Package ‘methylInheritance’

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Contents

methylInheritance-package ................................................................. 2
calculateSignificantLevel ................................................................. 3
createDataStructure ................................................................. 4
createOutputDir ................................................................. 5
demoForTransgenerationalAnalysis .................................................. 6
extractInfo ................................................................. 8
formatInputMethylData ................................................................. 9
getGRangesFromMethylDiff ............................................................ 10
interGeneration ................................................................. 11
isInterGenerationResults ............................................................. 12
loadAllRDSResults ................................................................. 13
loadConvergenceData ............................................................... 14
mergePermutationAndObservation ................................................... 15
methylInheritanceResults ............................................................ 18
plotConvergenceGraph ............................................................... 22
plotGraph ................................................................. 23
print.methylInheritanceAllResults ................................................. 24
readInterGenerationResults .......................................................... 25
runObservation ................................................................. 26
runOnePermutationOnAllGenerations ................................................. 28
runPermutation ................................................................. 31
samplesForTransgenerationalAnalysis ............................................. 35
saveInterGenerationResults .......................................................... 36
validateExtractInfo ................................................................. 37
validateLoadConvergenceData .......................................................... 38
validateMergePermutationAndObservation ....................................... 39
validateRunObservation .............................................................. 40
validateRunPermutation .............................................................. 43

Index  46
**calculateSignificantLevel**

**Description**

This package does a permutation analysis, based on Monte Carlo sampling, for testing the hypothesis that the number of conserved differentially methylated elements (sites or tiles), between several generations, is associated to an effect inherited from a treatment and that stochastic effect can be dismissed.

**Author(s)**

Astrid Deschênes, Pascal Belleau and Arnaud Droit

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**See Also**

- `runPermutation` for running a permutation analysis, and optionally an observation analysis, on a specified multi-generational dataset
- `runObservation` for running an observation analysis on a specified multi-generational dataset

---

**calculateSignificantLevel**

*Calculate significant level for hypo and hyper conserved elements*

**Description**

Calculate significant level for hypo and hyper conserved elements using permutation results as well as observed results

**Usage**

`calculateSignificantLevel(formatForGraphDataFrame)`

**Arguments**

- `formatForGraphDataFrame` a data.frame containing the observation results (using real data) and the permutation results (using shuffled data). Both hyper and hypo differentially conserved methylation results must be present. The data.frame must have 3 columns: "TYPE", "RESULT" and "SOURCE". The "TYPE" can be either "HYPER" or "HYPO". The "RESULT" is the number of conserved differentially elements. The "SOURCE" can be either "OBSERVATION" or "PERMUTATION".

**Value**

a list containing two elements:

- `HYPER` a double, the significant level for the hyper differentially methylated conserved elements
- `HYPO` a double, the significant level for the hypo differentially methylated conserved elements
createDataStructure

Author(s)
Astrid Deschenes, Pascal Belleau

Examples

```r
## Loading dataset containing all results
data(methylInheritanceResults)

## Extract information for the intersection between conserved differentially methylated sites (type = sites) between the intersection of 2 generations (inter = i2): F2 and F3 (position = 2)
info <- extractInfo(allResults = methylInheritanceResults, type = "sites", inter="i2", 2)

## Create graph
methylInheritance:::calculateSignificantLevel(info)
```

---

createDataStructure  Extract the number of conserved differentially methylated elements in GRanges.

Description

Extract the number of conserved differentially methylated elements in GRanges. Each GRanges is the result of one intersection between two or more consecutive generations for one analysis done on all generations. The hypo and hyper differentially methylated elements are counted separately.

Usage

`createDataStructure(interGenerationGR)`

Arguments

`interGenerationGR`

a list that contains the information for all differentially methylated analysis done on each generation present in the initial dataset. The list must contain the following elements:
- `i2` a list of GRanges. Each GRanges represents the intersection of analysis results between two consecutive generations. The first element represents the intersection of the first and second generations; the second element, the intersection of the second and third generations; etc. The number of entries depends on the number of generations.
- `iAll` a list of GRanges. Each GRanges represents the intersection of the analysis results between three or more consecutive generations. The first element represents the intersection of the first three generations; the second element, the intersection of the first fourth generations; etc. The number of entries depends on the number of generations.
Value

a list containing the following elements:

- i2 a list containing:
  - HYPER a list of integer, the number of conserved hyper differentially methylated sites between two consecutive generations. The first element represents the intersection of the first and second generations; the second element, the intersection of the second and third generations; etc..
  - HYPO a list of integer, the number of conserved hypo differentially methylated sites between two consecutive generations. The first element represents the intersection of the first and second generations; the second element, the intersection of the second and third generations; etc..

- iAll a list containing:
  - HYPER a list of integer, the number of conserved hyper differentially methylated sites between three or more consecutive generations. The first element represents the intersection of the first three generations; the second element, the intersection of the first four generations; etc..The number of entries depends of the number of generations.
  - HYPO a list of integer, the number of conserved hypo differentially methylated sites between three or more consecutive generations. The first element represents the intersection of the first three generations; the second element, the intersection of the first four generations; etc..The number of entries depends of the number of generations.

Author(s)

Astrid Deschenes, Pascal Belleau

Examples

```r
## Get the name of the directory where the file is stored
filesDir <- system.file("extdata", "TEST", package="methylInheritance")

## Load file containing results from a observation analysis
obsResults <- readRDS(file = paste0(filesDir, "/SITES/SITES_observed_results.RDS"))

## Create data structure using information form the observation analysis
formatedResults <- methylInheritance:::createDataStructure(obsResults)
```

createOutputDir

Create directories that will contained the results of the permutations in RDS format.

Description

Create directories that will contained the results of the permutations in RDS format.
Usage

createOutputDir(
  outputDir,
  doingSites = TRUE,
  doingTiles = FALSE,
  saveInfoByGeneration
)

Arguments

outputDir      a string of character, the name of the main directory to be created.
doingSites     a logical, a directory consecrated to contain the results of the permutation analysis for sites is created when doingSites = TRUE. Default: TRUE.
doingTiles     a logical, a directory consecrated to contain the results of the permutation analysis for tiles is created when doingTiles = TRUE. Default: FALSE.
saveInfoByGeneration
                a logical, when TRUE, the information about differentially methylated sites and tiles for each generation is saved in a RDS file. The information is saved in a different file for each permutation.

Value

0 when all directories are created without problem.

Author(s)

Astrid Deschenes

Examples

## Create an output directory for SITES only
methylInheritance:::createOutputDir(outputDir = "testSites",
                                      doingSites = TRUE, doingTiles = FALSE, saveInfoByGeneration = TRUE)

demoForTransgenerationalAnalysis

The methylation information from samples over three generations. Information for each generation is stored in a methylRawList format (for demo purpose).

Description

The object is a list with 3 entries. Each entry corresponds to the information for one generation (first entry = first generation, etc.) stored in a methylRawList object. There are 12 samples (6 controls and 6 cases) for each generation. Each sample information is stored in a methylRaw object.
Usage

data(demoForTransgenerationalAnalysis)

Format

A list containing three methylRawList objects. Each methylRawList contains the information for one generation (first entry = first generation, etc.). Each sample information is stored in a methylRaw object. There is methylRaw objects (6 controls and 6 cases) in each generation.

