

Package ‘dnaEPICO’

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Title dnaEPICO: Analysis Pipeline for Illumina DNA Methylation Array Data

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Description A modular and reproducible workflow for preprocessing and analysing Illumina DNA methylation array data from the EPICv2, EPIC, and 450K platforms. The package integrates quality control, probe filtering, cell-type deconvolution, phenotype preparation, generalised linear models, linear mixed-effects models, and automated report generation. It builds on established Bioconductor infrastructure and wraps commonly used tools including 'minfi', 'ENmix', and 'watermelon', with support for both local execution and high-performance computing workflows.

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URL <https://github.com/paulYRP/dnaEPICO>

BugReports <https://github.com/paulYRP/dnaEPICO/issues>

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analyzeSvaEnmix	<i>Analyze surrogate variables against Sentrix chip and position factors</i>
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Description

Fit linear models for each surrogate variable against Sentrix chip and Sentrix position, perform backward elimination with MASS::dropterm(), and return the in-memory analysis objects.

Usage

```
analyzeSvaEnmix(
  sva,
  RGSet,
  SentrixIDColumn = "Sentrix_ID",
  SentrixPositionColumn = "Sentrix_Position",
  verbose = FALSE,
  logs = FALSE,
  log_dir = NULL,
  log_file = "log_analyzeSvaEnmix.txt"
)
```

Arguments

sva	Numeric matrix of surrogate variables with samples in rows.
RGSet	An RGChannelSet aligned with sva.
SentrixIDColumn	Character. Name of the chip identifier column in SummarizedExperiment::colData(RGSet).
SentrixPositionColumn	Character. Name of the chip position column in SummarizedExperiment::colData(RGSet).
verbose	Logical. If TRUE, emit progress messages with message().
logs	Logical. If TRUE, write the same messages to a log file.
log_dir	Character or NULL. Directory used for the log file when logs = TRUE.
log_file	Character. File name used when logs = TRUE.

Value

A list with class "dnaEPICO_svaEnmix_analysis" containing the aligned Sentrix factors, full and reduced linear models, and ANOVA tables.

Examples

```
ex <- dnaEPICO:::exampleSvaAnalysisStateDnaEpico()
analysis_data <- analyzeSvaEnmix(
  sva = ex$sva,
  RGSet = ex$RGSet,
  SentrixIDColumn = "Sentrix_ID",
  SentrixPositionColumn = "Sentrix_Position",
  verbose = FALSE,
  logs = FALSE
)
analysis_data$K
```

annotateMethylationGLMM_T1T2Summaries

Annotate longitudinal mixed-effects summary tables with array annotation metadata

Description

Merge phenotype-specific longitudinal summary tables with probe annotation metadata and return a single annotated result table.

Usage

```
annotateMethylationGLMM_T1T2Summaries(
  modelSummaries,
  annotationObject,
  annotationCols = c("Name", "chr", "pos", "UCSC_RefGene_Group", "UCSC_RefGene_Name",
    "Relation_to_Island", "GencodeV41_Group"),
  verbose = FALSE,
  logs = FALSE,
```

```

log_dir = NULL,
log_file = "log_methylationGLMM_T1T2.txt"
)

```

Arguments

modelSummaries Object returned by summarizeMethylationGLMM_T1T2Models() or a named list of summary data frames.

annotationObject Character package/object name, annotation data frame, or annotation object understood by minfi::getAnnotation().

annotationCols Character vector or comma-separated string of annotation columns to append.

verbose Logical. If TRUE, emit progress messages with message().

logs Logical. If TRUE, write the same messages to a log file.

log_dir Character or NULL. Directory used for the log file when logs = TRUE.

log_file Character. File name used when logs = TRUE.

Value

A list with class "dnaEPICO_methylationGLMM_T1T2_annotation" containing the annotated summary table and any requested annotation columns that were unavailable in the chosen annotation object.

Examples

```

ex <- dnaEPICO::exampleMethylationGLMMStateDnaEpico()
annotation_data <- annotateMethylationGLMM_T1T2Summaries(
  modelSummaries = ex$modelSummaries,
  annotationObject = ex$annotationData,
  annotationCols = "Name,chr,pos",
  verbose = FALSE,
  logs = FALSE
)
names(annotation_data)

```

annotateMethylationGLM_T1Summaries

Annotate one-timepoint GLM summary tables with array annotation metadata

Description

Merge phenotype-specific CpG summary tables with probe annotation metadata and return a single annotated result table.

Usage

```

annotateMethylationGLM_T1Summaries(
  modelSummaries,
  annotationObject,
  annotationCols = c("Name", "chr", "pos", "UCSC_RefGene_Group", "UCSC_RefGene_Name",
    "Relation_to_Island", "GencodeV41_Group"),
  verbose = FALSE,
  logs = FALSE,
  log_dir = NULL,
  log_file = "log_methylationGLM_T1.txt"
)

```

Arguments

modelSummaries Object returned by `summarizeMethylationGLM_T1Models()` or a named list of CpG summary data frames.

annotationObject Character package/object name, annotation data frame, or annotation object understood by `minfi::getAnnotation()`.

annotationCols Character vector or comma-separated string of annotation columns to append.

verbose Logical. If TRUE, emit progress messages with `message()`.

logs Logical. If TRUE, write the same messages to a log file.

log_dir Character or NULL. Directory used for the log file when `logs = TRUE`.

log_file Character. File name used when `logs = TRUE`.

Value

A list with class "dnaEPICO_methylationGLM_T1_annotation" containing the annotated summary table and any requested annotation columns that were unavailable in the chosen annotation object.

Examples

```

ex <- dnaEPICO::exampleMethylationGLMStateDnaEpico()
annotation_data <- annotateMethylationGLM_T1Summaries(
  modelSummaries = ex$modelSummaries,
  annotationObject = ex$annotationData,
  annotationCols = "Name,chr,pos",
  verbose = FALSE,
  logs = FALSE
)
names(annotation_data)

```

 assessSamplesMinfiEwasWater

Assess sample quality before sample filtering

Description

Compute minfi QC metrics and detection P values, identify failed samples using the requested threshold, and return the assessment as a single object.

Usage

```
assessSamplesMinfiEwasWater(
  rawData,
  RGSet,
  qcCutoff = 10.5,
  detPtype = "m+u",
  detPThreshold = 0.05,
  verbose = FALSE,
  logs = FALSE,
  log_dir = NULL,
  log_file = "log_assessSamplesMinfiEwasWater.txt"
)
```

Arguments

rawData	Object returned by buildRawMinfiEwasWater().
RGSet	An RGChannelSet aligned with rawData.
qcCutoff	Numeric. Cutoff passed to minfi::plotQC() when the QC plot is drawn.
detPtype	Character. Detection P-value mode passed to minfi::detectionP(). Common values in minfi workflows are "m+u" and "negative". The default used here is "m+u".
detPThreshold	Numeric. Samples with mean detection P value above this threshold are marked as failed.
verbose	Logical. If TRUE, emit progress messages with message().
logs	Logical. If TRUE, write the same messages to a log file.
log_dir	Character or NULL. Directory used for the log file when logs = TRUE.
log_file	Character. File name used when logs = TRUE.

Value

A list with class "dnaEPICO_minfiEwasWater_assessment" containing the QC object, detection P matrix, mean detection P values, and failed sample identifiers.

Examples

```
ex <- dnaEPICO:::exampleMinfiBaseDataDnaEpico()
raw_data <- buildRawMinfiEwasWater(ex$RGSet, verbose = FALSE, logs = FALSE)
assessment <- assessSamplesMinfiEwasWater(
  rawData = raw_data,
```

```

RGSet = ex$RGSet,
detPThreshold = 1,
verbose = FALSE,
logs = FALSE
)
names(assessment)

```

```
buildClockFoundationInputsPreprocessingPheno
```

Build Clock Foundation input tables from preprocessingPheno data

Description

Prepare the beta and phenotype tables commonly exported for Clock Foundation style downstream workflows, without writing them to disk.

Usage

```

buildClockFoundationInputsPreprocessingPheno(
  beta,
  pheno,
  SampleID = "Sample_Name",
  sexColumn = "Sex",
  verbose = FALSE,
  logs = FALSE,
  log_dir = NULL,
  log_file = "log_buildClockFoundationInputsPreprocessingPheno.txt"
)

```

Arguments

beta	Numeric matrix of beta values with probes in rows and samples in columns.
pheno	Phenotype data frame aligned with the beta matrix columns.
SampleID	Character. Name of the phenotype sample identifier column.
sexColumn	Character. Name of the phenotype sex column.
verbose	Logical. If TRUE, emit progress messages with <code>message()</code> .
logs	Logical. If TRUE, write the same messages to a log file.
log_dir	Character or NULL. Directory used for the log file when <code>logs = TRUE</code> .
log_file	Character. File name used when <code>logs = TRUE</code> .

Value

A list with class `"dnaEPICO_preprocessingPheno_clock"` containing `betaCSV` and `phenoCF`.

Examples

```
ex <- dnaEPICO:::examplePreprocessingPhenoStateDnaEpico()
clock_inputs <- buildClockFoundationInputsPreprocessingPheno(
  beta = ex$timepointData$data[["1"]]$beta,
  pheno = ex$timepointData$data[["1"]]$pheno,
  SampleID = "Sample_Name",
  sexColumn = "Sex",
  verbose = FALSE,
  logs = FALSE
)
names(clock_inputs)
```

```
buildRawMinfiEwasWater
```

Build raw minfi preprocessing objects

Description

Create a raw MethylSet, RatioSet, and genome-mapped object from an RGChannelSet, and return them together in a single structured object.

Usage

```
buildRawMinfiEwasWater(
  RGSet,
  verbose = FALSE,
  logs = FALSE,
  log_dir = NULL,
  log_file = "log_buildRawMinfiEwasWater.txt"
)
```

Arguments

RGSet	An RGChannelSet.
verbose	Logical. If TRUE, emit progress messages with message().
logs	Logical. If TRUE, write the same messages to a log file.
log_dir	Character or NULL. Directory used for the log file when logs = TRUE.
log_file	Character. File name used when logs = TRUE.

Value

A list with class "dnaEPICO_minfiEwasWater_raw" containing MSet, RatioSet, and GSet.

Examples

```
ex <- dnaEPICO:::exampleMinfiBaseDataDnaEpico()
raw_data <- buildRawMinfiEwasWater(
  RGSet = ex$RGSet,
  verbose = FALSE,
  logs = FALSE
)
```

```
)
names(raw_data)
```

```
collectSignificantCpGsMethylationGLM_T1
  Collect significant CpG coefficient tables from fitted one-timepoint
  GLMs
```

Description

Collect the raw coefficient tables for CpGs whose phenotype main effect or interaction p-value passes the requested threshold.

Usage

```
collectSignificantCpGsMethylationGLM_T1(
  modelResults,
  pvalThreshold = 0.05,
  interactionTerm = NULL,
  verbose = FALSE,
  logs = FALSE,
  log_dir = NULL,
  log_file = "log_methylationGLM_T1.txt"
)
```

Arguments

modelResults	Object returned by fitMethylationGLM_T1Models().
pvalThreshold	Numeric. Threshold applied to phenotype main-effect or interaction p-values.
interactionTerm	Character or NULL. Optional interaction term.
verbose	Logical. If TRUE, emit progress messages with message().
logs	Logical. If TRUE, write the same messages to a log file.
log_dir	Character or NULL. Directory used for the log file when logs = TRUE.
log_file	Character. File name used when logs = TRUE.

Value

A list with class "dnaEPICO_methylationGLM_T1_significant_cpgs".

Examples

```
ex <- dnaEPICO:::exampleMethylationGLMStateDnaEpico()
significant_cpgs <- collectSignificantCpGsMethylationGLM_T1(
  modelResults = ex$modelResults,
  pvalThreshold = 1,
  verbose = FALSE,
  logs = FALSE
)
names(significant_cpgs)
```

```
collectSignificantInteractionsMethylationGLMM_T1T2
```

Collect significant longitudinal model terms from fitted mixed-effects models

Description

Collect raw coefficient tables for CpGs whose phenotype main effect or requested interaction p-value passes the chosen threshold.

Usage

```
collectSignificantInteractionsMethylationGLMM_T1T2(
  modelResults,
  pvalThreshold = 0.05,
  interactionTerm = NULL,
  verbose = FALSE,
  logs = FALSE,
  log_dir = NULL,
  log_file = "log_methylationGLMM_T1T2.txt"
)
```

Arguments

<code>modelResults</code>	Object returned by <code>fitMethylationGLMM_T1T2Models()</code> .
<code>pvalThreshold</code>	Numeric. Threshold applied to the extracted phenotype or interaction p-values.
<code>interactionTerm</code>	Character or NULL. Optional interaction term. When NULL, phenotype main effects are used.
<code>verbose</code>	Logical. If TRUE, emit progress messages with <code>message()</code> .
<code>logs</code>	Logical. If TRUE, write the same messages to a log file.
<code>log_dir</code>	Character or NULL. Directory used for the log file when <code>logs = TRUE</code> .
<code>log_file</code>	Character. File name used when <code>logs = TRUE</code> .

Value

A list with class `"dnaEPICO_methylationGLMM_T1T2_significant"` containing the retained coefficient tables for each phenotype.

Examples

```
ex <- dnaEPICO:::exampleMethylationGLMMStateDnaEpico()
significant_hits <- collectSignificantInteractionsMethylationGLMM_T1T2(
  modelResults = ex$modelResults,
  pvalThreshold = 1,
  verbose = FALSE,
  logs = FALSE
)
names(significant_hits)
```

```
combineTimepointsPreprocessingPheno
```

Combine selected timepoints for downstream longitudinal modeling

Description

Combine selected timepoints that were already aligned by `splitTimepointsPreprocessingPheno()` into the wide phenotype-plus-beta objects used by downstream longitudinal models.

Usage

```
combineTimepointsPreprocessingPheno(
  timepointData,
  combineTimepoints = "1,2",
  verbose = FALSE,
  logs = FALSE,
  log_dir = NULL,
  log_file = "log_combineTimepointsPreprocessingPheno.txt"
)
```

Arguments

<code>timepointData</code>	Object returned by <code>splitTimepointsPreprocessingPheno()</code> .
<code>combineTimepoints</code>	Character vector or comma-separated string of timepoints to combine.
<code>verbose</code>	Logical. If TRUE, emit progress messages with <code>message()</code> .
<code>logs</code>	Logical. If TRUE, write the same messages to a log file.
<code>log_dir</code>	Character or NULL. Directory used for the log file when <code>logs = TRUE</code> .
<code>log_file</code>	Character. File name used when <code>logs = TRUE</code> .

Value

A list with class "dnaEPICO_preprocessingPheno_combined" containing the combined phenotype table, merged phenotype-plus-beta table, selected timepoints, and output suffix.

Examples

```
ex <- dnaEPICO::examplePreprocessingPhenoStateDnaEpico()
combined_data <- combineTimepointsPreprocessingPheno(
  timepointData = ex$timepointData,
  combineTimepoints = "1,2",
  verbose = FALSE,
  logs = FALSE
)
combined_data$suffix
```

Description

The dnaEPICO package provides a structured workflow for preprocessing and analyzing Illumina DNA methylation array data, including quality control, normalization, cell-type estimation, surrogate-variable analysis, phenotype preparation, cross-sectional modeling, longitudinal mixed-effects modeling, and PDF reporting.

Details

The package supports two complementary usage styles:

- interactive use, where functions return structured in-memory result objects for inspection and composition; and
- file-based pipeline use, where the same functions can write logs, plots, tables, and serialized objects when `saveOutputs = TRUE`.

The main high-level entry points are:

- `preprocessingMinfiEwasWater()`
- `svaEnmix()`
- `preprocessingPheno()`
- `methylationGLM_T1()`
- `methylationGLMM_T1T2()`
- `dnamReport()`

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See Also

Useful links:

- <https://github.com/pauLYRP/dnaEPICO>
- Report bugs at <https://github.com/pauLYRP/dnaEPICO/issues>

 dnaEPICO_dnamReport-class

Result class returned by dnamReport

Description

Objects of class "dnaEPICO_dnamReport" are list-based results returned by [dnamReport\(\)](#). They combine the prepared report inputs, render result, and final status metadata into one convenience object.

Structure

preparedReport Object returned by [prepareDnamReportInputs\(\)](#).

renderResult Structured render metadata created by [dnamReport\(\)](#).

status Final status string such as "rendered", "skipped", or "failed".

outputFile Path to docs/index.html.

errorMessage Final render error or skip message when available, otherwise NULL.

logFile Resolved path to the optional log file, or NULL when logging was disabled.

See Also

[dnamReport\(\)](#)

dnaEPICO_dnamReport_prepared-class

Result class returned by prepareDnamReportInputs

Description

Objects of class "dnaEPICO_dnamReport_prepared" are list-based results returned by [prepareDnamReportInputs\(\)](#). They capture normalized report paths, available figures, and logging metadata before rendering.

Structure

output Requested output file name.

outputDir Normalized report output directory.

outputFile Normalized full path to the intended report output file.

figDir Normalized directory used by the report template for copied figures.

figureInventory Named list describing the available figures for each report section.

missingFigureDirectories Character vector of expected figure directories that were not present at preparation time.

logFile Resolved path to the optional log file, or NULL when logging was disabled.

