply`. See `?parallel::mclapply` for more details. Defaults to 1.

### Value

Returns the MultiAssayExperiment object given as the TENETMultiAssayExperiment argument with an additional list of information named 'step7TopGenesUserPeakOverlap' in its metadata with the output of this function. This list is subdivided into hypermethGplus or hypomethGplus results as selected by the user, which are further subdivided into data frames with data for the unique RE DNA methylation sites linked to the top overall genes, and for top TF genes only. Each of these data frames contain a row for each of the unique RE DNA methylation sites linked to the top genes/TFs for the specified analysis types. These note the locations of the RE DNA methylation sites and the specified search windows, whether the RE DNA methylation site is linked to each of the top



genes/TFs, and whether each RE DNA methylation site was found within the specified distance to any peak in each of the user's peak files.

## Examples

```
## This example uses the example MultiAssayExperiment provided in the
## TENET.ExperimentHub package to do peak overlapping for all unique RE DNA
## methylation sites linked to the top 10 genes, by number of linked hyper-
## or hypomethylated RE DNA methylation sites, using a GRanges object
## containing the genomic coordinates of peaks of interest. Gene names, and
## the locations of RE DNA methylation sites, will be retrieved from the
## rowRanges of the 'expression' and 'methylation' SummarizedExperiment
## objects in the example MultiAssayExperiment. The analysis will be
## performed using one CPU core.

## Load the example TENET MultiAssayExperiment object
## from the TENET.ExperimentHub package
exampleTENETMultiAssayExperiment <-
  TENET.ExperimentHub::exampleTENETMultiAssayExperiment()

## Load the example peak GRanges object from the TENET.ExperimentHub package
exampleTENETPeakRegions <- TENET.ExperimentHub::exampleTENETPeakRegions()

## Use the example datasets to do the peak overlapping
returnValue <- step7TopGenesUserPeakOverlap(
  TENETMultiAssayExperiment = exampleTENETMultiAssayExperiment,
  peakData = exampleTENETPeakRegions
)

## This example uses the example MultiAssayExperiment provided in the
## TENET.ExperimentHub package to do peak overlapping for all unique RE DNA
## methylation sites linked to the top 5 genes only, by number of linked
## hypomethylated RE DNA methylation sites only, using bed-like files
## containing the genomic coordinates of peaks of interest in the user's R
## working directory. Gene names will be retrieved from the rowRanges of the
## 'expression' SummarizedExperiment object in the example
## MultiAssayExperiment, and RE DNA methylation sites and their locations
## retrieved from the HM450 array via the sesameData package. A window of
## 500 base pairs will be used to identify if the RE DNA methylation sites
## lie within the vicinity of peaks. The analysis will be performed using 8
## CPU cores.

## Load the example TENET MultiAssayExperiment object from the
## TENET.ExperimentHub package
exampleTENETMultiAssayExperiment <-
  TENET.ExperimentHub::exampleTENETMultiAssayExperiment()

## Use the example dataset to do the peak overlapping
returnValue <- step7TopGenesUserPeakOverlap(
  TENETMultiAssayExperiment = exampleTENETMultiAssayExperiment,
  peakData = ".",
  DNAMethylationArray = "HM450",
  hypermethGplusAnalysis = FALSE,
```

```

    topGeneNumber = 5,
    distanceFromREDNAMethylationSites = 500,
    coreCount = 8
  )

```

---

TCGADownloader

*Download TCGA gene expression, DNA methylation, and clinical datasets and compile them into a MultiAssayExperiment object*

---

## Description

This function downloads and compiles TCGA gene expression and DNA methylation datasets, as well as clinical data primarily intended for use with the TENET package. This simplifies the TCGAbiolinks download functions, identifies samples with matching gene expression and DNA methylation data, and can also remove duplicate tumor samples taken from the same patient donor. Data are compiled into a MultiAssayExperiment object, which is returned and optionally saved in an .rda file at the path specified by the outputFile argument.