Details

This dataset can be used to test runPermutation and runObservation functions.

Value

A list containing three methylRawList objects. Each methylRawList contains the information for one generation (first entry = first generation, etc.). Each sample information is stored in a methylRaw object. There is methylRaw objects (6 controls and 6 cases) in each generation.

See Also

- runPermutation for running a permutation analysis, and optionally an observation analysis, using multi-generational dataset
- runObservation for running an observation analysis using methylKit info entry

Examples

```r
## Loading dataset
data(demoForTransgenerationalAnalysis)

## Run a permutation analysis
runObservation(methylKitData = demoForTransgenerationalAnalysis,
               outputDir = "test_demo", type = "tiles", vSeed = 2001)

## Get results
result <- loadAllRDSResults(analysisResultsDir = "test_demo",
                            permutationResultsDir = NULL, doingSites = FALSE,
                            doingTiles = TRUE)

## Remove result directory
if (dir.exists("test_demo")) {
  unlink("test_demo", recursive = TRUE)
}
```
extractInfo  
Extract the information specific to a subsection of the permutation analysis

Description

Extract the information specific to a subsection of the permutation analysis. The extracted information will be specific to one type of differential methylation analysis (tiles or sites), to one type of intersection (two consecutive generation or more) and to one specific group of generations.

Usage

```r
extractInfo(
  allResults,
  type = c("sites", "tiles"),
  inter = c("i2", "iAll"),
  position = 1
)
```

Arguments

- `allResults`: a list of class `methylInheritanceAllResults` as created by the `runPermutation` function. The list must contain two entries: "PERMUTATION" and "OBSERVATION". The "PERMUTATION" list must contain all results from all permutations while the "OBSERVATION" list must contain the result obtained with the observed dataset (not shuffled).

- `type`: One of the "sites" or "tiles" strings. Specifies the type of differentially methylated elements should be returned. For retrieving differentially methylated bases `type = "sites"`; for differentially methylated regions `type = "tiles"`. Default: "sites".

- `inter`: One of the "i2" or "iAll" strings. Specifies the type of intersection should be returned. For retrieving intersection results between two consecutive generations `inter = "i2"`; for intersection results between three generations or more `inter = "iAll"`. Default: "i2".

- `position`: a positive integer, the position in the list where the information will be extracted. Default=1.

Value

A `data.frame` containing the observation results (using real data) and the permutation results (using shuffled data). Both hyper and hypo differentially conserved methylation results are present.

Author(s)

Astrid Deschenes, Pascal Belleau
Examples

```r
## Get the name of the directory where files are storedilesDir <- system.file("extdata", "TEST", package="methylInheritance")

## Load information from files
results <- loadAllRDSResults(analysisResultsDir = filesDir,
                          permutationResultsDir = filesDir, doingSites = TRUE, doingTiles = TRUE)

## Extract information for the intersection between conserved differentially
## methylated sites (type = sites) between the intersection of 2
## generations (inter = i2): F1 and F2 (position = 1)
info <- extractInfo(allResults = results, type = "sites", inter="i2", 1)
```

formatInputMethylData  

**Description**

Permute dataset and format it to be ready for an analysis

**Usage**

```r
formatInputMethylData(methylKitData)
```

**Arguments**

- `methylKitData`  
  a list of methylRawList entries. Each methylRawList entry must contain all
  the methylRaw entries related to one generation (first entry = first generation,
  second entry = second generation, etc.). The number of generations must cor-
  respond to the number of entries in the methylKitData. At least 2 generations
  must be present to make a permutation analysis. More information can be found
  in the methylKit package.

**Value**

- a list of methylRawList entries.

**Author(s)**

Astrid Deschenes, Pascal Belleau

**Examples**

```r
## Load dataset
data("samplesForTransgenerationalAnalysis")
methylInheritance:::formatInputMethylData(samplesForTransgenerationalAnalysis)
```
getGRangesFromMethylDiff

Transform results from a CpG site or region analysis done on multiple generations into a list of GRanges objects

Description
Transform a list of methylDiff objects into a list of GRanges objects. Each methylDiff object represent a CpG site or region analysis done on one generation.

Usage
getGRangesFromMethylDiff(
  methDiff,
  pDiff,
  qvalue,
  type = c("all", "hyper", "hypo")
)

Arguments
methDiff a list of S4 methylDiff class objects, each entry of the list represents the differentially methylated results for one generation (first entry = first generation, second entry = second generation, etc.). Each methylDiff object holds statistics and locations for differentially methylated regions/bases.
pDiff a positive double between 0 and 100, the cutoff for absolute value of methylation percentage change between test and control.
qvalue a positive double inferior to 1, the cutoff for qvalue of differential methylation statistic.
type One of the "hyper", "hypo" or "all" strings, the string specifies what type of differentially methylated bases/tiles should be treated: For retrieving hypermethylated tiles/sites type = "hyper"; for hypo-methylated type = "hypo". Default: "all".

Value
a list of GRanges objects, each entry of the list represents the differentially methylated results for one generation (first entry = first generation, second entry = second generation, etc.). Each GRanges object holds statistics for differentially methylated regions/bases.

Author(s)
Pascal Belleau
Examples

```r
## Load permutation results on sites
permutationResultsFile <- system.file("extdata",
   "permutationResultsForSites.RDS", package="methylInheritance")
permutationResults <- readRDS(permutationResultsFile)

## Transform result to GRanges
resultsGR <- methylInheritance:::getGRangesFromMethylDiff(methDiff =
   permutationResults, pDiff = 10, qvalue = 0.01, type = "hyper")
```

---

**interGeneration**  
*Calculate the intersection of the differentially methylated results for two or more consecutive generations*

---

**Description**

Calculate the intersection of the differentially methylated results for two or more consecutive generations using a list of GRanges where each entry represents the results for one generation.

**Usage**

```r
interGeneration(resultAllGenGR)
```

**Arguments**

- `resultAllGenGR` a list of GRanges as created by the `getGRangesFromMethylDiff` function. Each entry of the list represents the differentially methylated results for one generation (first entry = first generation, second entry = second generation, etc.). Each GRanges object holds statistics for differentially methylated regions/bases.

**Value**

a list containing the following elements:

- `i2` a list of GRanges Each GRanges represents the intersection of analysis results between two consecutive generations. The first element represents the intersection of the first and second generations; the second element, the intersection of the second and third generations; etc. The number of entries depends of the number of generations.

- `iAll` a list of GRanges. Each GRanges represents the intersection of the analysis results between three or more consecutive generations. The first element represents the intersection of the first three generations; the second element, the intersection of the first fourth generations; etc.. The number of entries depends of the number of generations.