See Also

[prepareDnamReportInputs\(\)](#)

 dnaEPICO_dnamReport_render-class

Result class returned by renderDnamReport

Description

Objects of class "dnaEPICO_dnamReport_render" are list-based results returned by [renderDnamReport\(\)](#). They describe whether a prepared report was rendered, skipped, or failed.

Structure

preparedReport The input object supplied to [renderDnamReport\(\)](#).

status Render status string such as "rendered", "skipped", or "failed".

renderedFile Normalized path to the rendered PDF file when rendering succeeded, otherwise NULL.

errorMessage Render error or skip message when available, otherwise NULL.

logFile Resolved path to the optional log file, or NULL when logging was disabled.

See Also

[renderDnamReport\(\)](#)

dnaEPICO_methylationGLMM_T1T2-class

Result class returned by methylationGLMM_T1T2

Description

Objects of class "dnaEPICO_methylationGLMM_T1T2" are list-based results returned by [methylationGLMM_T1T2\(\)](#). They collect the prepared longitudinal analysis table, fitted mixed models, summaries, diagnostics, annotations, and optional saved files.

Structure

preparedData Object returned by [prepareMethylationGLMM_T1T2Data\(\)](#).

modelFits Object returned by [fitMethylationGLMM_T1T2Models\(\)](#).

modelSummaries Object returned by [summarizeMethylationGLMM_T1T2Models\(\)](#).

significantInteractions Object returned by [collectSignificantInteractionsMethylationGLMM_T1T2\(\)](#).

diagnosticPlots Object returned by [plotMethylationGLMM_T1T2Diagnostics\(\)](#).

annotation Object returned by [annotateMethylationGLMM_T1T2Summaries\(\)](#).

savedFiles Object returned by [writeMethylationGLMM_T1T2Outputs\(\)](#) when `saveOutputs = TRUE`, otherwise NULL.

See Also

[methylationGLMM_T1T2\(\)](#)

dnaEPICO_methylationGLM_T1-class

Result class returned by methylationGLM_T1

Description

Objects of class "dnaEPICO_methylationGLM_T1" are list-based results returned by [methylationGLM_T1\(\)](#). They collect the prepared analysis table, fitted models, summaries, diagnostics, annotations, and optional saved files.

Structure

preparedData Object returned by [prepareMethylationGLM_T1Data\(\)](#).
distributionPlots Object returned by [plotMethylationGLM_T1Distributions\(\)](#).
modelFits Object returned by [fitMethylationGLM_T1Models\(\)](#).
modelSummaries Object returned by [summarizeMethylationGLM_T1Models\(\)](#).
significantCpGs Object returned by [collectSignificantCpGsMethylationGLM_T1\(\)](#).
diagnosticPlots Object returned by [plotMethylationGLM_T1Diagnostics\(\)](#).
annotation Object returned by [annotateMethylationGLM_T1Summaries\(\)](#).
savedFiles Object returned by [writeMethylationGLM_T1Outputs\(\)](#) when `saveOutputs = TRUE`, otherwise NULL.

See Also

[methylationGLM_T1\(\)](#)

dnaEPICO_preprocessingMinfiEwasWater-class

Result class returned by preprocessingMinfiEwasWater

Description

Objects of class "dnaEPICO_preprocessingMinfiEwasWater" are list-based results returned by [preprocessingMinfiEwasWater\(\)](#). They are lightweight S3-style containers rather than formal S4 classes.

Structure

targets Filtered phenotype table aligned to the retained samples.
RGSet Filtered RGChannelSet used in downstream preprocessing.
rawData Object returned by [buildRawMinfiEwasWater\(\)](#).
assessment Object returned by [assessSamplesMinfiEwasWater\(\)](#).
sexData Object returned by [predictSexMinfiEwasWater\(\)](#).
normData Object returned by [normalizeMinfiEwasWater\(\)](#).
filterData Object returned by [filterProbesMinfiEwasWater\(\)](#).
metricsData Object returned by [extractMetricsMinfiEwasWater\(\)](#).
lcData Object returned by [estimateLCMinfiEwasWater\(\)](#).
logFile Resolved path to the optional log file, or NULL when logging was disabled.

See Also

[preprocessingMinfiEwasWater\(\)](#)

dnaEPICO_preprocessingPheno-class

Result class returned by preprocessingPheno

Description

Objects of class "dnaEPICO_preprocessingPheno" are list-based results returned by [preprocessingPheno\(\)](#). They describe the phenotype data, methylation matrices, timepoint splits, longitudinal merges, and optional exported files.

Structure

pheno Phenotype table read from phenoFile.

metricsData Object returned by [loadMetricsPreprocessingPheno\(\)](#).

timepointData Object returned by [splitTimepointsPreprocessingPheno\(\)](#).

combinedData Object returned by [combineTimepointsPreprocessingPheno\(\)](#).

clockFoundation Object returned by [buildClockFoundationInputsPreprocessingPheno\(\)](#).

savedFiles Object returned by [writePreprocessingPhenoOutputs\(\)](#) when saveOutputs = TRUE, otherwise NULL.

logFile Resolved path to the optional log file, or NULL when logging was disabled.

See Also

[preprocessingPheno\(\)](#)

dnaEPICO_svaEnmix-class

Result class returned by svaEnmix

Description

Objects of class "dnaEPICO_svaEnmix" are list-based results returned by [svaEnmix\(\)](#). They collect the loaded inputs, surrogate-variable results, association-analysis summaries, and optional file outputs.

Structure

targets Phenotype table read from phenoFile after any optional row subsetting.

RGSet Loaded RGChannelSet with sample names reset to match targets.

svaData Object returned by [estimateSvaEnmixControls\(\)](#).

mergedPheno Phenotype table returned by [mergeSvaTargetsEnmix\(\)](#) after surrogate variables were appended.

analysisData Object returned by [analyzeSvaEnmix\(\)](#).

plotFiles Named list of TIFF output paths for the SVA figures when `saveOutputs = TRUE`, otherwise NULL entries for plots not written.

savedFiles Object returned by `writeSvaEnmixOutputs()` when `saveOutputs = TRUE`, otherwise NULL.

logFile Resolved path to the optional log file, or NULL when logging was disabled.

See Also

[svaEnmix\(\)](#)

dnamReport

Generate a DNA methylation dashboard report

Description

Generate a DNA methylation dashboard report

Usage

```
dnamReport(
  outputDir = "reports",
  phenoTab = NULL,
  enmixTab = file.path("figures", "preprocessingMinfiEwasWater", "enmix"),
  qcTab = file.path("figures", "preprocessingMinfiEwasWater", "qc"),
  svaTab = file.path("figures", "svaEnmix"),
  metricTab = file.path("figures", "preprocessingMinfiEwasWater", "metrics"),
  glmTab = NULL,
  lmerTab = NULL,
  logTab = outputDir,
  verbose = FALSE,
  logs = FALSE,
  projectName = "dnaEPICO",
  detPPath = NULL,
  detPThreshold = 0.01,
  cpgDetectionPath = NULL,
  sampleDetectionPath = NULL,
  logoPath = system.file("extdata", "dnaEPICO.svg", package = "dnaEPICO"),
  imagePattern = "\\.(png|jpg|jpeg|gif|webp|svg|tif|tiff)$",
  recursive = TRUE
)
```

Arguments

<code>outputDir</code>	Character. Directory where the Quarto project is written.
<code>phenoTab</code>	Character or NULL. CSV file shown in the Data tab. When NULL, the path is inferred from the Makefile output layout.
<code>enmixTab</code>	Character. Directory containing ENmix quality-control figures.
<code>qcTab</code>	Character. Directory containing Quality Control figures.
<code>svaTab</code>	Character. Directory containing Batch Effect or SVA figures.

metricTab	Character. Directory containing Metrics figures.
glmTab	Character or NULL. CSV file shown in the GLM Analysis tab. When NULL, the path is inferred from the Makefile output layout.
lmerTab	Character or NULL. CSV file shown in the LMER Analysis tab. When NULL, the path is inferred from the Makefile output layout.
logTab	Character. Directory containing workflow logs shown in the Logs tab.
verbose	Logical. If TRUE, emit progress messages.
logs	Logical. If TRUE, write a report log.
projectName	Character. Name used for the generated Quarto project.
detPPath	Character or NULL. RData file containing the detection P-value matrix object detP, used to build the quality-control tables. When NULL, the path is inferred from the Makefile output layout.
detPThreshold	Numeric. Detection P-value threshold used when summarising the detP matrix.
cpgDetectionPath	Character or NULL. Optional fallback CpG detection summary CSV.
sampleDetectionPath	Character or NULL. Optional fallback sample detection summary CSV.
logoPath	Character. Path to the navbar logo. Defaults to the packaged inst/extdata/dnaEPICO.svg asset.
imagePattern	Character. Regular expression used to identify image files inside the section directories.
recursive	Logical. If TRUE, search section directories recursively.

Value

A list with class "dnaEPICO_dnamReport".

Examples

```
report_root <- file.path(tempdir(), "dnaepico-dnam-report")
pheno_file <- file.path(
  report_root,
  "data",
  "model1",
  "preprocessingMinfiEwasWater",
  "phenoLC.csv"
)
dir.create(dirname(pheno_file), recursive = TRUE, showWarnings = FALSE)
utils::write.csv(
  data.frame(
    UID = c("sample1", "sample2"),
    Timepoint = c(1, 2),
    Sex = c("F", "M")
  ),
  pheno_file,
  row.names = FALSE
)

result <- dnamReport(
  outputDir = file.path(report_root, "reports", "model1"),
  phenoTab = pheno_file,

```

```

enmixTab = file.path(
  report_root,
  "figures",
  "model1",
  "preprocessingMinfiEwasWater",
  "enmix"
),
qcTab = file.path(
  report_root,
  "figures",
  "model1",
  "preprocessingMinfiEwasWater",
  "qc"
),
svaTab = file.path(report_root, "figures", "model1", "svaEnmix"),
metricTab = file.path(
  report_root,
  "figures",
  "model1",
  "preprocessingMinfiEwasWater",
  "metrics"
),
logTab = file.path(report_root, "logs", "model1")
)
result$status

```

estimateLC

Estimate saliva cell proportions from DNA methylation beta values

Description

Estimate cell-type proportions with the saliva reference panels bundled in dnaEPIC0. This function keeps the original estimateLC() interface used by the package while using the internal reference files distributed in inst/extdata.

Usage

```
estimateLC(meth, ref, constrained = FALSE)
```

Arguments

meth	Numeric matrix of beta values with CpGs in rows and samples in columns. Row names must contain probe identifiers compatible with the selected reference.
ref	Character. Reference panel name. Supported values are "saliva" and "salivaEPIC".
constrained	Logical. If TRUE, estimated cell proportions are constrained to sum to one.

Value

A data.table with one row per sample and one column per estimated cell type.

References

Murat K, et al. Ewastools: Infinium Human Methylation BeadChip pipeline for population epigenetics integrated into Galaxy. *GigaScience*. 2020;9(5):giaa049. Houseman EA, Accomando WP, Koestler DC, et al. DNA methylation arrays as surrogate measures of cell mixture distribution. *BMC Bioinformatics*. 2012;13:86. Reinius LE, Acevedo N, Joerink M, et al. Differential DNA methylation in purified human blood cells: implications for cell lineage and studies on disease susceptibility. *PLoS One*. 2012;7(7):e41361. Bakulski KM, Feinberg JI, Andrews SV, et al. DNA methylation of cord blood cell types: applications for mixed cell birth studies. *Epigenetics*. 2016;11(5):354-362. de Goede OM, Razzaghi HR, Price EM, et al. Nucleated red blood cells impact DNA methylation and expression analyses of cord blood hematopoietic cells. *Clinical Epigenetics*. 2015;7:95. Gervin K, Salas LA, Bakulski KM, et al. Cell type specific DNA methylation in cord blood: a 450K reference data set and cell count-based validation of estimated cell type composition. *Epigenetics*. 2016;11(9):690-698. Gervin K, Salas LA, Bakulski KM, et al. Systematic evaluation and validation of reference and library selection methods for deconvolution of cord blood DNA methylation data. *bioRxiv*. 2019. doi:10.1101/570457. Salas LA, Koestler DC, Butler RA, et al. An optimized library for reference-based deconvolution of whole-blood biospecimens assayed using the Illumina HumanMethylationEPIC BeadArray. *Genome Biology*. 2018;19:64. Heiss JA, Just AC, Brenner H. Training a model for estimating leukocyte composition using whole-blood DNA methylation and cell counts as reference. *Epigenomics*. 2017;9(1):13-20. Middleton LYM, Dou J, Mill J, et al. Saliva cell type DNA methylation reference panel for epidemiology studies in children. 2020.

Examples

```
ref_file <- system.file("extdata", "saliva.txt", package = "dnaEPIC")
ref_panel <- as.matrix(utils::read.table(ref_file))
meth <- ref_panel[1:20, , drop = FALSE]
colnames(meth) <- c("sample1", "sample2")
estimateLC(
  meth = meth,
  ref = "saliva",
  constrained = FALSE
)
```

```
estimateLCMinfiEwasWater
```

Estimate cell composition for preprocessing MinfiEwasWater

Description

Estimate cell proportions from beta values using `estimateLC()` for saliva reference panels or `ENmix::estimateCellProp()` for other supported references, then merge the estimates into the phenotype table.

Usage

```
estimateLCMinfiEwasWater(
  beta,
  targets,
  lcRef = "salivaEPIC",
  phenoOrder = "Sample_Name;Timepoint;Sex;PredSex;Basename;Sentry_ID;Sentry_Position",
```

```

    constrained = FALSE,
    verbose = FALSE,
    logs = FALSE,
    log_dir = NULL,
    log_file = "log_estimateLCMinfiEwasWater.txt"
  )

```

Arguments

beta	Numeric matrix of beta values with probes in rows and samples in columns.
targets	Phenotype data frame aligned with the columns of beta.
lcRef	Character. Cell-composition reference. Internal saliva-based references supported through estimateLC() are "saliva" and "salivaEPIC". Other references are passed to ENmix::estimateCellProp().
phenoOrder	Character vector or semicolon-separated string describing the phenotype columns that should appear first in the merged phenoLC output.
constrained	Logical. Passed to estimateLC() when an internal saliva reference is used. If TRUE, estimated proportions are constrained to sum to one.
verbose	Logical. If TRUE, emit progress messages with message().
logs	Logical. If TRUE, write the same messages to a log file.
log_dir	Character or NULL. Directory used for the log file when logs = TRUE.
log_file	Character. File name used when logs = TRUE.

Value

A list with class "dnaEPICO_minfiEwasWater_lc" containing the cell proportion matrix, merged phenotype table, reference name, and method used.

Examples

```

ref_file <- system.file("extdata", "saliva.txt", package = "dnaEPICO")
beta <- as.matrix(utils::read.table(ref_file))[1:20, , drop = FALSE]
colnames(beta) <- c("sample1", "sample2")
targets <- data.frame(
  Sample_Name = colnames(beta),
  Timepoint = c("T1", "T2"),
  stringsAsFactors = FALSE
)
lc_data <- estimateLCMinfiEwasWater(
  beta = beta,
  targets = targets,
  lcRef = "saliva",
  phenoOrder = "Sample_Name;Timepoint"
)
stopifnot(is.data.frame(lc_data$phenoLC))

```

```
estimateSvaEnmixControls
```

Estimate surrogate variables from ENmix control probes

Description

Run `ENmix::ctrlsva()` on an `RGChannelSet` and return the surrogate variable matrix as an in-memory object.

Usage

```
estimateSvaEnmixControls(
  RGSet,
  ctrlSvaPercVar = 0.9,
  ctrlSvaFlag = 1,
  verbose = FALSE,
  logs = FALSE,
  log_dir = NULL,
  log_file = "log_estimateSvaEnmixControls.txt"
)
```

Arguments

<code>RGSet</code>	An <code>RGChannelSet</code> .
<code>ctrlSvaPercVar</code>	Numeric. Proportion of variance explained by control probes, passed to <code>ENmix::ctrlsva()</code> .
<code>ctrlSvaFlag</code>	Integer. Control-probe flag passed to <code>ENmix::ctrlsva()</code> .
<code>verbose</code>	Logical. If <code>TRUE</code> , emit progress messages with <code>message()</code> .
<code>logs</code>	Logical. If <code>TRUE</code> , write the same messages to a log file.
<code>log_dir</code>	Character or <code>NULL</code> . Directory used for the log file when <code>logs = TRUE</code> .
<code>log_file</code>	Character. File name used when <code>logs = TRUE</code> .

Value

A list with class `"dnaEPICO_svaEnmix_sva"` containing the surrogate variable matrix and the parameters used to estimate it.

Examples

```
ex <- dnaEPICO::exampleMinfiBaseDataDnaEpic()
sva_data <- estimateSvaEnmixControls(
  RGSet = ex$RGSet,
  ctrlSvaPercVar = 0.5,
  ctrlSvaFlag = 1,
  verbose = FALSE,
  logs = FALSE
)
sva_data$K
```

extractMake	<i>Copy dnaEPICO Makefile to a user directory</i>
-------------	---

Description

Copies the example Makefile pipeline shipped with dnaEPICO to a user-specified directory for local execution or modification.