## Usage

```

TCGADownloader(
  rawDataDownloadDirectory,
  GDCDownloadMethod = "api",
  filesPerChunk = 10,
  TCGAStudyAbbreviation,
  RNASeqWorkflow,
  RNASeqLog2Normalization = TRUE,
  removeDupTumor = TRUE,
  matchingExpAndMetSamples = TRUE,
  clinicalSurvivalData = "combined",
  outputFile = NA
)

```

## Arguments

rawDataDownloadDirectory

Specify the path to the directory where TCGAbiolinks should download data. Note that this dataset can be very sizable.

GDCDownloadMethod

The method to use when downloading data from the Genomic Data Commons (GDC). Passed as the method argument to TCGAbiolinks' GDCdownload function. The available options are "api" and "client"; the default is "api". The "api" method works on all operating systems, but it does not retry the download of incomplete or corrupted files, so TCGADownloader must be manually rerun in this case. The "client" method is more reliable, but it requires Windows, macOS (Apple Silicon only), or Ubuntu (64-bit x86 only), or manual installation of the GDC Data Transfer Tool Client (which must be in the command search path).

- filesPerChunk** The number of data files to download at once when using the "api" download method. Passed as the `files.per.chunk` argument to TCGAbiolinks' `GDCdownload` function. Lower values may improve download reliability, but higher values may increase download speed. Defaults to 10.
- TCGAStudyAbbreviation** Input a four-letter code for a TCGA dataset for which to download data. See <https://gdc.cancer.gov/resources-tcga-users/tcga-code-tables/tcga-study-abbreviations> for more information and a complete list of options.
- RNASeqWorkflow** Select the type of RNA-seq data to download. For TENET purposes, choose either "STAR - FPKM", "STAR - FPKM-UQ", "STAR - FPKM-UQ - old formula", or "STAR - TPM". "STAR - Counts" can also be used but is not recommended for TENET analyses. See [https://docs.gdc.cancer.gov/Data/Bioinformatics\\_Pipelines/Expression\\_mRNA\\_Pipeline/](https://docs.gdc.cancer.gov/Data/Bioinformatics_Pipelines/Expression_mRNA_Pipeline/) for more information on these.
- RNASeqLog2Normalization** Set to TRUE to do log2 normalization of RNA-seq expression values. Defaults to TRUE.
- removeDupTumor** Set to TRUE to remove duplicate tumor samples taken from the same subject, leaving only one sample per subject in alphanumeric order. **Note:** To properly create a dataset for use with TENET, both the `removeDupTumor` and `matchingExpAndMetSamples` arguments must be set to TRUE. Defaults to TRUE.
- matchingExpAndMetSamples** Select the type of expression and methylation sample data matching to perform. If set to TRUE, only samples with at least one methylation and expression sample annotated to their patient, will be kept. If set to FALSE, all samples will be kept, including those without matching expression and methylation data. **Note:** To properly create a dataset for use with TENET, both the `removeDupTumor` and `matchingExpAndMetSamples` arguments must be set to TRUE. Defaults to TRUE.
- clinicalSurvivalData** Select how clinical data should be prepared from the TCGA data, with respect to patient vital status and survival time. Valid options include "bcrBiotabPatient" to use survival data contained only in the 'patient' data in the BCR Biotab files downloaded using TCGAbiolinks, or "combined", which uses clinical information from the 'patient' and 'follow\_up' datasets in the BCR Biotab files, as well as data from the BCR XML files. Data from the same patient in each of the datasets are combined, and the data with the most recent (highest patient survival time) entry for each patient are kept. Additionally, for both options, the 'days\_to\_last\_followup' and 'days\_to\_death' variables are collapsed into a single time variable, which is combined with the other patient clinical data in the 'patient' BCR Biotab data. See <https://bioconductor.org/packages/devel/bioc/vignettes/TCGAbiolinks/inst/doc/clinical.html> for more information on how TCGAbiolinks prepares different clinical datasets. Defaults to "combined".
- outputFile** Specify the path to an .rda file in which to save the MultiAssayExperiment object with downloaded datasets. If set to NA or undefined, this results in the function only returning the MultiAssayExperiment object and not saving it. Defaults to NA.