**Author(s)**

Pascal Belleau, Astrid Deschenes
Examples

```r
## Load permutation results on sites
permutationResultsFile <- system.file("extdata", "permutationResultsForSites.RDS", package="methylInheritance")
permutationResults <- readRDS(permutationResultsFile)

## Transform result to GRanges
resultsGR <- methylInheritance:::getGRangesFromMethylDiff(methDiff = permutationResults, pDiff = 10, qvalue = 0.01, type = "hyper")

## Extract inter generational conserved sites
conservedSitesGR <- methylInheritance:::interGeneration(resultsGR)
```

isInterGenerationResults

Verify if a specific file containing intergenerational results exists or not.

Description

Verify if a specific file containing intergenerational results exists or not.

Usage

```r
isInterGenerationResults(outputDir, permutationID, type = c("sites", "tiles"))
```

Arguments

- **outputDir**: a string of character, the name of the directory that will contain the results of the permutation. The name should end with a slash. The directory should already exists.
- **permutationID**: an integer, the identifiant of the permutation. When the permutationID = 0, the results are considered as the observed results and are saved in a file with the ".observed_results.RDS" extension. When the permutationID != 0, the results are considered as permutation results and are saved in a file with the ".permutation_permutationID.RDS" extension.
- **type**: One of the "sites" or "tiles" strings. Specifies the type of differentially methylated elements should be saved. Default: "sites".

Value

TRUE when file present; otherwise FALSE.

Author(s)

Astrid Deschenes, Pascal Belleau
**loadAllRDSResults**

Load all RDS files created by the permutation and observation analysis. The function returns an object of class "methylInheritanceAllResults" that holds all the pertinent information.

### Examples

```r
## Get the name of the directory where the file is stored
filesDir <- system.file("extdata", "TEST", package="methylInheritance")

## Verify that DMS intergenerational results for the observed data exists
methylInheritance:::isInterGenerationResults(outputDir =
  paste0(filesDir, "/"), 0, "sites")
```

### Description

Load all RDS files created by the permutation and observation analysis. The function returns an object of class "methylInheritanceAllResults" that holds all the pertinent information.

### Usage

```r
loadAllRDSResults(
  analysisResultsDir,
  permutationResultsDir,
  doingSites = TRUE,
  doingTiles = FALSE,
  maxID = NA
)
```

### Arguments

- **analysisResultsDir**
  
  a character string, the path to the directory that contains the analysis results. The path can be the same as for the `permutationResultsDir` parameter. When NULL, the observation results are not loaded. Default = NULL.

- **permutationResultsDir**
  
  a character string, the path to the directory that contains the permutation results. The path can be the same as for the `analysisResultsDir` parameter. When NULL, the permutation results are not loaded. Default = NULL.

- **doingSites**
  
  a logical, the data related to differentially methylated sites are loaded when doingSites = TRUE. Default: TRUE.

- **doingTiles**
  
  a logical, the data related to differentially methylated tiles are loaded when doingTiles = TRUE. Default: TRUE.

- **maxID**
  
  NA or a positive integer, the maximum identification number of the permutation files to be loaded. When NA, all files present in the directory are loaded. Default: NA.
loadConvergenceData

Value

A list of class methylInheritanceAllResults containing the result of the observation analysis as well as the results of all the permutations.

Author(s)

Astrid Deschenes, Pascal Belleau

See Also

mergePermutationAndObservation for detail description, in the Value section, of the methylInheritanceAllResults object.

Examples

```r
## Get the name of the directory where files are stored
filesDir <- system.file("extdata", "TEST", package="methylInheritance")

## Load information from files
results <- loadAllRDSResults(analysisResultsDir = filesDir,
   permutationResultsDir = filesDir, doingSites = TRUE, doingTiles = TRUE)

## Print the observation results
results

## Access the results for the first permutation only for sites
results$PERMUTATION[[1]]$SITES
```

loadConvergenceData  Load convergence information from RDS files

Description

Load convergence information from RDS files.

Usage

```r
loadConvergenceData(
   analysisResultsDir, permutationResultsDir,
   type = c("sites", "tiles"),
   inter = c("i2", "iAll"),
   position,
   by = 100
)
```
Arguments

analysisResultsDir
a character string, the path to the directory that contains the analysis results. The path can be the same as for the permutationResultsDir parameter.

permutationResultsDir
a character string, the path to the directory that contains the permutation results. The path can be the same as for the analysisResultsDir parameter.

type
One of the "sites" or "tiles" strings. Specifies the type of differentially methylated elements should be returned. For retrieving differentially methylated bases type = "sites"; for differentially methylated regions type = "tiles". Default: "sites".

inter
One of the "i2" or "iAll" strings. Specifies the type of intersection should be returned. For retrieving intersection results between two consecutive generations inter = "i2"; for intersection results between three generations or more inter = "iAll". Default: "i2".

position
a positive integer, the position in the list where the information will be extracted.

by
a integer, the increment of the number of permutations where the significant level is tested. Default: 100.

Value

a graph showing the evolution of the significant level with the number of permutations

Author(s)

Astrid Deschenes, Pascal Belleau

Examples

```r
## Get the name of the directory where files are stored
filesDir <- system.file("extdata", "TEST", package="methylInheritance")

## Load convergence information
results <- loadConvergenceData(analysisResultsDir = filesDir,
permutationResultsDir = filesDir, type="sites", inter="i2", position=1,
by=1)
```

mergePermutationAndObservation

Merge the permutation results with the observation results.

Description

Merge the permutation results with the observation results. The merging is only needed when permutation and observation have been processed separately. The returned value is a methylInheritanceAllResults object that can be used by the extractInfo function.
Usage

mergePermutationAndObservation(permutationResults, observationResults)

Arguments

permutationResults

a list with 1 entry called PERMUTATION. The PERMUTATION entry is a list with
a number of entries corresponding to the number of permutations that have been
processed. Each entry contains the result of one permutation.

observationResults

a list with 1 entry called OBSERVATION. The OBSERVATION entry is a list
containing the result obtained with the observed dataset (not shuffled).

Value

a list of class methylInheritanceAllResults with 2 entries. The 2 entries are:

• PERMUTATION list with a number of entries corresponding to the number of permutations
  that have been processed. Each entry contains the result of one permutation. The elements in
each entry are:
  – SITES Only present when a sites analysis has been achieved, a list containing:
    * i2 a list containing:
      • HYPER a list of integer, the number of conserved hyper differentially methylated
        sites between two consecutive generations. The first element represents the inter-
        section of the first and second generations; the second element, the intersection of
        the second and third generations; etc.
      • HYPO a list of integer, the number of conserved hypo differentially methylated
        sites between two consecutive generations. The first element represents the inter-
        section of the first and second generations; the second element, the intersection of
        the second and third generations; etc.
    * iAll a list containing:
      • HYPER a list of integer, the number of conserved hyper differentially methylated
        sites between three or more consecutive generations. The first element represents the
        intersection of the first three generations; the second element, the intersection of
        the first fourth generations; etc. The number of entries depends on the number
        of generations.
      • HYPO a list of integer, the number of conserved hypo differentially methylated
        sites between three or more consecutive generations. The first element represents
        the intersection of the first three generations; the second element, the intersection
        of the first fourth generations; etc. The number of entries depends on the number
        of generations.
  – TILES Only present when a tiles analysis has been achieved, a list containing:
    * i2 a list containing:
      • HYPER a list of integer, the number of conserved hyper differentially methylated
        positions between two consecutive generations. The first element represents the
        intersection of the first and second generations; the second element, the inter-
        section of the second and third generations; etc.
- **HYP0** a list of integer, the number of conserved hypo differentially methylated positions between two consecutive generations. The first element represents the intersection of the first and second generations; the second element, the intersection of the second and third generations; etc.