Usage

```
extractMake(destDir, overwrite = FALSE)
```

Arguments

destDir	Character. Destination directory where the Makefile will be copied.
overwrite	Logical. Whether to overwrite an existing Makefile in destDir. The default is FALSE.

Value

Character scalar containing the path to the copied Makefile.

Examples

```
tmp <- file.path(tempdir(), "dnaEPICO-make-example")
dir.create(tmp, recursive = TRUE, showWarnings = FALSE)
makefile_path <- extractMake(
  destDir = tmp,
  overwrite = TRUE
)
stopifnot(file.exists(makefile_path))
```

extractMetricsMinfiEwasWater	<i>Extract beta, M, and copy-number matrices from a filtered object</i>
------------------------------	---

Description

Extract beta, M, and copy-number matrices from a filtered object

Usage

```
extractMetricsMinfiEwasWater(
  filteredData,
  verbose = FALSE,
  logs = FALSE,
  log_dir = NULL,
  log_file = "log_extractMetricsMinfiEwasWater.txt"
)
```

Arguments

filteredData	Object returned by filterProbesMinfiEwasWater().
verbose	Logical. If TRUE, emit progress messages with message().
logs	Logical. If TRUE, write the same messages to a log file.
log_dir	Character or NULL. Directory used for the log file when logs = TRUE.
log_file	Character. File name used when logs = TRUE.

Value

A list with class "dnaEPICO_minfiEwasWater_metrics" containing beta, m, and cn.

Examples

```
ex <- dnaEPICO:::exampleMinfiMetricsStateDnaEpico()
metrics_data <- extractMetricsMinfiEwasWater(
  filteredData = ex$filteredData,
  verbose = FALSE,
  logs = FALSE
)
names(metrics_data)
```

filterProbesMinfiEwasWater

Filter probes from a normalized methylation object

Description

Apply detection P-value, chromosome, SNP, and cross-reactive probe filters to the primary normalized object and return the filtered result.

Usage

```
filterProbesMinfiEwasWater(
  normData,
  RGSet,
  pvalThreshold = 0.01,
  chrToRemove = "chrX,chrY",
  snpsToRemove = "SBE,CpG",
  mafThreshold = 0.1,
  crossReactivePath,
  detPtype = "m+u",
  verbose = FALSE,
  logs = FALSE,
  log_dir = NULL,
  log_file = "log_filterProbesMinfiEwasWater.txt"
)
```

Arguments

normData	Object returned by <code>normalizeMinfiEwasWater()</code> .
RGSet	Filtered <code>RGChannelSet</code> aligned with <code>normData</code> .
pvalThreshold	Numeric. Probes must have detection P values below this threshold in all samples to be retained.
chrToRemove	Character vector or comma-separated string of chromosome names to remove, for example "chrX, chrY".
snpsToRemove	Character vector or comma-separated string of SNP probe types to remove, for example "SBE, CpG".
mafThreshold	Numeric. Minor allele frequency threshold passed to <code>minfi::dropLociWithSnps()</code> .
crossReactivePath	Character. Path to a CSV file containing a <code>ProbeID</code> column of cross-reactive probes to remove.
detPtype	Character. Detection P-value mode passed to <code>minfi::detectionP()</code> for the probe filter. Common values in minfi workflows are "m+u" and "negative".
verbose	Logical. If TRUE, emit progress messages with <code>message()</code> .
logs	Logical. If TRUE, write the same messages to a log file.
log_dir	Character or NULL. Directory used for the log file when <code>logs = TRUE</code> .
log_file	Character. File name used when <code>logs = TRUE</code> .

Value

A list with class "dnaEPICO_minfiEwasWater_filter" containing the filtered object and counts for each filtering stage.

Examples

```
ex <- dnaEPICO:::exampleMinfiWorkflowStateDnaEpico()
filtered_data <- filterProbesMinfiEwasWater(
  normData = ex$normData,
  RGSet = ex$sampleData$RGSet,
  pvalThreshold = 1,
  chrToRemove = "chrY",
  snpsToRemove = "SBE",
  mafThreshold = 1,
  crossReactivePath = ex$crossReactivePath,
  detPtype = "m+u",
  verbose = FALSE,
  logs = FALSE
)
filtered_data$counts[["crossReactive"]]
```

```
filterSamplesMinfiEwasWater
```

Filter failed samples from an RGSet and phenotype table

Description

Remove failed samples identified during sample assessment and return the filtered RGChannelSet together with the aligned phenotype table.

Usage

```
filterSamplesMinfiEwasWater(
  RGSet,
  targets,
  failedSamples = character(0),
  SampleID = "Sample_Name",
  verbose = FALSE,
  logs = FALSE,
  log_dir = NULL,
  log_file = "log_filterSamplesMinfiEwasWater.txt"
)
```

Arguments

RGSet	An RGChannelSet.
targets	Data frame containing phenotype information.
failedSamples	Character vector of sample identifiers to remove.
SampleID	Character. Name of the sample identifier column in targets.
verbose	Logical. If TRUE, emit progress messages with message().
logs	Logical. If TRUE, write the same messages to a log file.
log_dir	Character or NULL. Directory used for the log file when logs = TRUE.
log_file	Character. File name used when logs = TRUE.

Value

A list with class "dnaEPICO_minfiEwasWater_samples" containing the filtered RGSet, aligned phenotype table, and failed sample identifiers.

Examples

```
ex <- dnaEPICO::exampleMinfiBaseDataDnaEpico()
filtered_samples <- filterSamplesMinfiEwasWater(
  RGSet = ex$RGSet,
  targets = ex$targets,
  failedSamples = ex$targets$Sample_Name[1],
  SampleID = "Sample_Name",
  verbose = FALSE,
  logs = FALSE
)
nrow(filtered_samples$targets)
```

```
fitMethylationGLMM_T1T2Models
```

Fit CpG-wise mixed-effects models for longitudinal methylation analyses

Description

Fit one linear mixed-effects model per CpG for each phenotype requested in the object returned by `prepareMethylationGLMM_T1T2Data()`.

Usage

```
fitMethylationGLMM_T1T2Models(
  preparedData,
  nCores = 1L,
  libPath = NULL,
  lmeLibs = "lme4,lmerTest",
  verbose = FALSE,
  logs = FALSE,
  log_dir = NULL,
  log_file = "log_methylationGLMM_T1T2.txt"
)
```

Arguments

<code>preparedData</code>	Object returned by <code>prepareMethylationGLMM_T1T2Data()</code> .
<code>nCores</code>	Integer. Number of worker processes to use.
<code>libPath</code>	Character vector or NULL. Optional library paths forwarded to worker processes.
<code>lmeLibs</code>	Character vector or comma-separated string of package names to check on worker processes. The default is <code>"lme4,lmerTest"</code> .
<code>verbose</code>	Logical. If TRUE, emit progress messages with <code>message()</code> .
<code>logs</code>	Logical. If TRUE, write the same messages to a log file.
<code>log_dir</code>	Character or NULL. Directory used for the log file when <code>logs = TRUE</code> .
<code>log_file</code>	Character. File name used when <code>logs = TRUE</code> .

Value

A list with class `"dnaEPIC0_methylationGLMM_T1T2_models"` containing fitted model lists, model formulas, and counts of failed CpG fits.

Examples

```
ex <- dnaEPIC0::exampleMethylationGLMMStateDnaEpic0()
model_results <- fitMethylationGLMM_T1T2Models(
  preparedData = ex$preparedData,
  nCores = 1,
  verbose = FALSE,
  logs = FALSE
)
names(model_results$fits)
```

fitMethylationGLM_T1Models

Fit CpG-wise Gaussian GLMs for one-timepoint methylation analyses

Description

Fit one Gaussian GLM per CpG for each phenotype requested in the object returned by prepareMethylationGLM_T1Data.

Usage

```
fitMethylationGLM_T1Models(
  preparedData,
  nCores = 1L,
  libPath = NULL,
  glmLibs = "glm2",
  verbose = FALSE,
  logs = FALSE,
  log_dir = NULL,
  log_file = "log_methylationGLM_T1.txt"
)
```

Arguments

preparedData	Object returned by prepareMethylationGLM_T1Data().
nCores	Integer. Number of worker processes to use.
libPath	Character vector or NULL. Optional library paths forwarded to worker processes.
glmLibs	Character vector or comma-separated string of package names to check on worker processes. The default is "glm2".
verbose	Logical. If TRUE, emit progress messages with message().
logs	Logical. If TRUE, write the same messages to a log file.
log_dir	Character or NULL. Directory used for the log file when logs = TRUE.
log_file	Character. File name used when logs = TRUE.

Value

A list with class "dnaEPICO_methylationGLM_T1_models" containing fitted model lists, model formulas, and counts of failed CpG fits.

Examples

```
ex <- dnaEPICO::exampleMethylationGLMStateDnaEpico()
model_results <- fitMethylationGLM_T1Models(
  preparedData = ex$preparedData,
  nCores = 1,
  verbose = FALSE,
  logs = FALSE
)
names(model_results$fits)
```

```
loadMetricsPreprocessingPheno
```

Load methylation metric matrices for preprocessingPheno

Description

Load the metric matrices generated by `preprocessingMinfiEwasWater()` and return them as a single in-memory object for downstream phenotype alignment.

Usage

```
loadMetricsPreprocessingPheno(
  betaPath,
  mPath,
  cnPath,
  verbose = FALSE,
  logs = FALSE,
  log_dir = NULL,
  log_file = "log_loadMetricsPreprocessingPheno.txt"
)
```

Arguments

<code>betaPath</code>	Character. Path to the saved beta-value object.
<code>mPath</code>	Character. Path to the saved M-value object.
<code>cnPath</code>	Character. Path to the saved copy-number object.
<code>verbose</code>	Logical. If TRUE, emit progress messages with <code>message()</code> .
<code>logs</code>	Logical. If TRUE, write the same messages to a log file.
<code>log_dir</code>	Character or NULL. Directory used for the log file when <code>logs = TRUE</code> .
<code>log_file</code>	Character. File name used when <code>logs = TRUE</code> .

Value

A list with class `"dnaEPICO_preprocessingPheno_metrics"` containing beta, m, and cn.

Examples

```
ex <- dnaEPICO:::examplePreprocessingPhenoStateDnaEpico()
metrics_data <- loadMetricsPreprocessingPheno(
  betaPath = ex$betaPath,
  mPath = ex$mPath,
  cnPath = ex$cnPath,
  verbose = FALSE,
  logs = FALSE
)
names(metrics_data)
```

mergeSvaTargetsEnmix *Merge surrogate variables into the phenotype table*

Description

Merge the surrogate variable matrix back into the phenotype table while preserving the original row order of targets.

Usage

```
mergeSvaTargetsEnmix(  
  targets,  
  sva,  
  SampleID = "Sample_Name",  
  verbose = FALSE,  
  logs = FALSE,  
  log_dir = NULL,  
  log_file = "log_mergeSvaTargetsEnmix.txt"  
)
```

Arguments

targets	Phenotype data frame aligned with the samples in sva.
sva	Numeric matrix of surrogate variables with samples in rows.
SampleID	Character. Name of the phenotype sample identifier column.
verbose	Logical. If TRUE, emit progress messages with message().
logs	Logical. If TRUE, write the same messages to a log file.
log_dir	Character or NULL. Directory used for the log file when logs = TRUE.
log_file	Character. File name used when logs = TRUE.

Value

A phenotype data frame with the surrogate variables appended.

Examples

```
ex <- dnaEPICO:::exampleSvaAnalysisStateDnaEpico()  
merged_pheno <- mergeSvaTargetsEnmix(  
  targets = ex$targets,  
  sva = ex$sva,  
  SampleID = "Sample_Name",  
  verbose = FALSE,  
  logs = FALSE  
)  
colnames(merged_pheno)[seq_len(4)]
```

methylationGLMM_T1T2 *Fit CpG-wise linear mixed-effects models for longitudinal methylation analyses*

Description

methylationGLMM_T1T2() is the high-level coordinator for the longitudinal linear mixed-effects stage of the dnaEPIC0 workflow. It prepares the merged phenotype-plus-beta input, fits one mixed-effects model per CpG for each requested phenotype, extracts phenotype-specific coefficient summaries, optionally collects significant interaction tables, generates diagnostic plots, annotates the combined summary table, and optionally writes legacy-style outputs to disk. The default behavior is now in-memory and quiet, which makes the function easier to compose with other package functions and more aligned with typical Bioconductor usage.

Usage

```
methylationGLMM_T1T2(
  inputPheno = "rData/preprocessingPheno/mergeData/phenoBetaT1T2.RData",
  outputLogs = "logs",
  outputRData = "rData/methylationGLMM_T1T2/models",
  outputPlots = "figures/methylationGLMM_T1T2",
  personVar = "person",
  timeVar = "Timepoint",
  phenotypes = c("DASS_Depression", "DASS_Anxiety", "DASS_Stress", "PCL5_TotalScore",
    "MHCSF_TotalScore", "BRS_TotalScore"),
  covariates = "Sex, Age, Ethnicity, TraumaDefinition, Leukocytes, Epithelial.cells",
  factorVars = "Sex, Ethnicity, TraumaDefinition, Timepoint",
  lmeLibs = "lme4, lmerTest",
  prsMap = NULL,
  libPath = NULL,
  cpgPrefix = "cg",
  cpgLimit = NA,
  nCores = 32,
  summaryPval = NA,
  plotWidth = 2000,
  plotHeight = 1000,
  plotDPI = 150,
  interactionTerm = NULL,
  saveSignificantInteractions = TRUE,
  significantInteractionDir = "preliminaryResults/cpgs/methylationGLMM_T1T2",
  significantInteractionPval = 0.05,
  saveTxtSummaries = TRUE,
  chunkSize = NULL,
  summaryTxtDir = "preliminaryResults/summary/methylationGLMM_T1T2/lmer",
  fdrThreshold = 0.05,
  padjmethod = "fdr",
  annotationPackage = "IlluminaHumanMethylationEPICv2anno.20a1.hg38",
  annotationCols = c("Name", "chr", "pos", "UCSC_RefGene_Group", "UCSC_RefGene_Name",
    "Relation_to_Island", "GencodeV41_Group"),
  annotatedLMEOut = "data/methylationGLMM_T1T2",
  display = FALSE,
```

```

    verbose = FALSE,
    logs = FALSE,
    saveOutputs = FALSE
  )

```

Arguments

inputPheno	Character. Path to the merged longitudinal phenotype-plus-beta .RData or .rds object created by preprocessingPheno(). The default points to the combined timepoint object produced by the package workflow.
outputLogs	Character. Directory used for optional log files.
outputRData	Character. Directory used for optional serialized mixed-model and summary outputs.
outputPlots	Character. Directory used for optional TIFF diagnostic plots.
personVar	Character. Subject identifier variable used for the random intercept. When this column is missing, it is derived from SID using the package's existing sample naming convention.
timeVar	Character. Name of the longitudinal time variable included as a fixed effect in every model.
phenotypes	Character vector or comma-separated phenotype variables to model.
covariates	Character. Comma-separated fixed-effect covariates included in every mixed model.
factorVars	Character. Comma-separated variables that should be coerced to factors before modeling. This usually includes categorical covariates and timeVar.
lmeLibs	Character. Comma-separated package names to validate on worker processes. The default is "lme4, lmerTest".
prsMap	Character or NULL. Optional phenotype-to-PRS mapping in the form "Phenotype1:PRS_1, Phenotype2:PRS_2".
libPath	Character vector or NULL. Optional library paths forwarded to worker processes. By default, the current .libPaths() are used.
cpgPrefix	Character. Prefix used to identify methylation columns in the merged phenotype-plus-beta input object. The default is "cg".
cpgLimit	Integer or NA. Maximum number of CpGs to analyse. Use NA to keep all CpGs matching cpgPrefix.
nCores	Integer. Number of worker processes to use while fitting models and extracting summaries.
summaryPval	Numeric or NA. Optional p-value threshold applied to the returned longitudinal CpG summary tables. Use NA to keep all summary rows.
plotWidth	Integer. TIFF width in pixels when plots are written to disk.
plotHeight	Integer. TIFF height in pixels when plots are written to disk.
plotDPI	Integer. TIFF resolution in DPI when plots are written to disk.
interactionTerm	Character or NULL. Optional interaction term. When supplied and present in the input data, the phenotype is modeled together with its interaction against this variable.
saveSignificantInteractions	Logical. If TRUE, collect coefficient tables for CpGs passing significantInteractionPval in the returned object and optionally write them to disk when saveOutputs = TRUE.