**Value**

Returns and/or saves to an .rda file a MultiAssayExperiment object with expression and methylation data included SummarizedExperiment objects within the MultiAssayExperiment object, as well as clinical data included in the colData of the MultiAssayExperiment object.

**Examples**

```
## Download a TCGA LUAD dataset with log2-normalized
## FPKM-UQ expression values from tumor and adjacent normal tissue samples
## with matching expression and methylation data and keeping only one tumor
## sample from each patient. Additionally, survival data will be combined
## from three clinical datasets downloaded by TCGAbiolinks. Raw data files
## will be saved to the working directory, and the processed dataset will
## be returned as a variable.
TCGADataset <- TCGADownloader(
  rawDataDownloadDirectory = ".",
  TCGAStudyAbbreviation = "LUAD",
  RNASeqWorkflow = "STAR - FPKM-UQ"
)

## Another example, which downloads a TCGA BRCA dataset with FPKM expression
## values with no normalization and no duplicate samples removed. Survival
## data are derived from just the patient BCR Biotab file downloaded by
## TCGAbiolinks. Both raw data files and an .rda file containing the data
## as a MultiAssayExperiment object will be saved to the working directory.
## Note: This functionality is useful for downloading samples from
## TCGA but will *not* work for a TENET assay due to the lack of sample
## matching and duplicate tumor sample removal.
TCGADownloader(
  rawDataDownloadDirectory = ".",
  TCGAStudyAbbreviation = "BRCA",
  RNASeqWorkflow = "STAR - FPKM",
  RNASeqLog2Normalization = FALSE,
  removeDupTumor = FALSE,
  matchingExpAndMetSamples = FALSE,
  clinicalSurvivalData = "bcrBiotabPatient",
  outputFile = "BRCAMultiAssayExperimentObject.rda"
)
```

---

TENET

---

*TENET (Tracing regulatory Element Networks using Epigenetic Traits)*


---

**Description**

TENET identifies key transcription factors and regulatory elements linked to a specific cell type by finding significantly correlated differences in gene expression and regulatory element methylation between case and control input datasets, and identifying the top genes by number of significant RE

DNA methylation site links. It also includes many additional tools to aid in the visualization and analysis of the results, including plots displaying and comparing methylation and expression data and RE DNA methylation site link counts, survival analysis, TF motif searching in the vicinity of linked RE DNA methylation sites, custom TAD and peak overlap analysis, and UCSC Genome Browser track file generation. A utility function is also provided to download methylation, expression, and patient survival data from The Cancer Genome Atlas (TCGA) for use in TENET or other analyses.

### Author(s)

**Maintainer:** Rhie Lab at the University of Southern California <rhie@ucla.edu>

Authors:

- Daniel Mullen ([ORCID](#))
- Zexun Wu ([ORCID](#))
- Ethan Nelson-Moore ([ORCID](#))
- Suhm Rhie ([ORCID](#))

### See Also

Useful links:

- <https://github.com/rhie/TENET>
- Report bugs at <https://github.com/rhie/TENET/issues>

---

TENETCacheAllData	<i>Cache all online datasets required by TENET examples and optional features</i>
-------------------	---

---

### Description

This function locally caches all online TENET and SeSAmE datasets required by TENET examples and optional features (TENET.ExperimentHub objects used in examples, TENET.AnnotationHub datasets used in step 1, and SeSAmE datasets loaded via the DNAMethylationArray argument). The main purpose of this function is to enable the use of TENET in an HPC cluster environment where compute nodes do not have internet access. In this case, you must run TENETCacheAllData() once while connected to the internet before using TENET examples or these optional features.

### Usage

```
TENETCacheAllData()
```

### Value

Returns NULL.

### Examples

```
TENETCacheAllData()
```

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