- **iAll** a list containing:
  - **HYPER** a list of integer, the number of conserved hyper differentially methylated positions between three or more consecutive generations. The first element represents the intersection of the first three generations; the second element, the intersection of the first fourth generations; etc. The number of entries depends on the number of generations.
  - **HYP0** a list of integer, the number of conserved hypo differentially methylated positions between three or more consecutive generations. The first element represents the intersection of the first three generations; the second element, the intersection of the first fourth generations; etc. The number of entries depends on the number of generations.

- **OBSERVATION** a list containing the result obtained with the observed dataset (not shuffled). The elements are:
  - **SITES** Only present when a sites analysis has been achieved, a list containing:
    - **i2** a list containing:
      - **HYPER** a list of integer, the number of conserved hyper differentially methylated sites between two consecutive generations. The first element represents the intersection of the first and second generations; the second element, the intersection of the second and third generations; etc.
      - **HYP0** a list of integer, the number of conserved hypo differentially methylated sites between two consecutive generations. The first element represents the intersection of the first and second generations; the second element, the intersection of the second and third generations; etc.

  - **iAll** a list containing:
    - **HYPER** a list of integer, the number of conserved hyper differentially methylated sites between three or more consecutive generations. The first element represents the intersection of the first three generations; the second element, the intersection of the first fourth generations; etc. The number of entries depends on the number of generations.
    - **HYP0** a list of integer, the number of conserved hypo differentially methylated sites between three or more consecutive generations. The first element represents the intersection of the first three generations; the second element, the intersection of the first fourth generations; etc. The number of entries depends on the number of generations.

  - **TILES** Only present when a tiles analysis has been achieved, a list containing:
    - **i2** a list containing:
      - **HYPER** a list of integer, the number of conserved hyper differentially methylated positions between two consecutive generations. The first element represents the intersection of the first and second generations; the second element, the intersection of the second and third generations; etc.
      - **HYP0** a list of integer, the number of conserved hypo differentially methylated positions between two consecutive generations. The first element represents the in-
tersection of the first and second generations; the second element, the intersection of the second and third generations; etc.

* iAll a list containing:
  · HYPER a list of integer, the number of conserved hyper differentially methylated positions between three or more consecutive generations. The first element represents the intersection of the first three generations; the second element, the intersection of the first fourth generations; etc. The number of entries depends on the number of generations.
  · HYPO a list of integer, the number of conserved hypo differentially methylated positions between three or more consecutive generations. The first element represents the intersection of the first three generations; the second element, the intersection of the first fourth generations; etc. The number of entries depends on the number of generations.

**Author(s)**
Astrid Deschenes, Pascal Belleau

**Examples**

```r
## Create an observation result
observed <- list()
observed[["OBSERVATION"]][["SITES"]] <- list()
observed[["OBSERVATION"]][["SITES"]][["i2"]]
  <- list(HYPER = list(11, 10), HYPO = list(13, 12))
observed[["OBSERVATION"]][["SITES"]][["iAll"]] <- list(HYPER = list(1), HYPO = list(3))

## Create a permutation result containing only 1 permutation result
## Real permutation results would have more entries
permutated <- list()
permutated[["PERMUTATION"]][["SITES"]] <- list()
permutated[["PERMUTATION"]][["SITES"]][["i2"]]
  <- list(HYPER = list(11, 12), HYPO = list(8, 11))
permutated[["PERMUTATION"]][["SITES"]][["iAll"]] <- list(HYPER = list(0), HYPO = list(1))

## Merge permutation and observation results
mergePermutationAndObservation(permutationResults = permutated,
                                observationResults = observed)
```

**methylInheritanceResults**

All observed and permutation results formatted in a methylInheritanceResults class (for demo purpose).
**methylInheritanceResults**

**Description**

The object is a list with 2 entries: "OBSERVATION" and "PERMUTATION".

**Usage**

`data(methylInheritanceResults)`

**Format**

A list of class `methylInheritanceAllResults` containing the following elements:

- **OBSERVATION** a list containing:
  - **SITES** a list containing:
    - **i2** a list containing:
      - **HYPER** a list of integer with 2 entries, the number of conserved hyper differentially methylated sites between two consecutive generations. The first element represents the intersection of the first and second generations; the second element, the intersection of the second and third generations.
      - **HYPO** a list of integer with 2 entries, the number of conserved hypo differentially methylated sites between two consecutive generations. The first element represents the intersection of the first and second generations; the second element, the intersection of the second and third generations.
    - **iAll** a list containing:
      - **HYPER** a list of integer with 1 entry, the number of conserved hyper differentially methylated sites between the three consecutive generations.
      - **HYPO** a list of integer with 1 entry, the number of conserved hypo differentially methylated sites between the three consecutive generations.
  - **TILES** a list containing:
    - **i2** a list containing:
      - **HYPER** a list of integer with 2 entries, the number of conserved hyper differentially methylated positions between two consecutive generations. The first element represents the intersection of the first and second generations; the second element, the intersection of the second and third generations.
      - **HYPO** a list of integer with 2 entries, the number of conserved hypo differentially methylated positions between two consecutive generations. The first element represents the intersection of the first and second generations; the second element, the intersection of the second and third generations.
    - **iAll** a list containing:
      - **HYPER** a list of integer with 1 entry, the number of conserved hyper differentially methylated positions between the three consecutive generations.
      - **HYPO** a list of integer with 1 entry, the number of conserved hypo differentially methylated positions between the three consecutive generations.

- **PERMUTATION** a list containing `nbrPermutations` entries. Each entry is a list containing:
  - **SITES** a list containing:
    - **i2** a list containing:
· **HYPER** a list of integer with 2 entries, the number of conserved hyper differentially methylated sites between two consecutive generations. The first element represents the intersection of the first and second generations; the second element, the intersection of the second and third generations.

· **HYPO** a list of integer with 2 entries, the number of conserved hypo differentially methylated sites between two consecutive generations. The first element represents the intersection of the first and second generations; the second element, the intersection of the second and third generations.

* **iAll** a list containing:
  · **HYPER** a list of integer with 1 entry, the number of conserved hyper differentially methylated sites between the three consecutive generations.
  · **HYPO** a list of integer with 1 entry, the number of conserved hypo differentially methylated sites between the three consecutive generations.

  – **TILES** a list containing:
    * **i2** a list containing:
      · **HYPER** a list of integer with 2 entries, the number of conserved hyper differentially methylated positions between two consecutive generations. The first element represents the intersection of the first and second generations; the second element, the intersection of the second and third generations.
      · **HYPO** a list of integer with 2 entries, the number of conserved hypo differentially methylated positions between two consecutive generations. The first element represents the intersection of the first and second generations; the second element, the intersection of the second and third generations.