<code>significantInteractionDir</code>	Character. Directory used for optional significant-interaction coefficient tables.
<code>significantInteractionPval</code>	Numeric. P-value threshold used to collect or write significant interaction coefficient tables.
<code>saveTxtSummaries</code>	Logical. If TRUE and <code>saveOutputs = TRUE</code> , write tab-delimited summary tables to <code>summaryTxtDir</code> .
<code>chunkSize</code>	Integer or NULL. Number of CpGs processed per summary extraction chunk. NULL chooses a value automatically.
<code>summaryTxtDir</code>	Character. Directory used for optional tab-delimited LME summary tables.
<code>fdrThreshold</code>	Numeric. False-discovery-rate threshold used to highlight CpGs in the residual-significance diagnostic plots.
<code>padjmethod</code>	Character. P-value adjustment method passed to <code>stats::p.adjust()</code> . The default is "fdr".
<code>annotationPackage</code>	Character. Annotation package or object name passed to <code>minfi::getAnnotation()</code> , for example "IlluminaHumanMethylationEPICv2anno.20a1.hg38".
<code>annotationCols</code>	Character vector or comma-separated annotation columns to append to the combined LME summary table. Available columns depend on the selected annotation package.
<code>annotatedLMEOut</code>	Character. Directory used for the optional annotated LME summary CSV file.
<code>display</code>	Logical. If TRUE, draw diagnostic plots on the active graphics device.
<code>verbose</code>	Logical. If TRUE, emit progress messages with <code>message()</code> . The default is FALSE, so the function is quiet unless requested.
<code>logs</code>	Logical. If TRUE, write the same progress messages to <code>file.path(outputLogs, "log_methylationGLMM_T1T2.txt")</code> .
<code>saveOutputs</code>	Logical. If TRUE, write optional serialized model files, summary tables, significant-interaction tables, annotated results, and TIFF plots to the requested output directories. The default is FALSE, so the function returns in-memory results without writing files.

Value

A list with class "dnaEPICO_methylationGLMM_T1T2".

preparedData Object returned by `prepareMethylationGLMM_T1T2Data()` containing the merged longitudinal phenotype-plus-beta analysis table and modeling metadata.

modelFits Object returned by `fitMethylationGLMM_T1T2Models()` containing the per-phenotype CpG mixed-effects model fits.

modelSummaries Object returned by `summarizeMethylationGLMM_T1T2Models()` containing the combined CpG summary tables used for reporting and annotation.

significantInteractions Object returned by `collectSignificantInteractionsMethylationGLMM_T1T2()` containing optional phenotype-specific significant-interaction tables.

diagnosticPlots Object returned by `plotMethylationGLMM_T1T2Diagnostics()` describing the diagnostic plot objects and any written TIFF files.

annotation Object returned by `annotateMethylationGLMM_T1T2Summaries()` containing the annotated combined summary table.

savedFiles Object returned by `writeMethylationGLMM_T1T2Outputs()` when `saveOutputs = TRUE`, otherwise `NULL`.

See [dnaEPICO_methylationGLMM_T1T2](#) for a class-level overview.

See Also

[dnaEPICO_methylationGLMM_T1T2](#)

Examples

```
if (
  requireNamespace("IlluminaHumanMethylation450kanno.ilmn12.hg19", quietly = TRUE) &&
  requireNamespace("lmerTest", quietly = TRUE)
) {
  tmp <- tempdir()
  toy_path <- file.path(tmp, "phenoBetaT1T2.RData")
  phenoBT1T2 <- data.frame(
    SID = c("P1A", "P1B", "P2A", "P2B", "P3A", "P3B", "P4A", "P4B"),
    person = c(1, 1, 2, 2, 3, 3, 4, 4),
    Timepoint = factor(c("1", "2", "1", "2", "1", "2", "1", "2")),
    score = c(10, 12, 9, 11, 13, 14, 8, 9),
    sex = factor(c("F", "F", "M", "M", "F", "F", "M", "M")),
    cg00000029 = c(0.25, 0.27, 0.20, 0.22, 0.30, 0.31, 0.18, 0.20),
    cg00000108 = c(0.50, 0.53, 0.55, 0.57, 0.48, 0.49, 0.60, 0.61),
    check.names = FALSE
  )
  save(phenoBT1T2, file = toy_path)

  result <- methylationGLMM_T1T2(
    inputPheno = toy_path,
    phenotypes = "score",
    covariates = "sex",
    factorVars = "sex,Timepoint",
    cpGLimit = 2,
    nCores = 1,
    summaryPval = 1,
    annotationPackage = "IlluminaHumanMethylation450kanno.ilmn12.hg19",
    annotationCols = "Name,chr,pos",
    display = FALSE,
    verbose = FALSE,
    logs = FALSE,
    saveOutputs = FALSE
  )

  class(result)
}
```

Description

`methylationGLM_T1()` is the high-level coordinator for the one-timepoint GLM stage of the `dnaEPIC0` workflow. It prepares the merged phenotype-plus-beta input, optionally creates exploratory plots, fits one Gaussian GLM per CpG for each requested phenotype, extracts CpG-level summaries, optionally collects significant CpG coefficient tables, generates diagnostic plots, annotates the combined summary table, and optionally writes legacy-style outputs to disk. The default behavior is now in-memory and quiet, which makes the function easier to compose with other package functions and more aligned with typical Bioconductor usage.

Usage

```
methylationGLM_T1(
  inputPheno = "rData/preprocessingPheno/mergeData/phenoBetaT1.RData",
  outputLogs = "logs",
  outputRData = "rData/methylationGLM_T1/models",
  outputPlots = "figures/methylationGLM_T1",
  phenotypes = c("DASS_Depression", "DASS_Anxiety", "DASS_Stress", "PCL5_TotalScore",
    "MHCSF_TotalScore", "BRS_TotalScore"),
  covariates = "Sex, Age, Ethnicity, TraumaDefinition, Leukocytes, Epithelial.cells",
  factorVars = "Sex, Ethnicity, TraumaDefinition",
  cpGPrefix = "cg",
  cpGLimit = NA,
  nCores = 32,
  plotWidth = 2000,
  plotHeight = 1000,
  plotDPI = 150,
  interactionTerm = NULL,
  libPath = NULL,
  glmLibs = "glm2",
  prsMap = NULL,
  summaryPval = NA,
  summaryResidualSD = TRUE,
  saveSignificantCpGs = FALSE,
  significantCpGDir = "preliminaryResults/cpGs/methylationGLM_T1",
  significantCpGPval = 0.05,
  saveTxtSummaries = TRUE,
  chunkSize = NULL,
  summaryTxtDir = "preliminaryResults/summary/methylationGLM_T1/glm",
  fdrThreshold = 0.05,
  padjmethod = "fdr",
  annotationPackage = "IlluminaHumanMethylationEPICv2anno.20a1.hg38",
  annotationCols = c("Name", "chr", "pos", "UCSC_RefGene_Group", "UCSC_RefGene_Name",
    "Relation_to_Island", "GencodeV41_Group"),
  annotatedGLMOut = "data/methylationGLM_T1",
  display = FALSE,
  verbose = FALSE,
  logs = FALSE,
  saveOutputs = FALSE
)
```

Arguments

inputPheno	Character. Path to the merged phenotype-plus-beta .RData or .rds object created by preprocessingPheno(). The default points to the timepoint-1 object produced by the package workflow.
outputLogs	Character. Directory used for optional log files.
outputRData	Character. Directory used for optional serialized model and summary outputs.
outputPlots	Character. Directory used for optional TIFF plots.
phenotypes	Character vector or comma-separated phenotype variables to model.
covariates	Character. Comma-separated covariate variables included in each GLM.
factorVars	Character. Comma-separated variables that should be treated as factors before modeling.
cpgPrefix	Character. Prefix used to identify methylation columns in the merged phenotype-plus-beta input object. The default is "cg".
cpgLimit	Integer or NA. Maximum number of CpGs to analyse. Use NA to keep all CpGs matching cpgPrefix.
nCores	Integer. Number of worker processes to use while fitting models and extracting summaries.
plotWidth	Integer. TIFF width in pixels when plots are written to disk.
plotHeight	Integer. TIFF height in pixels when plots are written to disk.
plotDPI	Integer. TIFF resolution in DPI when plots are written to disk.
interactionTerm	Character or NULL. Optional interaction term. When supplied and present in the input data, the phenotype is modeled together with its interaction against this variable.
libPath	Character vector or NULL. Optional library paths forwarded to worker processes. By default, the current .libPaths() are used.
glmLibs	Character. Comma-separated package names to validate on worker processes. The default is "glm2".
prsMap	Character or NULL. Optional phenotype-to-PRS mapping in the form "Phenotype1:PRS_1,Phenotype2:PRS_2".
summaryPval	Numeric or NA. Optional p-value threshold applied to the returned CpG summary tables. Use NA to keep all summary rows.
summaryResidualSD	Logical. If TRUE, append residual standard deviations to the CpG summary tables and residual diagnostic plots.
saveSignificantCpGs	Logical. If TRUE, collect significant CpG coefficient tables in the returned object and optionally write them to disk when saveOutputs = TRUE.
significantCpGDir	Character. Directory used for optional significant CpG coefficient tables.
significantCpGPval	Numeric. P-value threshold used to collect or write significant CpG coefficient tables.
saveTxtSummaries	Logical. If TRUE and saveOutputs = TRUE, write tab-delimited summary tables to summaryTxtDir.

chunkSize	Integer or NULL. Number of CpGs processed per summary extraction chunk. NULL chooses a value automatically.
summaryTxtDir	Character. Directory used for optional tab-delimited GLM summary tables.
fdrThreshold	Numeric. False-discovery-rate threshold used to highlight CpGs in the residual-significance diagnostic plots.
padjmethod	Character. P-value adjustment method passed to <code>stats::p.adjust()</code> . The default is "fdr".
annotationPackage	Character. Annotation package or object name passed to <code>minfi::getAnnotation()</code> , for example "IlluminaHumanMethylationEPICv2anno.20a1.hg38".
annotationCols	Character vector or comma-separated annotation columns to append to the combined GLM summary table. Available columns depend on the selected annotation package.
annotatedGLMOut	Character. Directory used for the optional annotated GLM summary CSV file.
display	Logical. If TRUE, draw exploratory and diagnostic plots on the active graphics device.
verbose	Logical. If TRUE, emit progress messages with <code>message()</code> . The default is FALSE, so the function is quiet unless requested.
logs	Logical. If TRUE, write the same progress messages to <code>file.path(outputLogs, "log_methylationGLM_T1.txt")</code> .
saveOutputs	Logical. If TRUE, write optional serialized model files, summary tables, significant-CpG tables, annotated results, and TIFF plots to the requested output directories. The default is FALSE, so the function returns in-memory results without writing files.

Value

A list with class "dnaEPICO_methylationGLM_T1".

preparedData Object returned by `prepareMethylationGLM_T1Data()` containing the merged phenotype-plus-beta analysis table and modeling metadata.

distributionPlots Object returned by `plotMethylationGLM_T1Distributions()` describing any exploratory plots that were generated or written.

modelFits Object returned by `fitMethylationGLM_T1Models()` containing the per-phenotype CpG model fits.

modelSummaries Object returned by `summarizeMethylationGLM_T1Models()` containing the combined CpG summary tables used for reporting and annotation.

significantCpGs Object returned by `collectSignificantCpGsMethylationGLM_T1()` containing optional phenotype-specific significant-CpG tables.

diagnosticPlots Object returned by `plotMethylationGLM_T1Diagnostics()` describing the diagnostic plot objects and any written TIFF files.

annotation Object returned by `annotateMethylationGLM_T1Summaries()` containing the annotated combined summary table.

savedFiles Object returned by `writeMethylationGLM_T1Outputs()` when `saveOutputs = TRUE`, otherwise NULL.

See `dnaEPICO_methylationGLM_T1` for a class-level overview.

See Also[dnaEPICO_methylationGLM_T1](#)**Examples**

```

if (requireNamespace("IlluminaHumanMethylation450kanno.ilmn12.hg19", quietly = TRUE)) {
  tmp <- tempdir()
  toy_path <- file.path(tmp, "phenoBetaT1.RData")
  phenoBT1 <- data.frame(
    Sample_Name = c("S1", "S2", "S3", "S4"),
    status = factor(c("Case", "Case", "Control", "Control")),
    sex = factor(c("F", "M", "F", "M")),
    cg00000029 = c(0.20, 0.25, 0.22, 0.27),
    cg00000108 = c(0.60, 0.55, 0.52, 0.58),
    check.names = FALSE
  )
  save(phenoBT1, file = toy_path)

  result <- methylationGLM_T1(
    inputPheno = toy_path,
    phenotypes = "status",
    covariates = "sex",
    factorVars = "status,sex",
    cpGLimit = 2,
    nCores = 1,
    summaryPval = 1,
    annotationPackage = "IlluminaHumanMethylation450kanno.ilmn12.hg19",
    annotationCols = "Name,chr,pos",
    display = FALSE,
    verbose = FALSE,
    logs = FALSE,
    saveOutputs = FALSE
  )

  class(result)
}

```

normalizeMinfiEwasWater

Normalize filtered samples with minfi and wateRmelon methods

Description

Apply one or more supported normalization methods to a filtered RGSets and return all normalized objects together in a single result object.

Usage

```

normalizeMinfiEwasWater(
  sampleData,
  sexColumn = "Sex",
  normMethods = "adjustedfunnorm",

```

```

    verbose = FALSE,
    logs = FALSE,
    log_dir = NULL,
    log_file = "log_normalizeMinfiEwasWater.txt"
  )

```

Arguments

sampleData	Object returned by filterSamplesMinfiEwasWater().
sexColumn	Character. Name of the phenotype column used as the optional sex covariate for normalization methods that support it.
normMethods	Character vector or semicolon-separated string of normalization methods. Supported values are "adjustedfunnorm", "funnorm", "illumina", "quantile", and "swan".
verbose	Logical. If TRUE, emit progress messages with message().
logs	Logical. If TRUE, write the same messages to a log file.
log_dir	Character or NULL. Directory used for the log file when logs = TRUE.
log_file	Character. File name used when logs = TRUE.

Value

A list with class "dnaEPICO_minfiEwasWater_norm" containing the requested normalized objects and the first method as primary.

Examples

```

ex <- dnaEPICO:::exampleMinfiBaseDataDnaEpico()
sample_data <- filterSamplesMinfiEwasWater(
  RGSet = ex$RGSet,
  targets = ex$targets,
  failedSamples = character(0),
  SampleID = "Sample_Name",
  verbose = FALSE,
  logs = FALSE
)
norm_data <- normalizeMinfiEwasWater(
  sampleData = sample_data,
  sexColumn = "Sex",
  normMethods = "quantile",
  verbose = FALSE,
  logs = FALSE
)
names(norm_data$normalized)

```

plotAssessmentMinfiEwasWater

Plot quality-assessment outputs for preprocessingMinfiEwasWater

Description

Draw either the minfi QC plot or the detection P-value plot from an assessment object returned by `assessSamplesMinfiEwasWater()`.

Usage

```
plotAssessmentMinfiEwasWater(
  assessment,
  plot = c("qc", "detection"),
  display = FALSE,
  file = NULL,
  width = 2000L,
  height = 1000L,
  res = 150L,
  verbose = FALSE,
  logs = FALSE,
  log_dir = NULL,
  log_file = "log_plotAssessmentMinfiEwasWater.txt"
)
```

Arguments

<code>assessment</code>	Object returned by <code>assessSamplesMinfiEwasWater()</code> .
<code>plot</code>	Character. Plot type: "qc" or "detection".
<code>display</code>	Logical. If TRUE, draw the plot on the active graphics device.
<code>file</code>	Character or NULL. TIFF file written when supplied.
<code>width</code>	Integer. TIFF width in pixels when file is supplied.
<code>height</code>	Integer. TIFF height in pixels when file is supplied.
<code>res</code>	Integer. TIFF resolution in DPI when file is supplied.
<code>verbose</code>	Logical. If TRUE, emit progress messages with <code>message()</code> .
<code>logs</code>	Logical. If TRUE, write the same messages to a log file.
<code>log_dir</code>	Character or NULL. Directory used for the log file when <code>logs = TRUE</code> .
<code>log_file</code>	Character. File name used when <code>logs = TRUE</code> .

Value

Invisibly returns the saved TIFF path when `file` is supplied, otherwise NULL.

Examples

```
assessment <- list(
  meanDetP = c(S1 = 0.01, S2 = 0.02, S3 = 0.04),
  detPThreshold = 0.05
)
plotAssessmentMinfiEwasWater(
  assessment = assessment,
  plot = "detection",
  display = FALSE,
  verbose = FALSE,
  logs = FALSE
)
```

 plotCtrlMinfiEwasWater

Plot ENmix control images from an RGSet

Description

Call `ENmix::plotCtrl()` for a supplied `RGSet`. This function only writes files when `output_dir` is provided because `ENmix::plotCtrl()` produces JPG files on disk rather than returning a plot object.

Usage

```
plotCtrlMinfiEwasWater(
  RGSet,
  output_dir = NULL,
  verbose = FALSE,
  logs = FALSE,
  log_dir = NULL,
  log_file = "log_plotCtrlMinfiEwasWater.txt"
)
```

Arguments

<code>RGSet</code>	An <code>RGChannelSet</code> .
<code>output_dir</code>	Character or <code>NULL</code> . Directory where ENmix control JPG files should be written. If <code>NULL</code> , the function returns without writing files.
<code>verbose</code>	Logical. If <code>TRUE</code> , emit progress messages with <code>message()</code> .
<code>logs</code>	Logical. If <code>TRUE</code> , write the same messages to a log file.
<code>log_dir</code>	Character or <code>NULL</code> . Directory used for the log file when <code>logs = TRUE</code> .
<code>log_file</code>	Character. File name used when <code>logs = TRUE</code> .

Value

Invisibly returns `output_dir`.