Details

This dataset can be used to test the `extractInfo` function. The extracted information can be used to calculate the significant level or to create a graph.

Value

a list of class `methylInheritanceAllResults` containing the following elements:

· **OBSERVATION** a list containing:
  
  – **SITES** a list containing:
    * **i2** a list containing:
      · **HYPER** a list of integer with 2 entries, the number of conserved hyper differentially methylated sites between two consecutive generations. The first element represents the intersection of the first and second generations; the second element, the intersection of the second and third generations.
- HYPO a list of integer with 2 entries, the number of conserved hypo differentially methylated sites between two consecutive generations. The first element represents the intersection of the first and second generations; the second element, the intersection of the second and third generations.

* iAll a list containing:
  - HYPER a list of integer with 1 entry, the number of conserved hyper differentially methylated sites between the three consecutive generations.
  - HYPO a list of integer with 1 entry, the number of conserved hypo differentially methylated sites between the three consecutive generations.

- TILES a list containing:
  * i2 a list containing:
    - HYPER a list of integer with 2 entries, the number of conserved hyper differentially methylated positions between two consecutive generations. The first element represents the intersection of the first and second generations; the second element, the intersection of the second and third generations.
    - HYPO a list of integer with 2 entries, the number of conserved hypo differentially methylated positions between two consecutive generations. The first element represents the intersection of the first and second generations; the second element, the intersection of the second and third generations.

* iAll a list containing:
  - HYPER a list of integer with 1 entry, the number of conserved hyper differentially methylated positions between the three consecutive generations.
  - HYPO a list of integer with 1 entry, the number of conserved hypo differentially methylated positions between the three consecutive generations.

- PERMUTATION a list containing a number of entries corresponding to the number of permutations that have been produced. Each entry is a list containing:

  - SITES a list containing:
    * i2 a list containing:
      - HYPER a list of integer with 2 entries, the number of conserved hyper differentially methylated sites between two consecutive generations. The first element represents the intersection of the first and second generations; the second element, the intersection of the second and third generations.
      - HYPO a list of integer with 2 entries, the number of conserved hypo differentially methylated sites between two consecutive generations. The first element represents the intersection of the first and second generations; the second element, the intersection of the second and third generations.

  * iAll a list containing:
    - HYPER a list of integer with 1 entry, the number of conserved hyper differentially methylated sites between the three consecutive generations.
    - HYPO a list of integer with 1 entry, the number of conserved hypo differentially methylated sites between the three consecutive generations.

  - TILES a list containing:
    * i2 a list containing:
      - HYPER a list of integer with 2 entries, the number of conserved hyper differentially methylated positions between two consecutive generations. The first element
plotConvergenceGraph

Generate a graph showing the convergence for a permutation analysis

Description

Generate a graph showing the convergence for a permutation analysis using observed and permuted results.

Usage

plotConvergenceGraph(dataFrameConvergence)

Arguments

dataFrameConvergence

a data.frame containing the significant levels at different number of cycles (total number of permuted data analysed). The data.frame must have 6 columns: "NBR_PERMUTATIONS", "ELEMENT", "ANALYSIS", "POSITION", "TYPE" and "SIGNIFICANT_LEVEL". The "ELEMENT" can be either "SITES" or "TILES". The "TYPE" can be either "HYPER" or "HYPO".
**plotGraph**

*Generate a graph for a permutation analysis*

**Value**

a ggplot object.

**Author(s)**

Astrid Deschenes, Pascal Belleau

**Examples**

```r
## Get the name of the directory where files are stored
filesDir <- system.file("extdata", "TEST", package="methylInheritance")

## Extract convergence information for F1 and F2 and F3
data <- loadConvergenceData(analysisResultsDir = filesDir,
permutationResultsDir = filesDir, type = "sites", inter = "iAll",
position = 1, by = 1)

## Create convergence graph
plotConvergenceGraph(data)
```

**Description**

Generate a graph for a permutation analysis using observed and shuffled results.

**Usage**

`plotGraph(formatForGraphDataFrame)`

**Arguments**

`formatForGraphDataFrame`

a data.frame containing the observation results (using real data) and the permutation results (using shuffled data). Both hyper and hypo differentially conserved methylation results must be present. The data.frame must have 3 columns: "TYPE", "RESULT" and "SOURCE". The "TYPE" can be either "HYPER" or "HYPO". The "RESULT" is the number of conserved differentially elements. The "SOURCE" can be either "OBSERVATION" or "PERMUTATION".

**Value**

a graph showing the permutation analysis results

**Author(s)**

Astrid Deschenes, Pascal Belleau
## Loading dataset containing all results
```r
data(methylInheritanceResults)
```

## Extract information for the intersection between conserved differentially methylated sites (type = sites) between the intersection of 2 generations (inter = i2): F2 and F3 (position = 2)
```r
info <- extractInfo(allResults = methylInheritanceResults, 
                     type = "sites", inter="i2", 2)
```

## Create graph
```r
plotGraph(info)
```

### Description
Print a `methylInheritanceAllResults` object

### Usage

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

### Arguments

- `x`  
  the output object from `mergePermutationAndObservation` function, `runPermutationUsingRDSFile` function (when `runObservationAnalysis = TRUE` and `runPermutationUsingMethylKitInfo` function (when `runObservationAnalysis = TRUE` to be printed

- `...`  
  arguments passed to or from other methods

### Value
an object of class `methylInheritanceAllResults`

### Examples

```r
## Load dataset
data("methylInheritanceResults")

## Print dataset
print(methylInheritanceResults)
```
**readInterGenerationResults**

*Read and return intergenerational results contained in a RDS file*

---

**Description**

Read and return intergenerational results contained in a RDS file

**Usage**

```r
greadInterGenerationResults(
  outputDir,
  permutationID,
  type = c("sites", "tiles")
)
```

**Arguments**

- `outputDir` a string of character, the name of the directory that will contain the results of the permutation. The name should end with a slash. The directory should already exist.
- `permutationID` an integer, the identifier of the permutation. When the `permutationID` = 0, the results are considered as the observed results and are saved in a file with the ".observed_results.RDS" extension. When the `permutationID` != 0, the results are considered as permutation results and are saved in a file with the ".permutation_permutationID.RDS" extension.
- `type` One of the "sites" or "tiles" strings. Specifies the type of differentially methylated elements should be saved. Default: "sites".

**Value**

a list containing the intergenerational results for the specified permutation.

**Author(s)**

Astrid Deschenes, Pascal Belleau

**Examples**

```r
## Get the name of the directory where the file is stored
filesDir <- system.file("extdata", "TEST", package="methylInheritance")

## Read DMS intergenerational results for the observed data
methylInheritance:::readInterGenerationResults(outputDir =
paste0(filesDir, "/"), 0, "sites")
```
runObservation  Run a differential methylation analysis on multi-generational dataset

Description

Run a differential methylation analysis on each generation present in a dataset. The number of conserved differentially methylated elements (sites, tile or both) between generations is then calculated. The methylKit package is used to identify the differentially methylated elements.

The multi-generational dataset or the name of the RDS file that contains the dataset can be used as input.

The results can also be saved in RDS file (optional).

Usage

runObservation(
  methylKitData,
  type = c("both", "sites", "tiles"),
  outputDir = "output",
  nbrCoresDiffMeth = 1,
  minReads = 10,
  minMethDiff = 10,
  qvalue = 0.01,
  maxPercReads = 99.9,
  destrand = FALSE,
  minCovBasesForTiles = 0,
  tileSize = 1000,
  stepSize = 1000,
  vSeed = -1,
  restartCalculation = FALSE,
  saveInfoByGeneration = FALSE
)

Arguments

methylKitData  a list of methylRawList entries or the name of the RDS file containing the list. Each methylRawList contains all the methylRaw entries related to one generation (first entry = first generation, second entry = second generation, etc.). The number of generations must correspond to the number of entries in the methylKitData. At least 2 generations must be present to calculate the conserved elements. More information can be found in the methylKit package.

type  One of the "sites", "tiles" or "both" strings. Specifies the type of differentially methylated elements should be returned. For retrieving differentially methylated bases type="sites"; for differentially methylated regions type="tiles". Default: "both".

outputDir  a string, the name of the directory that will contain the results of the analysis. If the directory does not exist, it will be created. Default: "output".
**runObservation**

**nbrCoresDiffMeth**
- A positive integer, the number of cores to use for parallel differential methylation calculations. The parameter is used for both sites and tiles analysis. The parameter corresponds to the `num.cores` parameter in the package `methylKit`. Default: 1 and always 1 for Windows.

**minReads**
- A positive integer. Bases and regions having lower coverage than this count are discarded. The parameter corresponds to the `lo.count` parameter in the package `methylKit`.

**minMethDiff**
- A positive double between [0,100], the absolute value of methylation percentage change between cases and controls. The parameter corresponds to the `difference` parameter in the `methylKit` package. Default: 10.

**qvalue**
- A positive double between [0,1], the cutoff for qvalue of differential methylation statistics. Default: 0.01.

**maxPercReads**
- A double between [0,100], the percentile of read counts that is going to be used as an upper cutoff. Bases or regions having higher coverage than this percentile are discarded. The parameter is used for both CpG sites and tiles analysis. The parameter corresponds to the `hi.perc` parameter in the package `methylKit`. Default: 99.9.

**destrand**
- A logical, when `TRUE` will merge reads on both strands of a CpG dinucleotide to provide better coverage. Only advised when looking at CpG methylation. Parameter used for both CpG sites and tiles analysis. Default: `FALSE`.

**minCovBasesForTiles**
- A non-negative integer, the minimum number of bases to be covered in a given tiling window. The parameter corresponds to the `cov.bases` parameter in the package `methylKit`. Only used when `doingTiles = TRUE`. Default: 0.

**tileSize**
- A positive integer, the size of the tiling window. The parameter corresponds to the `win.size` parameter in the package `methylKit`. Only used when `doingTiles = TRUE`. Default: 1000.

**stepSize**
- A positive integer, the step size of tiling windows. The parameter corresponds to the `stepSize` parameter in the package `methylKit`. Only used when `doingTiles = TRUE`. Default: 1000.

**vSeed**
- A integer, a seed used when reproducible results are needed. When a value inferior or equal to zero is given, a random integer is used. Default: -1.

**restartCalculation**
- A logical, when `TRUE`, only permutations that don’t have a RDS result final are run. Useful to restart a permutation analysis that has been interrupted. Beware that the parameters have to be identical except for this one.

**saveInfoByGeneration**
- A logical, when `TRUE`, the information about differentially methylated sites and tiles for each generation is saved in a RDS file. The files are saved in the directory specified by the `outputDir` parameter.

**Value**
0.
runOnePermutationOnAllGenerations

**Author(s)**

Astrid Deschenes, Pascal Belleau

**See Also**

`mergePermutationAndObservation` for detail description, in the Value section, of the OBSERVATION section of the methylInheritanceAllResults object.

**Examples**

```r
## Load methylation information
data(samplesForTransgenerationalAnalysis)

## Run an observation analysis
runObservation(methylKitData = samplesForTransgenerationalAnalysis,
               outputDir = "test", type = "sites", vSeed = 221)

## Load the results
results <- loadAllRDSResults(analysisResultsDir = "test",
                              permutationResultsDir = NULL, doingSites = TRUE,
                              doingTiles = FALSE)

## Print the results
results

## Remove directory
if (dir.exists("test")) {
  unlink("test", recursive = TRUE, force = FALSE)
}
```

---

**Description**

Run CpG site or region analysis using the methylKit package for each generation present in the dataset. The intersection of conserved elements is obtained for each group of two consecutive generations, as well as, for larger group subset. The output of the analysis is saved in a RDS file when an directory is specified.