Examples

```
ex <- dnaEPICO::exampleMinfiBaseDataDnaEpico()
output_dir <- file.path(tempdir(), "enmix-control-plots")
plotCtrlMinfiEwasWater(
  RGSet = ex$RGSet,
  output_dir = output_dir,
  verbose = FALSE,
  logs = FALSE
)
dir.exists(output_dir)
```

plotMethylationGLMM_T1T2Diagnostics

Plot longitudinal mixed-effects model diagnostics

Description

Create Q-Q and standard-error diagnostic plots from the mixed-effects summary tables returned by `summarizeMethylationGLMM_T1T2Models()`.

Usage

```
plotMethylationGLMM_T1T2Diagnostics(
  modelSummaries,
  preparedData,
  fdrThreshold = 0.05,
  padjmethod = "fdr",
  outputDir = NULL,
  plotWidth = 2000L,
  plotHeight = 1000L,
  plotDPI = 150L,
  display = FALSE,
  verbose = FALSE,
  logs = FALSE,
  log_dir = NULL,
  log_file = "log_methylationGLMM_T1T2.txt"
)
```

Arguments

<code>modelSummaries</code>	Object returned by <code>summarizeMethylationGLMM_T1T2Models()</code> .
<code>preparedData</code>	Object returned by <code>prepareMethylationGLMM_T1T2Data()</code> .
<code>fdrThreshold</code>	Numeric. False-discovery-rate threshold used to highlight CpGs in the diagnostic plots.
<code>padjmethod</code>	Character. P-value adjustment method passed to <code>stats::p.adjust()</code> .
<code>outputDir</code>	Character or NULL. Directory used for TIFF files. When NULL, plots are returned in memory only.
<code>plotWidth</code>	Integer. TIFF width in pixels when plots are written to disk.
<code>plotHeight</code>	Integer. TIFF height in pixels when plots are written to disk.
<code>plotDPI</code>	Integer. TIFF resolution in DPI when plots are written to disk.
<code>display</code>	Logical. If TRUE, draw plots on the active graphics device.
<code>verbose</code>	Logical. If TRUE, emit progress messages with <code>message()</code> .
<code>logs</code>	Logical. If TRUE, write the same messages to a log file.
<code>log_dir</code>	Character or NULL. Directory used for the log file when <code>logs = TRUE</code> .
<code>log_file</code>	Character. File name used when <code>logs = TRUE</code> .

Value

A list with class `"dnaEPIC0_methylationGLMM_T1T2_diagnostic_plots"` containing the generated `ggplot2` objects, genomic inflation factors, and any saved TIFF file paths.

Examples

```
ex <- dnaEPICO:::exampleMethylationGLMMStateDnaEpico()
diagnostic_plots <- plotMethylationGLMM_T1T2Diagnostics(
  modelSummaries = ex$modelSummaries,
  preparedData = ex$preparedData,
  display = FALSE,
  verbose = FALSE,
  logs = FALSE
)
names(diagnostic_plots$plots)
```

```
plotMethylationGLM_T1Diagnostics
```

Plot diagnostic summaries for one-timepoint methylation GLMs

Description

Create Q-Q and residual-diagnostic plots from the CpG summary tables returned by `summarizeMethylationGLM_T1Models()`.

Usage

```
plotMethylationGLM_T1Diagnostics(
  modelSummaries,
  preparedData,
  fdrThreshold = 0.05,
  padjmethod = "fdr",
  outputDir = NULL,
  plotWidth = 2000L,
  plotHeight = 1000L,
  plotDPI = 150L,
  display = FALSE,
  verbose = FALSE,
  logs = FALSE,
  log_dir = NULL,
  log_file = "log_methylationGLM_T1.txt"
)
```

Arguments

<code>modelSummaries</code>	Object returned by <code>summarizeMethylationGLM_T1Models()</code> .
<code>preparedData</code>	Object returned by <code>prepareMethylationGLM_T1Data()</code> .
<code>fdrThreshold</code>	Numeric. False-discovery-rate threshold used to highlight CpGs in the diagnostic plots.
<code>padjmethod</code>	Character. P-value adjustment method passed to <code>stats::p.adjust()</code> .
<code>outputDir</code>	Character or NULL. Directory used for TIFF files. When NULL, plots are returned in memory only.
<code>plotWidth</code>	Integer. TIFF width in pixels when plots are written to disk.
<code>plotHeight</code>	Integer. TIFF height in pixels when plots are written to disk.

plotDPI	Integer. TIFF resolution in DPI when plots are written to disk.
display	Logical. If TRUE, draw plots on the active graphics device.
verbose	Logical. If TRUE, emit progress messages with message().
logs	Logical. If TRUE, write the same messages to a log file.
log_dir	Character or NULL. Directory used for the log file when logs = TRUE.
log_file	Character. File name used when logs = TRUE.

Value

A list with class "dnaEPICO_methylationGLM_T1_diagnostic_plots" containing the generated ggplot2 objects, genomic inflation factors, and any saved TIFF file paths.

Examples

```
ex <- dnaEPICO::exampleMethylationGLMStateDnaEpico()
diagnostic_plots <- plotMethylationGLM_T1Diagnostics(
  modelSummaries = ex$modelSummaries,
  preparedData = ex$preparedData,
  display = FALSE,
  verbose = FALSE,
  logs = FALSE
)
names(diagnostic_plots$plots)
```

plotMethylationGLM_T1Distributions

Plot phenotype and covariate distributions for one-timepoint GLM analyses

Description

Create phenotype, factor-variable, and numeric-covariate distribution plots from the object returned by prepareMethylationGLM_T1Data().

Usage

```
plotMethylationGLM_T1Distributions(
  preparedData,
  plotWidth = 2000L,
  plotHeight = 1000L,
  plotDPI = 150L,
  outputDir = NULL,
  display = FALSE,
  verbose = FALSE,
  logs = FALSE,
  log_dir = NULL,
  log_file = "log_methylationGLM_T1.txt"
)
```

Arguments

preparedData	Object returned by prepareMethylationGLM_T1Data().
plotWidth	Integer. TIFF width in pixels when plots are written to disk.
plotHeight	Integer. TIFF height in pixels when plots are written to disk.
plotDPI	Integer. TIFF resolution in DPI when plots are written to disk.
outputDir	Character or NULL. Directory used for TIFF files. When NULL, plots are returned in memory only.
display	Logical. If TRUE, draw plots on the active graphics device.
verbose	Logical. If TRUE, emit progress messages with message().
logs	Logical. If TRUE, write the same messages to a log file.
log_dir	Character or NULL. Directory used for the log file when logs = TRUE.
log_file	Character. File name used when logs = TRUE.

Value

A list with class "dnaEPICO_methylationGLM_T1_distribution_plots" containing the generated ggplot2 objects and any saved TIFF file paths.

Examples

```
ex <- dnaEPICO::exampleMethylationGLMStateDnaEpico()
distribution_plots <- plotMethylationGLM_T1Distributions(
  preparedData = ex$preparedData,
  display = FALSE,
  verbose = FALSE,
  logs = FALSE
)
names(distribution_plots)
```

plotMetricsMinfiEwasWater

Plot multidimensional scaling or density summaries from final metrics

Description

Plot multidimensional scaling or density summaries from final metrics

Usage

```
plotMetricsMinfiEwasWater(
  metricsData,
  targets,
  plot = c("mds", "density"),
  plotGroupVar = "Sex",
  sexColumn = "Sex",
  display = FALSE,
  file = NULL,
  width = 2000L,
```

```

    height = 1000L,
    res = 150L,
    verbose = FALSE,
    logs = FALSE,
    log_dir = NULL,
    log_file = "log_plotMetricsMinfiEwasWater.txt"
  )

```

Arguments

metricsData	Object returned by extractMetricsMinfiEwasWater().
targets	Filtered phenotype data aligned with metricsData.
plot	Character. Plot type: "mds" or "density".
plotGroupVar	Character. Phenotype column used for the main grouping.
sexColumn	Character. Phenotype column used for the sex grouping in the MDS plot.
display	Logical. If TRUE, draw the plot on the active graphics device.
file	Character or NULL. TIFF file written when supplied.
width	Integer. TIFF width in pixels when file is supplied.
height	Integer. TIFF height in pixels when file is supplied.
res	Integer. TIFF resolution in DPI when file is supplied.
verbose	Logical. If TRUE, emit progress messages with message().
logs	Logical. If TRUE, write the same messages to a log file.
log_dir	Character or NULL. Directory used for the log file when logs = TRUE.
log_file	Character. File name used when logs = TRUE.

Value

Invisibly returns the saved TIFF path when file is supplied, otherwise NULL.

Examples

```

ex <- dnaEPICO:::exampleMinfiMetricsStateDnaEpico()
plotMetricsMinfiEwasWater(
  metricsData = ex$metricsData,
  targets = ex$targets,
  plot = "density",
  plotGroupVar = "Sex",
  sexColumn = "Sex",
  display = FALSE,
  verbose = FALSE,
  logs = FALSE
)

```

```
plotNormalizationMinfiEwasWater
```

Plot raw and normalized methylation distributions

Description

Draw the density comparison plot used to inspect raw versus normalized data.

Usage

```
plotNormalizationMinfiEwasWater(
  RGSet,
  normData,
  targets,
  sexColumn = "Sex",
  display = FALSE,
  file = NULL,
  width = 2000L,
  height = 1000L,
  res = 150L,
  verbose = FALSE,
  logs = FALSE,
  log_dir = NULL,
  log_file = "log_plotNormalizationMinfiEwasWater.txt"
)
```

Arguments

RGSet	An RGChannelSet aligned with targets.
normData	Object returned by <code>normalizeMinfiEwasWater()</code> .
targets	Filtered phenotype data aligned with RGSet.
sexColumn	Character. Name of the phenotype column used to colour the density curves.
display	Logical. If TRUE, draw the plot on the active graphics device.
file	Character or NULL. TIFF file written when supplied.
width	Integer. TIFF width in pixels when file is supplied.
height	Integer. TIFF height in pixels when file is supplied.
res	Integer. TIFF resolution in DPI when file is supplied.
verbose	Logical. If TRUE, emit progress messages with <code>message()</code> .
logs	Logical. If TRUE, write the same messages to a log file.
log_dir	Character or NULL. Directory used for the log file when <code>logs = TRUE</code> .
log_file	Character. File name used when <code>logs = TRUE</code> .

Value

Invisibly returns the saved TIFF path when file is supplied, otherwise NULL.

Examples

```

ex <- dnaEPICO:::exampleMinfiMetricsStateDnaEpico()
plotNormalizationMinfiEwasWater(
  RGSet = ex$beta,
  normData = ex$normData,
  targets = ex$targets,
  sexColumn = "Sex",
  display = FALSE,
  verbose = FALSE,
  logs = FALSE
)

```

plotRawDensityMinfiEwasWater

Plot raw beta-value density from a raw preprocessing object

Description

Draw the pre-normalization beta density plot from a raw minfi object and a grouping variable in the phenotype table.

Usage

```

plotRawDensityMinfiEwasWater(
  rawData,
  targets,
  plotGroupVar = "Sex",
  display = FALSE,
  file = NULL,
  width = 2000L,
  height = 1000L,
  res = 150L,
  verbose = FALSE,
  logs = FALSE,
  log_dir = NULL,
  log_file = "log_plotRawDensityMinfiEwasWater.txt"
)

```

Arguments

rawData	Object returned by buildRawMinfiEwasWater().
targets	Filtered phenotype data aligned with rawData.
plotGroupVar	Character. Phenotype column used to group samples in the density plot.
display	Logical. If TRUE, draw the plot on the active graphics device.
file	Character or NULL. TIFF file written when supplied.
width	Integer. TIFF width in pixels when file is supplied.
height	Integer. TIFF height in pixels when file is supplied.
res	Integer. TIFF resolution in DPI when file is supplied.

verbose	Logical. If TRUE, emit progress messages with message().
logs	Logical. If TRUE, write the same messages to a log file.
log_dir	Character or NULL. Directory used for the log file when logs = TRUE.
log_file	Character. File name used when logs = TRUE.

Value

Invisibly returns the saved TIFF path when file is supplied, otherwise NULL.

Examples

```
ex <- dnaEPICO::exampleMinfiMetricsStateDnaEpico()
plotRawDensityMinfiEwasWater(
  rawData = ex$rawData,
  targets = ex$targets,
  plotGroupVar = "Sex",
  display = FALSE,
  verbose = FALSE,
  logs = FALSE
)
```

plotSexMinfiEwasWater *Plot predicted or clinical sex from predictSexMinfiEwasWater()*

Description

Plot predicted or clinical sex from predictSexMinfiEwasWater()

Usage

```
plotSexMinfiEwasWater(
  sexData,
  type = c("predicted", "clinical"),
  display = FALSE,
  file = NULL,
  width = 2000L,
  height = 1000L,
  res = 70L,
  verbose = FALSE,
  logs = FALSE,
  log_dir = NULL,
  log_file = "log_plotSexMinfiEwasWater.txt"
)
```

Arguments

sexData	Object returned by predictSexMinfiEwasWater().
type	Character. Plot type: "predicted" for methylation-predicted sex or "clinical" for reported sex.
display	Logical. If TRUE, draw the plot on the active graphics device.

file	Character or NULL. TIFF file written when supplied.
width	Integer. TIFF width in pixels when file is supplied.
height	Integer. TIFF height in pixels when file is supplied.
res	Integer. TIFF resolution in DPI when file is supplied.
verbose	Logical. If TRUE, emit progress messages with message().
logs	Logical. If TRUE, write the same messages to a log file.
log_dir	Character or NULL. Directory used for the log file when logs = TRUE.
log_file	Character. File name used when logs = TRUE.

Value

Invisibly returns the saved TIFF path when file is supplied, otherwise NULL.

Examples

```
ex <- dnaEPICO:::exampleSexPlotStateDnaEpico()
plotSexMinfiEwasWater(
  sexData = ex,
  type = "predicted",
  display = FALSE,
  verbose = FALSE,
  logs = FALSE
)
```

plotSvaEnmix

Plot surrogate variables for svaEnmix

Description

Draw one of the standard surrogate-variable plots used by svaEnmix().

Usage

```
plotSvaEnmix(
  analysisData,
  plot = c("sentrix_id", "sentrix_position", "matrix"),
  display = FALSE,
  file = NULL,
  width = 2000L,
  height = 1000L,
  res = 150L,
  verbose = FALSE,
  logs = FALSE,
  log_dir = NULL,
  log_file = "log_plotSvaEnmix.txt"
)
```

Arguments

analysisData	Object returned by analyzeSvaEnmix().
plot	Character. Plot type: "sentrrix_id", "sentrrix_position", or "matrix".
display	Logical. If TRUE, draw the plot on the active graphics device.
file	Character or NULL. TIFF file path used for saved output.
width	Integer. Plot width in pixels when file is supplied.
height	Integer. Plot height in pixels when file is supplied.
res	Integer. TIFF resolution in DPI when file is supplied.
verbose	Logical. If TRUE, emit progress messages with message().
logs	Logical. If TRUE, write the same messages to a log file.
log_dir	Character or NULL. Directory used for the log file when logs = TRUE.
log_file	Character. File name used when logs = TRUE.

Value

Invisibly returns file when a TIFF is written, otherwise NULL.

Examples

```
ex <- dnaEPICO:::exampleSvaAnalysisStateDnaEpicco()
plotSvaEnmix(
  analysisData = ex$analysisData,
  plot = "sentrrix_id",
  display = FALSE,
  verbose = FALSE,
  logs = FALSE
)
```

predictSexMinfiEwasWater

Predict biological sex from a filtered raw-data object

Description

Predict sample sex from a genome-mapped methylation object, align the predictions with phenotype data, and return a structured object that can be plotted or merged into downstream phenotype tables.

Usage

```
predictSexMinfiEwasWater(
  rawData,
  targets,
  SampleID = "Sample_Name",
  sexColumn = "Sex",
  verbose = FALSE,
  logs = FALSE,
  log_dir = NULL,
  log_file = "log_predictSexMinfiEwasWater.txt"
)
```

Arguments

rawData	Object returned by buildRawMinfiEwasWater().
targets	Filtered phenotype data frame aligned with rawData.
SampleID	Character. Name of the sample identifier column in targets.
sexColumn	Character. Name of the phenotype column containing reported sex.
verbose	Logical. If TRUE, emit progress messages with message().
logs	Logical. If TRUE, write the same messages to a log file.
log_dir	Character or NULL. Directory used for the log file when logs = TRUE.
log_file	Character. File name used when logs = TRUE.

Value

A list with class "dnaEPICO_minfiEwasWater_sex" containing the sex prediction result, aligned phenotype data, plotting data, and mismatch table.