**Usage**

```r
runOnePermutationOnAllGenerations(
  id,
  methylInfoForAllGenerations,
  type = c("both", "sites", "tiles"),
```
```r
runOnePermutationOnAllGenerations

outputDir = NULL,
nbrCoresDiffMeth = 1,
minReads = 10,
minMethDiff = 10,
qvalue = 0.01,
maxPercReads = 99.9,
destrand = FALSE,
minCovBasesForTiles = 0,
tileSize = 1000,
stepSize = 1000,
restartCalculation,
saveInfoByGeneration
)

Arguments

id

an integer, the unique identification of the permutation. When id is 0, the analysis is done on the real dataset.

methylInfoForAllGenerations

a list of methylRawList entries. Each methylRawList entry must contain all the methylRaw entries related to one generation (first entry = first generation, second entry = second generation, etc..). The number of generations must correspond to the number of entries in the methylKitData. At least 2 generations must be present to make a permutation analysis. More information can be found in the methylKit package.

type

One of the "sites","tiles" or "both" strings. Specifies the type of differentially methylated elements should be returned. For retrieving differentially methylated bases type="sites"; for differentially methylated regions type="tiles". Default: "both".

outputDir

a string, the name of the directory that will contain the results of the permutation. If the directory does not exist, it will be created.

nbrCoresDiffMeth

a positive integer, the number of cores to use for parallel differential methylation calculations.Parameter used for both sites and tiles analysis. The parameter corresponds to the num.cores parameter in the package methylKit. Default: 1 and always 1 for Windows.

minReads

a positive integer Bases and regions having lower coverage than this count are discarded. The parameter correspond to the lo.count parameter in the methylKit package.

minMethDiff

a positive integer between [0,100], the absolute value of methylation percentage change between cases and controls. The parameter correspond to the difference parameter in the package methylKit. Default: 10.

qvalue

a positive double inferior to 1, the cutoff for qvalue of differential methylation statistic. Default: 0.01.

maxPercReads

a double between [0-100], the percentile of read counts that is going to be used as upper cutoff. Bases ore regions having higher coverage than this percentile are
discarded. Parameter used for both CpG sites and tiles analysis. The parameter correspond to the hi.perc parameter in the methylKit package. Default: 99.9.

destrand a logical, when TRUE will merge reads on both strands of a CpG dinucleotide to provide better coverage. Only advised when looking at CpG methylation. Parameter used for both sites and tiles analysis. Default: FALSE.

minCovBasesForTiles a non-negative integer, the minimum number of bases to be covered in a given tiling window. The parameter corresponds to the cov.bases parameter in the methylKit package. Only used when doingTiles = TRUE. Default: 0.

tileSize a positive integer, the size of the tiling window. The parameter corresponds to the win.size parameter in the methylKit package. Only used when doingTiles = TRUE. Default: 1000.

stepSize a positive integer, the step size of tiling windows. The parameter corresponds to the stepSize parameter in the methylKit package. Only used when doingTiles = TRUE. Default: 1000.

restartCalculation a logical, when TRUE, only permutations that don’t have a RDS result final are run.

saveInfoByGeneration a logical, when TRUE, the information about differentially methylated sites and tiles for each generation is saved in a RDS file. The information is saved in a different file for each permutation. The files are = saved in the outputDir.

Value

a list containing the following elements:

- **SITES** Only present when type = "sites" or "both", a list containing:
  - i2 a list containing:
    - HYPER a list of integer, the number of conserved hyper differentially methylated sites between two consecutive generations. The first element represents the intersection of the first and second generations; the second element, the intersection of the second and third generations; etc..
    - HYPO a list of integer, the number of conserved hypo differentially methylated sites between two consecutive generations. The first element represents the intersection of the first and second generations; the second element, the intersection of the second and third generations; etc..
  - iAll a list containing:
    - HYPER a list of integer, the number of conserved hyper differentially methylated sites between three or more consecutive generations. The first element represents the intersection of the first three generations; the second element, the intersection of the first fourth generations; etc.. The number of entries depends of the number of generations.
    - HYPO a list of integer, the number of conserved hypo differentially methylated sites between three or more consecutive generations. The first element represents the intersection of the first three generations; the second element, the intersection of the first fourth generations; etc.. The number of entries depends of the number of generations.
- **TILES**: Only present when type = "tiles" or "both", a list containing: itemize
  - **HYPER**: a list of integer, the number of conserved hyper differentially methylated positions between two consecutive generations. The first element represents the intersection of the first and second generations; the second element, the intersection of the second and third generations; etc..
  - **HYPO**: a list of integer, the number of conserved hypo differentially methylated positions between two consecutive generations. The first element represents the intersection of the first and second generations; the second element, the intersection of the second and third generations; etc..

**iAll** a list containing:
- **HYPER**: a list of integer, the number of conserved hyper differentially methylated positions between three or more consecutive generations. The first element represents the intersection of the first three generations; the second element, the intersection of the first four generations; etc.. The number of entries depends of the number of generations.
- **HYPO**: a list of integer, the number of conserved hypo differentially methylated positions between three or more consecutive generations. The first element represents the intersection of the first three generations; the second element, the intersection of the first fourth generations; etc.. The number of entries depends of the number of generations.

**Author(s)**
Astrid Deschenes, Pascal Belleau

**Examples**
```r
## Load methyl information
data(samplesForTransgenerationalAnalysis)

## Run a permutation analysis
methylInheritance:::runOnePermutationOnAllGenerations(id = 2,
methylationInfoForAllGenerations = samplesForTransgenerationalAnalysis,
type = "tiles", outputDir = NULL,
nbrCoresDiffMeth = 1, minReads = 10, minMethDiff = 10, qvalue = 0.01,
maxPercReads = 99.9, destrand = FALSE, minCovBasesForTiles = 0,
tileSize = 1000, stepSize = 1000, restartCalculation = FALSE)
```

---

**runPermutation**  
Run all permutations on the specified multi-generational dataset

**Description**
Run a permutation analysis, based on Monte Carlo sampling, for testing the hypothesis that the number of conserved differentially methylated elements (sites, tiles or both), between several generations, is associated to an effect inherited from a treatment and that stochastic effect can be dismissed.
The multi-generational dataset or the name of the RDS file that contains the dataset can be used as input. The observation analysis can also be run (optional). All permutation results are saved in RDS files.

Usage