Examples

```
ex <- dnaEPICO::exampleMinfiWorkflowStateDnaEpico()
sex_data <- predictSexMinfiEwasWater(
  rawData = ex$rawFiltered,
  targets = ex$sampleData$targets,
  SampleID = "Sample_Name",
  sexColumn = "Sex",
  verbose = FALSE,
  logs = FALSE
)
names(sex_data)
```

```
prepareDnamReportInputs
```

Prepare inputs for a DNA methylation report

Description

Prepare inputs for a DNA methylation report

Usage

```
prepareDnamReportInputs(
  outputDir = "reports",
  qcDir = file.path("figures", "preprocessingMinfiEwasWater", "enmix"),
  preprocessingDir = file.path("figures", "preprocessingMinfiEwasWater", "qc"),
  postprocessingDir = file.path("figures", "preprocessingMinfiEwasWater", "metrics"),
  svaDir = file.path("figures", "svaEnmix"),
  glmDir = file.path("figures", "methylationGLM_T1"),
  glmmDir = file.path("figures", "methylationGLMM_T1T2"),
  figDir = file.path(outputDir, "assets", "figures"),
  verbose = FALSE,
  logs = FALSE,
  logDir = outputDir
)
```

Arguments

outputDir	Character. Directory where the report project is written.
qcDir	Character. Directory containing ENmix quality-control figures.
preprocessingDir	Character. Directory containing preprocessing quality-control figures.
postprocessingDir	Character. Directory containing postprocessing metric figures.
svaDir	Character. Directory containing SVA or batch-effect figures.
glmDir	Character. Directory containing GLM figures.
glmmDir	Character. Directory containing GLMM figures.
figDir	Character. Directory used for generated report figure assets.
verbose	Logical. If TRUE, emit progress messages with message().
logs	Logical. If TRUE, write progress messages to file.path(logDir, "log_dnamReport.txt").
logDir	Character. Directory for optional log files.

Value

A list with class "dnaEPICO_dnamReport_prepared".

Examples

```
report_root <- file.path(tempdir(), "dnaepico-report-inputs")
prepared <- prepareDnamReportInputs(
  outputDir = file.path(report_root, "reports"),
  qcDir = file.path(
    report_root,
    "figures",
    "preprocessingMinfiEwasWater",
    "enmix"
  ),
  preprocessingDir = file.path(
    report_root,
    "figures",
    "preprocessingMinfiEwasWater",
    "qc"
  ),
  postprocessingDir = file.path(
    report_root,
    "figures",
    "preprocessingMinfiEwasWater",
    "metrics"
  ),
  svaDir = file.path(report_root, "figures", "svaEnmix")
)
inherits(prepared, "dnaEPICO_dnamReport_prepared")
```

```
prepareMethylationGLMM_T1T2Data
```

Prepare longitudinal phenotype-plus-beta data for mixed-effects analyses

Description

Load the merged longitudinal phenotype-plus-beta object, ensure that a subject identifier column is available, validate the requested modeling variables, convert selected variables to factors, and return a single in-memory object for downstream mixed-effects modeling helpers.

Usage

```
prepareMethylationGLMM_T1T2Data(
  inputPheno,
  personVar = "person",
  timeVar = "Timepoint",
  phenotypes,
  covariates,
  factorVars,
  prsMap = NULL,
  cpGPrefix = "cg",
  cpGLimit = NA,
  interactionTerm = NULL,
  verbose = FALSE,
  logs = FALSE,
  log_dir = NULL,
  log_file = "log_methylationGLMM_T1T2.txt"
)
```

Arguments

inputPheno	Character. Path to the merged longitudinal phenotype-plus- beta object created by preprocessingPheno().
personVar	Character. Name of the subject identifier column.
timeVar	Character. Name of the time variable.
phenotypes	Character vector or comma-separated string of phenotype variables to model.
covariates	Character vector or comma-separated string of covariate variables to adjust for.
factorVars	Character vector or comma-separated string of variables that should be converted to factors before modeling.
prsMap	Character vector or comma-separated string of phenotype-to-PRS mappings in the form "Phenotype:PRS".
cpGPrefix	Character. Prefix used to identify methylation columns.
cpGLimit	Integer or NA. Maximum number of CpGs to retain. NA keeps all matching CpGs.
interactionTerm	Character or NULL. Optional interaction term.
verbose	Logical. If TRUE, emit progress messages with message().
logs	Logical. If TRUE, write the same messages to a log file.

log_dir Character or NULL. Directory used for the log file when logs = TRUE.
 log_file Character. File name used when logs = TRUE.

Value

A list with class "dnaEPICO_methylationGLMM_T1T2_data" containing the prepared analysis data, parsed variable selections, CpG columns, timepoint summaries, and subject-ID diagnostics.

Examples

```
ex <- dnaEPICO:::exampleMethylationGLMMStateDnaEpico()
prepared_data <- prepareMethylationGLMM_T1T2Data(
  inputPheno = ex$inputPath,
  personVar = "person",
  timeVar = "Timepoint",
  phenotypes = "score",
  covariates = "sex",
  factorVars = "sex,Timepoint",
  cpGLimit = 2,
  verbose = FALSE,
  logs = FALSE
)
names(prepared_data)
```

prepareMethylationGLM_T1Data

Prepare phenotype-plus-beta data for one-timepoint GLM analyses

Description

Load the merged phenotype-plus-beta input object, validate the requested modeling variables, convert selected variables to factors, and return a single in-memory object for downstream helpers.

Usage

```
prepareMethylationGLM_T1Data(
  inputPheno,
  phenotypes,
  covariates,
  factorVars,
  cpGLimit = NA,
  interactionTerm = NULL,
  prsMap = NULL,
  verbose = FALSE,
  logs = FALSE,
  log_dir = NULL,
  log_file = "log_methylationGLM_T1.txt"
)
```

Arguments

inputPheno	Character. Path to the merged phenotype-plus-beta object created by preprocessingPheno().
phenotypes	Character vector or comma-separated string of phenotype variables to model.
covariates	Character vector or comma-separated string of covariate variables to adjust for.
factorVars	Character vector or comma-separated string of variables that should be converted to factors before modeling.
cpgPrefix	Character. Prefix used to identify methylation columns.
cpgLimit	Integer or NA. Maximum number of CpGs to retain. NA keeps all matching CpGs.
interactionTerm	Character or NULL. Optional interaction term.
prsMap	Character vector or comma-separated string of phenotype-to-PRS mappings in the form "Phenotype:PRS".
verbose	Logical. If TRUE, emit progress messages with message().
logs	Logical. If TRUE, write the same messages to a log file.
log_dir	Character or NULL. Directory used for the log file when logs = TRUE.
log_file	Character. File name used when logs = TRUE.

Value

A list with class "dnaEPICO_methylationGLM_T1_data" containing the prepared analysis data, parsed variable selections, CpG columns, and exploratory summaries.

Examples

```
ex <- dnaEPICO::exampleMethylationGLMStateDnaEpico()
prepared_data <- prepareMethylationGLM_T1Data(
  inputPheno = ex$inputPath,
  phenotypes = "status",
  covariates = "sex,age",
  factorVars = "status,sex",
  cpgLimit = 2,
  verbose = FALSE,
  logs = FALSE
)
names(prepared_data)
```

```
preprocessingMinfiEwasWater
```

Convenience preprocessing pipeline for Illumina methylation arrays

Description

Run the dnaEPICO preprocessing workflow as a convenience wrapper around the smaller minfi/ENmix/wateRmelon helper functions in this package. The wrapper now returns a structured result object containing the in-memory outputs from each stage. Legacy files are written only when saveOutputs = TRUE.

Usage

```
preprocessingMinfiEwasWater(
  phenoFile = "data/preprocessingMinfiEwasWater/pheno.csv",
  idatFolder = "data/preprocessingMinfiEwasWater/idats",
  outputLogs = "logs",
  nSamples = NA,
  SampleID = "Sample_Name",
  arrayType = "IlluminaHumanMethylationEPICv2",
  annotationVersion = "20a1.hg38",
  scriptLabel = "preprocessingMinfiEwasWater",
  baseDataFolder = "rData",
  figureBaseDir = "figures",
  sepType = "",
  tiffWidth = 2000,
  tiffHeight = 1000,
  tiffRes = 150,
  qcCutoff = 10.5,
  detPtype = "m+u",
  detPThreshold = 0.05,
  normMethods = "adjustedfunnorm",
  sexColumn = "Sex",
  pvalThreshold = 0.01,
  chrToRemove = "chrX,chrY",
  snpsToRemove = "SBE,CpG",
  mafThreshold = 0.1,
  crossReactivePath = "data/preprocessingMinfiEwasWater/12864_2024_10027_MOESM8_ESM.csv",
  plotGroupVar = "Sex",
  lcRef = "salivaEPIC",
  phenoOrder = "Sample_Name;Timepoint;Sex;PredSex;Basename;Sentrix_ID;Sentrix_Position",
  lcPhenoDir = "data/preprocessingMinfiEwasWater",
  display = FALSE,
  verbose = FALSE,
  logs = FALSE,
  saveOutputs = FALSE
)
```

Arguments

<code>phenoFile</code>	Character. Path to the phenotype CSV file.
<code>idatFolder</code>	Character. Directory containing the IDAT files.
<code>outputLogs</code>	Character. Directory used for log files when <code>logs = TRUE</code> .
<code>nSamples</code>	Integer or NA. Number of rows to keep from the phenotype table. Use NA to keep all samples.
<code>SampleID</code>	Character. Name of the phenotype column containing sample identifiers.
<code>arrayType</code>	Character. Illumina array identifier passed to <code>Biobase::annotation()</code> , for example "IlluminaHumanMethylationEPICv2".
<code>annotationVersion</code>	Character. Annotation build passed to <code>Biobase::annotation()</code> , for example "20a1.hg38" or "ilmn12.hg19".
<code>scriptLabel</code>	Character. Label used to name output folders when <code>saveOutputs = TRUE</code> .

baseDataFolder	Character. Base directory used for saved .RData outputs when saveOutputs = TRUE.
figureBaseDir	Character. Base directory used for saved figure outputs when saveOutputs = TRUE.
sepType	Character. Field separator used in phenoFile. Use "," for a comma-separated file, "\\t" for a tab-delimited file, or another separator accepted by <code>utils::read.csv()</code> .
tiffWidth	Integer. Width of saved TIFF plots in pixels.
tiffHeight	Integer. Height of saved TIFF plots in pixels.
tiffRes	Integer. Resolution in DPI for saved TIFF plots.
qcCutoff	Numeric. QC cutoff passed to <code>minfi::plotQC()</code> .
detPtype	Character. Detection P-value mode passed to <code>minfi::detectionP()</code> . Common values in minfi workflows are "m+u" and "negative". The default here is "m+u".
detPThreshold	Numeric. Samples with mean detection P value above this threshold are removed.
normMethods	Character vector or semicolon-separated string of normalization methods. Supported values are "adjustedfunnorm", "funnorm", "illumina", "quantile", and "swan".
sexColumn	Character. Name of the phenotype column containing reported sex.
pvalThreshold	Numeric. Probe-level detection P-value threshold used in the probe filter.
chrToRemove	Character vector or comma-separated string of chromosome names to remove, for example "chrX, chrY".
snpsToRemove	Character vector or comma-separated string of SNP probe types to remove, for example "SBE, CpG".
mafThreshold	Numeric. Minor allele frequency threshold passed to <code>minfi::dropLociWithSnps()</code> .
crossReactivePath	Character. Path to a CSV file containing a ProbeID column of cross-reactive probes to remove.
plotGroupVar	Character. Phenotype column used for density and MDS grouping plots.
lcRef	Character. Reference panel used for cell composition estimation. "saliva" and "salivaEPIC" use <code>estimateLC()</code> . Other values are passed to <code>ENmix::estimateCellProp()</code> .
phenoOrder	Character vector or semicolon-separated string describing which phenotype columns should appear first in the merged phenoLC table.
lcPhenoDir	Character. Directory used for the saved phenoLC.csv file when saveOutputs = TRUE.
display	Logical. If TRUE, draw plots on the active graphics device.
verbose	Logical. If TRUE, emit progress messages with <code>message()</code> . The default is FALSE.
logs	Logical. If TRUE, write log messages to outputLogs. The default is FALSE.
saveOutputs	Logical. If TRUE, write the legacy .RData, figure, and phenoLC.csv outputs to disk. The default is FALSE, so the function can be used in the more traditional in-memory Bioconductor style.

Value

A list with class "dnaEPICO_preprocessingMinfiEwasWater".

targets Filtered phenotype table aligned to the retained samples.

RGSet Filtered RGChannelSet used in downstream preprocessing and available for direct interactive inspection.

rawData Object returned by `buildRawMinfiEwasWater()` containing the raw MSet, RatioSet, and genome-mapped object derived from RGSet.

assessment Object returned by `assessSamplesMinfiEwasWater()` containing detection P values, QC summaries, and failed-sample tracking.

sexData Object returned by `predictSexMinfiEwasWater()` containing predicted sex labels, mismatch summaries, and plotting data.

normData Object returned by `normalizeMinfiEwasWater()` containing the requested normalized objects and metadata on the methods that were run.

filterData Object returned by `filterProbesMinfiEwasWater()` containing the probe-filtered methylation objects at each filtering stage.

metricsData Object returned by `extractMetricsMinfiEwasWater()` containing the beta-value, M-value, and copy-number matrices used by later workflow steps.

lcData Object returned by `estimateLCMinfiEwasWater()` containing the estimated cell-type proportions and the phenotype table augmented with those proportions.

logFile Resolved path to the optional log file, or NULL when logging was disabled.

See [dnaEPICO_preprocessingMinfiEwasWater](#) for a class-level overview.

See Also

[dnaEPICO_preprocessingMinfiEwasWater](#)

Examples

```
if (requireNamespace("minfiData", quietly = TRUE) &&
    requireNamespace("IlluminaHumanMethylation450kmanifest", quietly = TRUE) &&
    requireNamespace("IlluminaHumanMethylation450kanno.ilmn12.hg19", quietly = TRUE)) {
  ex <- dnaEPICO::exampleMinfiIdatInputsDnaEpico(n = 4)
  result <- preprocessingMinfiEwasWater(
    phenoFile = ex$phenoFile,
    idatFolder = ex$idatFolder,
    outputLogs = file.path(ex$tempDir, "logs"),
    nSamples = 4,
    SampleID = "Sample_Name",
    arrayType = ex$arrayType,
    annotationVersion = ex$annotationVersion,
    scriptLabel = "preprocessingMinfiEwasWater",
    baseDataFolder = file.path(ex$tempDir, "rData"),
    figureBaseDir = file.path(ex$tempDir, "figures"),
    detPThreshold = 1,
    normMethods = "quantile",
    sexColumn = "Sex",
    pvalThreshold = 1,
    chrToRemove = "",
    snpsToRemove = "SBE",
    mafThreshold = 1,
```

```

    crossReactivePath = ex$crossReactivePath,
    plotGroupVar = "Sex",
    lcRef = "saliva",
    phenoOrder = "Sample_Name;Sex;Basename;Sentrrix_ID;Sentrrix_Position",
    lcPhenoDir = ex$tempDir,
    saveOutputs = FALSE,
    verbose = FALSE,
    logs = FALSE
  )
  inherits(result, "dnaEPICO_preprocessingMinfiEwasWater")
}

```

preprocessingPheno	<i>Prepare phenotype and methylation matrices for downstream modeling</i>
--------------------	---

Description

Read the phenotype table and the preprocessed beta, M-value, and copy-number matrices; align them by sample identifier; split them by timepoint; prepare combined longitudinal objects; and build Clock Foundation export tables. The function returns a structured in-memory result, while legacy files are written only when `saveOutputs = TRUE`.

Usage

```

preprocessingPheno(
  phenoFile = "data/preprocessingMinfiEwasWater/phenoLC.csv",
  sepType = "",
  betaPath =
    "rData/preprocessingMinfiEwasWater/metrics/beta_NomFilt_MSetF_Flt_Rxy_Ds_Rc.RData",
  mPath = "rData/preprocessingMinfiEwasWater/metrics/m_NomFilt_MSetF_Flt_Rxy_Ds_Rc.RData",
  cnPath =
    "rData/preprocessingMinfiEwasWater/metrics/cn_NomFilt_MSetF_Flt_Rxy_Ds_Rc.RData",
  SampleID = "Sample_Name",
  timeVar = "Timepoint",
  timepoints = "1,2",
  combineTimepoints = "1,2",
  outputPheno = "data/preprocessingPheno",
  outputRData = "rData/preprocessingPheno/metrics",
  outputRDataMerge = "rData/preprocessingPheno/mergeData",
  sexColumn = "Sex",
  outputLogs = "logs",
  outputDir = "data/preprocessingPheno",
  verbose = FALSE,
  logs = FALSE,
  saveOutputs = FALSE
)

```

Arguments

phenoFile	Character. Path to the phenotype CSV file.
-----------	--

<code>sepType</code>	Character. Field separator used in <code>phenoFile</code> . Use <code>","</code> for a comma-separated file, <code>"\\t"</code> for a tab-delimited file, or another separator accepted by <code>utils::read.csv()</code> .
<code>betaPath</code>	Character. Path to the saved beta-value object. Both <code>.RData</code> and <code>.rds</code> files are supported.
<code>mPath</code>	Character. Path to the saved M-value object. Both <code>.RData</code> and <code>.rds</code> files are supported.
<code>cnPath</code>	Character. Path to the saved copy-number object. Both <code>.RData</code> and <code>.rds</code> files are supported.
<code>SampleID</code>	Character. Name of the phenotype column containing sample identifiers used to align phenotype and methylation data.
<code>timeVar</code>	Character. Name of the phenotype column containing timepoint labels.
<code>timepoints</code>	Character vector or comma-separated string of timepoints to retain and split into separate in-memory subsets.
<code>combineTimepoints</code>	Character vector or comma-separated string of timepoints to combine into the longitudinal phenotype-plus-beta object.
<code>outputPheno</code>	Character. Directory used for saved phenotype CSV files when <code>saveOutputs = TRUE</code> .
<code>outputRData</code>	Character. Directory used for saved metric <code>.RData</code> files when <code>saveOutputs = TRUE</code> .
<code>outputRDataMerge</code>	Character. Directory used for saved merged phenotype-plus-beta <code>.RData</code> files when <code>saveOutputs = TRUE</code> .
<code>sexColumn</code>	Character. Name of the phenotype sex column used when building Clock Foundation exports.
<code>outputLogs</code>	Character. Directory used for log files when <code>logs = TRUE</code> .
<code>outputDir</code>	Character. Directory used for Clock Foundation export files when <code>saveOutputs = TRUE</code> .
<code>verbose</code>	Logical. If <code>TRUE</code> , emit progress messages with <code>message()</code> . The default is <code>FALSE</code> .
<code>logs</code>	Logical. If <code>TRUE</code> , write the same progress messages to <code>outputLogs</code> . The default is <code>FALSE</code> .
<code>saveOutputs</code>	Logical. If <code>TRUE</code> , write the legacy CSV, ZIP, and <code>.RData</code> outputs to disk. The default is <code>FALSE</code> , so the function can be used in the more traditional in-memory Bioconductor style.

Value

A list with class `"dnaEPICO_preprocessingPheno"`.

pheno Phenotype table read from `phenoFile`.

metricsData Object returned by `loadMetricsPreprocessingPheno()` containing the beta-value, M-value, and copy-number matrices loaded from `betaPath`, `mPath`, and `cnPath`.

timepointData Object returned by `splitTimepointsPreprocessingPheno()` containing per-timepoint phenotype tables and methylation matrices.

combinedData Object returned by `combineTimepointsPreprocessingPheno()` containing the merged longitudinal phenotype-plus-beta object and the timepoint combination metadata.

clockFoundation Object returned by `buildClockFoundationInputsPreprocessingPheno()` containing the beta table and phenotype table prepared for Clock Foundation export.

savedFiles Object returned by `writePreprocessingPhenoOutputs()` when `saveOutputs = TRUE`, otherwise `NULL`.

logFile Resolved path to the optional log file, or `NULL` when logging was disabled.

See [dnaEPICO_preprocessingPheno](#) for a class-level overview.

See Also

[dnaEPICO_preprocessingPheno](#)

Examples

```
tmp <- tempdir()
pheno <- data.frame(
  Sample_Name = c("S1", "S2", "S3"),
  Timepoint = c("1", "1", "2"),
  Sex = c(0, 1, 0),
  stringsAsFactors = FALSE
)
beta <- matrix(
  c(0.10, 0.20, 0.30, 0.40, 0.50, 0.60),
  nrow = 2,
  dimnames = list(c("cg1", "cg2"), pheno$Sample_Name)
)
m <- beta * 10
cn <- beta * 100
pheno_file <- file.path(tmp, "pheno.csv")
beta_path <- file.path(tmp, "beta.RData")
m_path <- file.path(tmp, "m.RData")
cn_path <- file.path(tmp, "cn.RData")
utils::write.csv(pheno, pheno_file, row.names = FALSE)
save(beta, file = beta_path)
save(m, file = m_path)
save(cn, file = cn_path)
result <- preprocessingPheno(
  phenoFile = pheno_file,
  betaPath = beta_path,
  mPath = m_path,
  cnPath = cn_path,
  SampleID = "Sample_Name",
  timeVar = "Timepoint",
  timepoints = "1,2",
  combineTimepoints = "1,2",
  outputPheno = file.path(tmp, "data", "preprocessingPheno"),
  outputRData = file.path(tmp, "rData", "preprocessingPheno", "metrics"),
  outputRDataMerge = file.path(tmp, "rData", "preprocessingPheno", "mergeData"),
  sexColumn = "Sex",
  outputLogs = file.path(tmp, "logs"),
  outputDir = file.path(tmp, "clockFoundation"),
  saveOutputs = FALSE
)
stopifnot(inherits(result, "dnaEPICO_preprocessingPheno"))
```

```
print.dnaEPICO_dnamReport  
Print a DNA methylation report result
```

Description

Print a DNA methylation report result

Usage

```
## S3 method for class 'dnaEPICO_dnamReport'  
print(x, ...)
```

Arguments

x	Object returned by <code>dnamReport()</code> .
...	Additional arguments ignored.

Value

Invisibly returns x.

```
readPhenotypeTargets Read phenotype targets for shared dnaEPICO workflows
```

Description

Read the phenotype table used by shared dnaEPICO workflows, validate the sample identifier column, optionally subset the first nSamples, and return the targets as a base data.frame.

Usage

```
readPhenotypeTargets(  
  phenoFile,  
  sepType = "",  
  nSamples = NA,  
  SampleID = "Sample_Name",  
  verbose = FALSE,  
  logs = FALSE,  
  log_dir = NULL,  
  log_file = "log_readPhenotypeTargets.txt"  
)
```

Arguments

phenoFile	Character. Path to the phenotype table on disk.
sepType	Character. Field separator used in phenoFile. Use "" (default) for a standard comma-separated file, "\\t" for a tab-delimited file, or another single-character separator accepted by <code>utils::read.csv()</code> .
nSamples	Integer or NA. Number of rows to keep from the start of the phenotype table. The default NA reads and returns all rows.
SampleID	Character. Name of the column containing sample identifiers that will later be used to name methylation-array samples.
verbose	Logical. If TRUE, emit progress and preview messages with <code>message()</code> . The default is FALSE, so the function is quiet unless the user explicitly requests messages.
logs	Logical. If TRUE, write the same progress messages to a log file. The default is FALSE.
log_dir	Character or NULL. Directory where the log file should be written when <code>logs = TRUE</code> . If NULL, the current working directory is used.
log_file	Character. File name used when <code>logs = TRUE</code> . The default is "log_readPhenotypeTargets.txt".

Value

A data.frame containing the phenotype targets.

Examples

```
tmp <- tempdir()
pheno <- data.frame(
  Sample_Name = c("S1", "S2"),
  Sex = c("F", "M"),
  stringsAsFactors = FALSE
)
pheno_file <- file.path(tmp, "pheno.csv")
utils::write.csv(pheno, pheno_file, row.names = FALSE)
targets <- readPhenotypeTargets(
  phenoFile = pheno_file,
  SampleID = "Sample_Name"
)
stopifnot(is.data.frame(targets))
stopifnot(nrow(targets) == 2L)
```

readRGSetMinfiEwasWater

Read IDAT files into an annotated RGChannelSet

Description

Read methylation-array IDAT files with `minfi::read.metharray.exp()`, set sample names from the phenotype table, apply the requested annotation, and return the resulting `RGChannelSet`.

Usage

```
readRGSetMinfiEwasWater(
  idatFolder,
  targets,
  SampleID = "Sample_Name",
  arrayType = "IlluminaHumanMethylationEPICv2",
  annotationVersion = "20a1.hg38",
  verbose = FALSE,
  logs = FALSE,
  log_dir = NULL,
  log_file = "log_readRGSetMinfiEwasWater.txt"
)
```

Arguments

idatFolder	Character. Directory containing the IDAT files.
targets	Data frame returned by readPhenotypeTargets() or an equivalent phenotype table.
SampleID	Character. Name of the phenotype column containing sample identifiers used to label the RGChannelSet.
arrayType	Character. Array name passed to Biobase::annotation(RGSet), for example "IlluminaHumanMethylationEPICv2".
annotationVersion	Character. Annotation build passed to Biobase::annotation(RGSet), for example "20a1.hg38" for EPIC v2 hg38 annotations or "ilmn12.hg19" for 450K hg19 annotations.
verbose	Logical. If TRUE, emit progress messages with message().
logs	Logical. If TRUE, write the same messages to a log file.
log_dir	Character or NULL. Directory used for the log file when logs = TRUE.
log_file	Character. File name used when logs = TRUE.

Value

An annotated RGChannelSet.

Examples

```
if (requireNamespace("minfiData", quietly = TRUE) &&
    requireNamespace("IlluminaHumanMethylation450kmanifest", quietly = TRUE) &&
    requireNamespace("IlluminaHumanMethylation450kanno.ilmn12.hg19", quietly = TRUE)) {
  ex <- dnaEPICo::exampleMinfiIdatInputsDnaEpico(n = 4)
  rgset <- readRGSetMinfiEwasWater(
    idatFolder = ex$idatFolder,
    targets = ex$targets,
    SampleID = "Sample_Name",
    arrayType = ex$arrayType,
    annotationVersion = ex$annotationVersion
  )
  class(rgset)
}
```

renderDnamReport	<i>Render a prepared DNA methylation report</i>
------------------	---

Description

Render a prepared DNA methylation report

Usage

```
renderDnamReport(
  preparedReport,
  verbose = FALSE,
  logs = FALSE,
  logDir = NULL,
  clean = TRUE
)
```

Arguments

preparedReport	Object returned by prepareDnamReportInputs().
verbose	Logical. If TRUE, emit progress messages.
logs	Logical. If TRUE, write progress messages to a log file.
logDir	Character or NULL. Directory for optional log files.
clean	Logical. Retained for backwards compatibility.

Value

A list with class "dnaEPICO_dnamReport_render".

Examples

```
report_root <- file.path(tempdir(), "dnaepico-render-example")
prepared <- prepareDnamReportInputs(
  outputDir = file.path(report_root, "reports")
)
rendered <- renderDnamReport(prepared)
rendered$status
```

splitTimepointsPreprocessingPheno	<i>Split phenotype and methylation data by timepoint</i>
-----------------------------------	--

Description

Align phenotype rows and metric matrices for each requested timepoint, and precompute the per-timepoint phenotype-plus-beta objects used by downstream modeling functions.

Usage

```
splitTimepointsPreprocessingPheno(
  pheno,
  metricsData,
  SampleID = "Sample_Name",
  timeVar = "Timepoint",
  timepoints = "1,2",
  verbose = FALSE,
  logs = FALSE,
  log_dir = NULL,
  log_file = "log_splitTimepointsPreprocessingPheno.txt"
)
```

Arguments

pheno	Data frame containing phenotype information.
metricsData	Object returned by <code>loadMetricsPreprocessingPheno()</code> .
SampleID	Character. Name of the sample identifier column in pheno.
timeVar	Character. Name of the timepoint column in pheno.
timepoints	Character vector or comma-separated string of timepoints to retain.
verbose	Logical. If TRUE, emit progress messages with <code>message()</code> .
logs	Logical. If TRUE, write the same messages to a log file.
log_dir	Character or NULL. Directory used for the log file when <code>logs = TRUE</code> .
log_file	Character. File name used when <code>logs = TRUE</code> .

Value

A list with class "dnaEPICO_preprocessingPheno_timepoints" containing the parsed timepoints and aligned per-timepoint subsets.

Examples

```
ex <- dnaEPICO:::examplePreprocessingPhenoStateDnaEpico()
timepoint_data <- splitTimepointsPreprocessingPheno(
  pheno = ex$pheno,
  metricsData = ex$metricsData,
  SampleID = "Sample_Name",
  timeVar = "Timepoint",
  timepoints = "1,2",
  verbose = FALSE,
  logs = FALSE
)
timepoint_data$timepoints
```

```
summarizeMethylationGLMM_T1T2Models
```

Summarize CpG-wise mixed-effects model fits for longitudinal analyses

Description

Extract phenotype-specific fixed-effect tables from the fitted mixed-effects model object returned by `fitMethylationGLMM_T1T2Models()`.

Usage

```
summarizeMethylationGLMM_T1T2Models(
  modelResults,
  preparedData,
  summaryPval = NA,
  nCores = 1L,
  chunkSize = NULL,
  verbose = FALSE,
  logs = FALSE,
  log_dir = NULL,
  log_file = "log_methylationGLMM_T1T2.txt"
)
```

Arguments

<code>modelResults</code>	Object returned by <code>fitMethylationGLMM_T1T2Models()</code> .
<code>preparedData</code>	Object returned by <code>prepareMethylationGLMM_T1T2Data()</code> .
<code>summaryPval</code>	Numeric or NA. Optional p-value filter applied to the returned summary tables. NA keeps all rows.
<code>nCores</code>	Integer. Number of worker processes to use while extracting summary rows.
<code>chunkSize</code>	Integer or NULL. Number of CpGs processed per parallel chunk. NULL chooses a value automatically.
<code>verbose</code>	Logical. If TRUE, emit progress messages with <code>message()</code> .
<code>logs</code>	Logical. If TRUE, write the same messages to a log file.
<code>log_dir</code>	Character or NULL. Directory used for the log file when <code>logs = TRUE</code> .
<code>log_file</code>	Character. File name used when <code>logs = TRUE</code> .

Value

A list with class `"dnaEPICO_methylationGLMM_T1T2_summaries"` containing one CpG-level summary data frame per phenotype.

Examples

```
ex <- dnaEPICO:::exampleMethylationGLMMStateDnaEpiCo()
summary_results <- summarizeMethylationGLMM_T1T2Models(
  modelResults = ex$modelResults,
  preparedData = ex$preparedData,
```

```

summaryPval = NA,
nCores = 1,
verbose = FALSE,
logs = FALSE
)
names(summary_results$summaries)

```

```
summarizeMethylationGLM_T1Models
```

Summarize CpG-wise Gaussian GLM fits for one-timepoint analyses

Description

Extract phenotype-specific CpG coefficient tables from the fitted model object returned by `fitMethylationGLM_T1Models()`.

Usage

```

summarizeMethylationGLM_T1Models(
  modelResults,
  preparedData,
  summaryResidualSD = TRUE,
  summaryPval = NA,
  nCores = 1L,
  libPath = NULL,
  glmLibs = "glm2",
  chunkSize = NULL,
  verbose = FALSE,
  logs = FALSE,
  log_dir = NULL,
  log_file = "log_methylationGLM_T1.txt"
)

```

Arguments

<code>modelResults</code>	Object returned by <code>fitMethylationGLM_T1Models()</code> .
<code>preparedData</code>	Object returned by <code>prepareMethylationGLM_T1Data()</code> .
<code>summaryResidualSD</code>	Logical. If TRUE, add residual standard deviations to each CpG summary row.
<code>summaryPval</code>	Numeric or NA. Optional p-value filter applied to the returned summary tables. NA keeps all rows.
<code>nCores</code>	Integer. Number of worker processes to use while extracting summary rows.
<code>libPath</code>	Character vector or NULL. Optional library paths forwarded to worker processes.
<code>glmLibs</code>	Character vector or comma-separated string of package names to check on worker processes. The default is "glm2".
<code>chunkSize</code>	Integer or NULL. Number of CpGs to process per parallel chunk. NULL chooses a value automatically.
<code>verbose</code>	Logical. If TRUE, emit progress messages with <code>message()</code> .
<code>logs</code>	Logical. If TRUE, write the same messages to a log file.
<code>log_dir</code>	Character or NULL. Directory used for the log file when <code>logs = TRUE</code> .
<code>log_file</code>	Character. File name used when <code>logs = TRUE</code> .

Value

A list with class "dnaEPICO_methylationGLM_T1_summaries" containing one CpG-level summary data frame per phenotype.

Examples

```
ex <- dnaEPICO::exampleMethylationGLMStateDnaEpico()
summary_results <- summarizeMethylationGLM_T1Models(
  modelResults = ex$modelResults,
  preparedData = ex$preparedData,
  summaryResidualSD = TRUE,
  summaryPval = NA,
  nCores = 1,
  verbose = FALSE,
  logs = FALSE
)
names(summary_results$summaries)
```

```
summarizeTimepointsMethylationGLMM_T1T2
```

Summarize phenotype values by timepoint for longitudinal methylation analyses

Description

Summarize the requested phenotype variables by timepoint. Numeric phenotypes are reported with mean, standard deviation, and non-missing counts; non-numeric phenotypes are reported with non-missing counts and the observed levels.

Usage

```
summarizeTimepointsMethylationGLMM_T1T2(
  data,
  timeVar = "Timepoint",
  phenotypes,
  verbose = FALSE,
  logs = FALSE,
  log_dir = NULL,
  log_file = "log_methylationGLMM_T1T2.txt"
)
```

Arguments

data	Data frame containing the longitudinal phenotype-plus-beta data.
timeVar	Character. Name of the time variable.
phenotypes	Character vector or comma-separated string of phenotype variables to summarize.
verbose	Logical. If TRUE, emit progress messages with message().
logs	Logical. If TRUE, write the same messages to a log file.
log_dir	Character or NULL. Directory used for the log file when logs = TRUE.
log_file	Character. File name used when logs = TRUE.

Value

A data frame with one row per timepoint and summary columns for each requested phenotype.

Examples

```
ex <- dnaEPICO::exampleMethylationGLMMStateDnaEpico()
timepoint_summary <- summarizeTimepointsMethylationGLMM_T1T2(
  data = ex$preparedData$data,
  timeVar = "Timepoint",
  phenotypes = "score",
  verbose = FALSE,
  logs = FALSE
)
nrow(timepoint_summary)
```

svaEnmix

Estimate surrogate variables from ENmix control probes

Description

Read the phenotype table and a saved RGChannelSet, estimate surrogate variables from ENmix control probes, analyze their association with Sentrix chip and position factors, and return a structured in-memory result. Legacy CSV, .RData, text-summary, and figure outputs are written only when saveOutputs = TRUE.

Usage

```
svaEnmix(
  phenoFile = "data/preprocessingMinfiEwasWater/phenoLC.csv",
  rgsetData = "rData/preprocessingMinfiEwasWater/objects/RGSet.RData",
  sepType = "",
  outputLogs = "logs",
  nSamples = NA,
  SampleID = "Sample_Name",
  arrayType = "IlluminaHumanMethylationEPICv2",
  annotationVersion = "20a1.hg38",
  SentrixIDColumn = "Sentrix_ID",
  SentrixPositionColumn = "Sentrix_Position",
  ctrlSvaPercVar = 0.9,
  ctrlSvaFlag = 1,
  scriptLabel = "svaEnmix",
  tiffWidth = 2000,
  tiffHeight = 1000,
  tiffRes = 150,
  figureBaseDir = "figures",
  dataBaseDir = "data",
  rBaseDir = "rData",
  display = FALSE,
  verbose = FALSE,
  logs = FALSE,
  saveOutputs = FALSE
)
```

Arguments

phenoFile	Character. Path to the phenotype file with cell-composition data.
rgsetData	Character. Path to a saved RGChannelSet object. Both .RData and .rds files are supported.
sepType	Character. Field separator used in phenoFile. Use "," for a comma-separated file, "\\t" for a tab-delimited file, or another separator accepted by <code>utils::read.csv()</code> .
outputLogs	Character. Directory used for log files when <code>logs = TRUE</code> .
nSamples	Integer or NA. Number of rows to keep from the phenotype table. Use NA to keep all samples.
SampleID	Character. Name of the phenotype column containing sample identifiers.
arrayType	Character. Illumina array identifier assigned to <code>Biobase::annotation(RGSet)</code> .
annotationVersion	Character. Annotation build assigned to <code>Biobase::annotation(RGSet)</code> .
SentrixIDColumn	Character. Name of the chip identifier column in the phenotype data.
SentrixPositionColumn	Character. Name of the chip position column in the phenotype data.
ctrlSvaPercVar	Numeric. Proportion of control-probe variance explained when running <code>ENmix::ctrlsva()</code> .
ctrlSvaFlag	Integer. Control-probe flag passed to <code>ENmix::ctrlsva()</code> .
scriptLabel	Character. Label used to name output folders when <code>saveOutputs = TRUE</code> .
tiffWidth	Integer. Width of saved TIFF plots in pixels.
tiffHeight	Integer. Height of saved TIFF plots in pixels.
tiffRes	Integer. Resolution in DPI for saved TIFF plots.
figureBaseDir	Character. Base directory used for saved figure outputs when <code>saveOutputs = TRUE</code> .
dataBaseDir	Character. Base directory used for saved CSV and text outputs when <code>saveOutputs = TRUE</code> .
rBaseDir	Character. Base directory used for saved .RData outputs when <code>saveOutputs = TRUE</code> .
display	Logical. If TRUE, draw plots on the active graphics device.
verbose	Logical. If TRUE, emit progress messages with <code>message()</code> . The default is FALSE.
logs	Logical. If TRUE, write the same progress messages to outputLogs. The default is FALSE.
saveOutputs	Logical. If TRUE, write the legacy CSV, .RData, text, and TIFF outputs to disk. The default is FALSE.

Value

A list with class "dnaEPICO_svaEnmix".

targets Phenotype table read from `phenoFile` after any optional row subsetting.

RGSet Loaded `RGChannelSet` with sample names realigned to `targets[[SampleID]]`.

svaData Object returned by `estimateSvaEnmixControls()` containing the surrogate-variable matrix and the control-probe settings used to estimate it.

- mergedPheno** Phenotype table returned by `mergeSvaTargetsEnmix()` after the surrogate variables were appended as additional columns.
- analysisData** Object returned by `analyzeSvaEnmix()` containing the surrogate-variable association models, ANOVA tables, and Sentrix metadata.
- plotFiles** Named list describing the plot file paths requested for the SVA figures. When `saveOutputs = FALSE`, the entries are typically NULL.
- savedFiles** Object returned by `writeSvaEnmixOutputs()` when `saveOutputs = TRUE`, otherwise NULL.
- logFile** Resolved path to the optional log file, or NULL when logging was disabled.
- See [dnaEPICO_svaEnmix](#) for a class-level overview.

See Also

[dnaEPICO_svaEnmix](#)

Examples

```
tmp <- tempdir()
stopifnot(dir.exists(tmp))

if (requireNamespace("minfiData", quietly = TRUE)) {
  ex <- dnaEPICO::exampleMinfiBaseDataDnaEpico()
  pheno_file <- file.path(tmp, "pheno.csv")
  rgset_path <- file.path(tmp, "RGSet.RData")
  RGSet <- ex$RGSet
  utils::write.csv(ex$targets, pheno_file, row.names = FALSE)
  save(RGSet, file = rgset_path)
  sva_result <- svaEnmix(
    phenoFile = pheno_file,
    rgsetData = rgset_path,
    SampleID = "Sample_Name",
    arrayType = "IlluminaHumanMethylation450k",
    annotationVersion = "ilmn12.hg19",
    SentrixIDColumn = "Sentrix_ID",
    SentrixPositionColumn = "Sentrix_Position",
    outputLogs = file.path(tmp, "logs"),
    figureBaseDir = file.path(tmp, "figures"),
    dataBaseDir = file.path(tmp, "data"),
    rBaseDir = file.path(tmp, "rData"),
    saveOutputs = FALSE
  )
  stopifnot(inherits(sva_result, "dnaEPICO_svaEnmix"))
}
```

writeMethylationGLMM_T1T2Outputs

Write optional disk outputs for longitudinal mixed-effects analyses

Description

Write optional serialized outputs, summary tables, significant interaction tables, and annotated results from the longitudinal mixed-effects workflow.

Usage

```
writeMethylationGLMM_T1T2Outputs(
  modelResults,
  modelSummaries,
  annotatedResults,
  significantInteractions = NULL,
  outputRData,
  summaryTxtDir,
  significantInteractionDir,
  annotatedLMEOut,
  saveTxtSummaries = TRUE,
  saveSignificantInteractions = FALSE,
  verbose = FALSE,
  logs = FALSE,
  log_dir = NULL,
  log_file = "log_methylationGLMM_T1T2.txt"
)
```

Arguments

`modelResults` Object returned by `fitMethylationGLMM_T1T2Models()`.

`modelSummaries` Object returned by `summarizeMethylationGLMM_T1T2Models()`.

`annotatedResults` Object returned by `annotateMethylationGLMM_T1T2Summaries()` or a compatible data frame.

`significantInteractions` Object returned by `collectSignificantInteractionsMethylationGLMM_T1T2()` or `NULL`.

`outputRData` Character. Directory used for serialized model and summary outputs.

`summaryTxtDir` Character. Directory used for tab-delimited summary tables.

`significantInteractionDir` Character. Directory used for significant interaction coefficient tables.

`annotatedLMEOut` Character. Directory used for the annotated summary CSV file.

`saveTxtSummaries` Logical. If `TRUE`, write tab-delimited summary tables.

`saveSignificantInteractions` Logical. If `TRUE`, write significant interaction coefficient tables.

`verbose` Logical. If `TRUE`, emit progress messages with `message()`.

`logs` Logical. If `TRUE`, write the same messages to a log file.

`log_dir` Character or `NULL`. Directory used for the log file when `logs = TRUE`.

`log_file` Character. File name used when `logs = TRUE`.

Value

A list with class `"dnaEPIC0_methylationGLMM_T1T2_paths"` containing the paths of the files written to disk.

Examples

```

ex <- dnaEPICO:::exampleMethylationGLMMStateDnaEpico()
annotation_data <- annotateMethylationGLMM_T1T2Summaries(
  modelSummaries = ex$modelSummaries,
  annotationObject = ex$annotationData,
  annotationCols = "Name,chr,pos",
  verbose = FALSE,
  logs = FALSE
)
significant_hits <- collectSignificantInteractionsMethylationGLMM_T1T2(
  modelResults = ex$modelResults,
  pvalThreshold = 1,
  verbose = FALSE,
  logs = FALSE
)
output_paths <- writeMethylationGLMM_T1T2Outputs(
  modelResults = ex$modelResults,
  modelSummaries = ex$modelSummaries,
  annotatedResults = annotation_data,
  significantInteractions = significant_hits,
  outputRData = file.path(ex$tempDir, "models"),
  summaryTxtDir = file.path(ex$tempDir, "summary"),
  significantInteractionDir = file.path(ex$tempDir, "significant"),
  annotatedLMEOut = file.path(ex$tempDir, "annotated"),
  saveTxtSummaries = TRUE,
  saveSignificantInteractions = TRUE,
  verbose = FALSE,
  logs = FALSE
)
names(output_paths)

```

```
writeMethylationGLM_T1Outputs
```

Write optional disk outputs for one-timepoint GLM analyses

Description

Write optional serialized outputs, summary tables, significant-CpG tables, and annotated results from the one-timepoint GLM workflow.

Usage

```

writeMethylationGLM_T1Outputs(
  modelResults,
  modelSummaries,
  annotatedResults,
  significantCpGs = NULL,
  outputRData,
  summaryTxtDir,
  significantCpGDir,
  annotatedGLMOut,
  saveTxtSummaries = TRUE,

```

```

    saveSignificantCpGs = FALSE,
    verbose = FALSE,
    logs = FALSE,
    log_dir = NULL,
    log_file = "log_methylationGLM_T1.txt"
  )

```

Arguments

modelResults Object returned by `fitMethylationGLM_T1Models()`.

modelSummaries Object returned by `summarizeMethylationGLM_T1Models()`.

annotatedResults Object returned by `annotateMethylationGLM_T1Summaries()` or a compatible data frame.

significantCpGs Object returned by `collectSignificantCpGsMethylationGLM_T1()` or `NULL`.

outputRData Character. Directory used for serialized model and summary outputs.

summaryTxtDir Character. Directory used for tab-delimited summary tables.

significantCpGDir Character. Directory used for significant-CpG coefficient tables.

annotatedGLMOut Character. Directory used for the annotated summary CSV file.

saveTxtSummaries Logical. If `TRUE`, write tab-delimited summary tables.

saveSignificantCpGs Logical. If `TRUE`, write significant-CpG coefficient tables.

verbose Logical. If `TRUE`, emit progress messages with `message()`.

logs Logical. If `TRUE`, write the same messages to a log file.

log_dir Character or `NULL`. Directory used for the log file when `logs = TRUE`.

log_file Character. File name used when `logs = TRUE`.

Value

A list with class `"dnaEPICO_methylationGLM_T1_paths"` containing the paths of the files written to disk.

Examples

```

ex <- dnaEPICO::exampleMethylationGLMStateDnaEpico()
annotation_data <- annotateMethylationGLM_T1Summaries(
  modelSummaries = ex$modelSummaries,
  annotationObject = ex$annotationData,
  annotationCols = "Name,chr,pos",
  verbose = FALSE,
  logs = FALSE
)
significant_cpGs <- collectSignificantCpGsMethylationGLM_T1(
  modelResults = ex$modelResults,
  pvalThreshold = 1,
  verbose = FALSE,
  logs = FALSE
)

```

```

)
output_paths <- writeMethylationGLM_T1Outputs(
  modelResults = ex$modelResults,
  modelSummaries = ex$modelSummaries,
  annotatedResults = annotation_data,
  significantCpGs = significant_cpGs,
  outputRData = file.path(ex$tempDir, "models"),
  summaryTxtDir = file.path(ex$tempDir, "summary"),
  significantCpGDir = file.path(ex$tempDir, "significant"),
  annotatedGLMOut = file.path(ex$tempDir, "annotated"),
  saveTxtSummaries = TRUE,
  saveSignificantCpGs = TRUE,
  verbose = FALSE,
  logs = FALSE
)
names(output_paths)

```

```
writePhenoLCMinfiEwasWater
```

Write the merged phenotype plus cell-composition table

Description

Write the merged phenotype plus cell-composition table

Usage

```

writePhenoLCMinfiEwasWater(
  lcData,
  file,
  verbose = FALSE,
  logs = FALSE,
  log_dir = NULL,
  log_file = "log_writePhenoLCMinfiEwasWater.txt"
)

```

Arguments

lcData	Object returned by estimateLCMinfiEwasWater().
file	Character. Path to the CSV file to write.
verbose	Logical. If TRUE, emit progress messages with message().
logs	Logical. If TRUE, write the same messages to a log file.
log_dir	Character or NULL. Directory used for the log file when logs = TRUE.
log_file	Character. File name used when logs = TRUE.

Value

Invisibly returns file.

Examples

```

ref_file <- system.file("extdata", "saliva.txt", package = "dnaEPIC0")
beta <- as.matrix(utils::read.table(ref_file))[1:20, , drop = FALSE]
colnames(beta) <- c("sample1", "sample2")
targets <- data.frame(
  Sample_Name = colnames(beta),
  Timepoint = c("T1", "T2"),
  stringsAsFactors = FALSE
)
lc_data <- estimateLCMinfiEwasWater(
  beta = beta,
  targets = targets,
  lcRef = "saliva",
  phenoOrder = "Sample_Name;Timepoint"
)
output_file <- file.path(tempdir(), "phenoLC.csv")
writePhenoLCMinfiEwasWater(lcData = lc_data, file = output_file)
file.exists(output_file)

```

```
writePreprocessingPhenoOutputs
```

Write legacy preprocessingPheno outputs to disk

Description

Write the legacy CSV, ZIP, and .RData outputs produced by preprocessingPheno(). This helper keeps file writing separate from the in-memory preprocessing steps.

Usage

```

writePreprocessingPhenoOutputs(
  preprocessingData,
  outputPheno = "data/preprocessingPheno",
  outputRData = "rData/preprocessingPheno/metrics",
  outputRDataMerge = "rData/preprocessingPheno/mergeData",
  outputDir = "data/preprocessingPheno",
  verbose = FALSE,
  logs = FALSE,
  log_dir = NULL,
  log_file = "log_writePreprocessingPhenoOutputs.txt"
)

```

Arguments

preprocessingData	Object returned by preprocessingPheno() or a list with the same components.
outputPheno	Character. Directory used for saved phenotype CSV files.
outputRData	Character. Directory used for saved metric .RData files.
outputRDataMerge	Character. Directory used for saved merged phenotype-plus-beta .RData files.

outputDir	Character. Directory used for the Clock Foundation export files.
verbose	Logical. If TRUE, emit progress messages with message().
logs	Logical. If TRUE, write the same messages to a log file.
log_dir	Character or NULL. Directory used for the log file when logs = TRUE.
log_file	Character. File name used when logs = TRUE.

Value

A list with class "dnaEPICO_preprocessingPheno_paths" containing the paths written to disk.

Examples

```
ex <- dnaEPICO:::examplePreprocessingPhenoStateDnaEpico()
output_paths <- writePreprocessingPhenoOutputs(
  preprocessingData = ex$preprocessingData,
  outputPheno = file.path(ex$tempDir, "pheno"),
  outputRData = file.path(ex$tempDir, "metrics"),
  outputRDataMerge = file.path(ex$tempDir, "merge"),
  outputDir = file.path(ex$tempDir, "clock"),
  verbose = FALSE,
  logs = FALSE
)
names(output_paths)
```

writeSvaEnmixOutputs *Write svaEnmix outputs to disk*

Description

Write the legacy CSV, .RData, and text-summary outputs used by the original svaEnmix() workflow.

Usage

```
writeSvaEnmixOutputs(
  svaData,
  mergedPheno,
  analysisData = NULL,
  phenoFile = NULL,
  dataBaseDir = "data",
  rBaseDir = "rData",
  scriptLabel = "svaEnmix",
  verbose = FALSE,
  logs = FALSE,
  log_dir = NULL,
  log_file = "log_writeSvaEnmixOutputs.txt"
)
```

Arguments

svaData	Object returned by estimateSvaEnmixControls().
mergedPheno	Phenotype data frame returned by mergeSvaTargetsEnmix().
analysisData	Optional object returned by analyzeSvaEnmix().
phenoFile	Character or NULL. When supplied, mergedPheno is written back to this path for legacy compatibility.
dataBaseDir	Character. Base directory used for saved data outputs.
rBaseDir	Character. Base directory used for saved .RData outputs.
scriptLabel	Character. Label used to create the output subdirectory.
verbose	Logical. If TRUE, emit progress messages with message().
logs	Logical. If TRUE, write the same messages to a log file.
log_dir	Character or NULL. Directory used for the log file when logs = TRUE.
log_file	Character. File name used when logs = TRUE.

Value

A list with class "dnaEPICO_svaEnmix_paths" containing the paths written to disk.

Examples

```
ex <- dnaEPICO:::exampleSvaAnalysisStateDnaEpico()
temp_dir <- tempdir()
output_paths <- writeSvaEnmixOutputs(
  svaData = list(sva = ex$sva),
  mergedPheno = ex$mergedPheno,
  analysisData = ex$analysisData,
  phenoFile = file.path(temp_dir, "phenoLC.csv"),
  dataBaseDir = file.path(temp_dir, "data"),
  rBaseDir = file.path(temp_dir, "rData"),
  scriptLabel = "svaEnmixExample",
  verbose = FALSE,
  logs = FALSE
)
names(output_paths)
```

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