```r
runPermutation(
methylKitData,
type = c("both", "sites", "tiles"),
outputDir = "output",
runObservationAnalysis = TRUE,
nbrPermutations = 1000,
nbrCores = 1,
nbrCoresDiffMeth = 1,
minReads = 10,
minMethDiff = 10,
qvalue = 0.01,
maxPercReads = 99.9,
destrand = FALSE,
minCovBasesForTiles = 0,
tileSize = 1000,
stepSize = 1000,
vSeed = -1,
restartCalculation = FALSE,
saveInfoByGeneration = FALSE
)
```

Arguments

- `methylKitData`: a list of methylRawList entries or the name of the RDS file containing the list. Each methylRawList entry must contain all the methylRaw entries related to one generation (first entry = first generation, second entry = second generation, etc.). The number of generations must correspond to the number of entries in the methylKitData. At least 2 generations must be present to make a permutation analysis. More information can be found in the methylKit package.

- `type`: One of the "sites", "tiles" or "both" strings. Specifies the type of differentially methylated elements should be returned. For retrieving differentially methylated bases type="sites"; for differentially methylated regions type="tiles". Default: "both".

- `outputDir`: a string, the name of the directory that will contain the results of the permutation. If the directory does not exist, it will be created. Default: "output".

- `runObservationAnalysis`: a logical. when runObservationAnalysis = TRUE, a CpG analysis on the observed dataset is done. Default: TRUE.

- `nbrPermutations`: a positive integer, the total number of permutations that is going to be done. Default: 1000.
runPermutation

nbrCores a positive integer, the number of cores to use when processing the analysis. Default: 1 and always 1 for Windows.

nbrCoresDiffMeth a positive integer, the number of cores to use for parallel differential methylation calculations. The parameter is used for both sites and tiles analysis. The parameter corresponds to the num.cores parameter in the package methylKit. Default: 1 and always 1 for Windows.

minReads a positive integer Bases and regions having lower coverage than this count are discarded. The parameter corresponds to the lo.count parameter in the package methylKit.

minMethDiff a positive double between [0,100], the absolute value of methylation percentage change between cases and controls. The parameter corresponds to the difference parameter in the methylKit package. Default: 10.

qvalue a positive double between [0,1], the cutoff for qvalue of differential methylation statistics. Default: 0.01.

maxPercReads a double between [0,100], the percentile of read counts that is going to be used as an upper cutoff. Bases or regions having higher coverage than this percentile are discarded. The parameter is used for both CpG sites and tiles analysis. The parameter corresponds to the hi.perc parameter in the package methylKit. Default: 99.9.

destrand a logical, when TRUE will merge reads on both strands of a CpG dinucleotide to provide better coverage. Only advised when looking at CpG methylation. The parameter is used for both CpG sites and tiles analysis. Default: FALSE.

minCovBasesForTiles a non-negative integer, the minimum number of bases to be covered in a given tiling window. The parameter corresponds to the cov.bases parameter in the package methylKit. Only used when doingTiles = TRUE. Default: 0.

tileSize a positive integer, the size of the tiling window. The parameter corresponds to the win.size parameter in the package methylKit. Only used when doingTiles = TRUE. Default: 1000.

stepSize a positive integer, the step size of tiling windows. The parameter corresponds to the stepSize parameter in the package methylKit. Only used when doingTiles = TRUE. Default: 1000.

vSeed a integer, a seed used when reproducible results are needed. When a value inferior or equal to zero is given, a random integer is used. Default: -1.

restartCalculation a logical, when TRUE, only permutations that don’t have an associated RDS result file are run. Useful to restart a permutation analysis that has been interrupted. Beware that the parameters have to be identical except for this one.

saveInfoByGeneration a logical, when TRUE, the information about differentially methylated sites and tiles for each generation is saved in a RDS file. The information is saved in a different file for each permutation. The files are saved in the directory specified by the outputDir parameter.
runPermutation

Value
0.

Author(s)
Astrid Deschenes, Pascal Belleau

See Also
mergePermutationAndObservation for detail description, in the Value section, of the methylInheritanceAllResults object as well as its PERMUTATION section.

Examples
## Load methylKit information
data(samplesForTransgenerationalAnalysis)

## Run a permutation analysis using the methylKit dataset
## A real analysis would require a much higher number of permutations
runPermutation(methylKitData = samplesForTransgenerationalAnalysis, 
   outputDir = "test_01", runObservationAnalysis = FALSE, type = "sites", 
   nbrPermutations = 2, vSeed = 221)

## Get results
results_01 <- loadAllRDSResults(analysisResultsDir = NULL, 
   permutationResultsDir = "test_01", doingSites = TRUE, 
   doingTiles = FALSE)

## Remove results directory
if (dir.exists("test_01")) {
   unlink("test_01", recursive = TRUE, force = TRUE)
}

## Path to a methylKit RDS file
methylFile <- system.file("extdata", "methylObj_001.RDS", 
   package = "methylInheritance")

## Run a permutation analysis using RDS file name
## A real analysis would require a much higher number of permutations
runPermutation(methylKitData = methylFile, type = "tiles", 
   outputDir = "test_02", nbrPermutations = 2, minCovBasesForTiles = 10, 
   vSeed = 2001)

## Get results
results_02 <- loadAllRDSResults(analysisResultsDir = NULL, 
   permutationResultsDir = "test_02", doingSites = FALSE, 
   doingTiles = TRUE)

## Remove results directory
if (dir.exists("test_02")) {
   unlink("test_02", recursive = TRUE, force = TRUE)
}
All samples information, formatted by methylKit, in a methylRawList format (for demo purpose).

Description

The object is a list with 3 entries. Each entry corresponds to the information for one generation (first entry = first generation, etc.) stored in a methylRawList. There are 12 samples (6 controls and 6 cases) for each generation. Each sample information is stored in a methylRaw object.

Usage

data(samplesForTransgenerationalAnalysis)

Format

A list containing three methylRawList objects. Each methylRawList contains the information for one generation (first entry = first generation, etc...). Each sample information is stored in a methylRaw object. There is methylRaw objects (6 controls and 6 cases) in each generation.

Details

This dataset can be used to test the runPermutation function.

Value

A list containing three methylRawList objects. Each methylRawList contains the information for one generation (first entry = first generation, etc...). Each sample information is stored in a methylRaw object. There is methylRaw objects (6 controls and 6 cases) in each generation.

See Also

- runPermutation for running a permutation analysis, and optionally an observation analysis, using multi-generational dataset

Examples

```r
## Loading dataset
data(samplesForTransgenerationalAnalysis)

## Run a permutation analysis
runPermutation(methylKitData = samplesForTransgenerationalAnalysis,
              type = "tiles", nbrPermutations = 2, vSeed = 2332)
```
saveInterGenerationResults

Save the result of on CpG site or tile analysis on all generations. The analysis can come from observed or shuffled dataset. Each case is saved with a different extension.

Description

Save the result of on CpG site or tile analysis on all generations. The results are saved in a RDS file. The analysis can have been done on the observed or shuffled dataset. Each permutation is saved using its identifiant in the file name.

Usage

```r
saveInterGenerationResults(
  outputDir,
  permutationID,
  type = c("sites", "tiles"),
  interGenerationResult
)
```

Arguments

- **outputDir** a string of character, the name of the directory that will contain the results of the permutation. The name should end with a slash. The directory should already exists.
- **permutationID** an integer, the identifiant of the permutation. When the permutationID = 0, the results are considered as the observed results and are saved in a file with the "_observed_results.RDS" extension. When the permutationID != 0, the results are considered as permutation results and are saved in a file with the "_permutation_permutationID.RDS" extension.
- **type** One of the "sites" or "tiles" strings. Specifies the type of differentially methylated elements should be saved. Default: "sites".
- **interGenerationResult** a list that corresponds to the output of the interGeneration function, the result of on CpG site or tile analysis on all generations.

Value

0 indicating that all parameters validations have been successful.

Author(s)

Astrid Deschenes, Pascal Belleau
Examples

```r
## Load permutation results on sites
permutationResultsFile <- system.file("extdata", "permutationResultsForSites.RDS", package="methylInheritance")
permutationResults <- readRDS(permutationResultsFile)

## Transform result to GRanges
resultsGR <- methylInheritance:::getGRangesFromMethylDiff(methDiff = permutationResults, pDiff = 10, qvalue = 0.01, type = "hyper")

## Extract inter-generationally conserved sites
interGenerationResult <- methylInheritance:::interGeneration(resultsGR)

## Create directories
dir.create("TEST", showWarnings = TRUE)
dir.create("TEST/SITES", showWarnings = TRUE)

## Save results
methylInheritance:::saveInterGenerationResults(
outputDir = "TEST/", permutationID=100, type = "sites",
interGenerationResult = interGenerationResult)
```

validateExtractInfo  Validation of some parameters of the extractInfo function

Description

Validation of some parameters needed by the public extractInfo function.

Usage

`validateExtractInfo(allResults, type, inter, position)`

Arguments

- `allResults` a list as created by the runPermutation or the loadAllRDSResults functions.
- `type` One of the "sites" or "tiles" strings. Specifies the type of differentially methylated elements should be returned. For retrieving differentially methylated bases type = "sites"; for differentially methylated regions type = "tiles".
- `inter` One of the "i2" or "iAll" strings. Specifies the type of intersection should be returned. For retrieving intersection results between two consecutive generations inter = "i2"; for intersection results between three generations or more inter = "iAll".
- `position` a positive integer, the position in the list where the information will be extracted. The position must be an existing position inside allResults.
validateLoadConvergenceData

Validation of some parameters of the \texttt{loadConvergenceData} function

Description

Validation of some parameters needed by the public \texttt{loadConvergenceData} function.

Usage

\begin{verbatim}
validateLoadConvergenceData(
    analysisResultsDir,
    permutationResultsDir,
    position,
    by
)
\end{verbatim}

Arguments

- \texttt{analysisResultsDir} \\
a character string, the path to the directory that contains the analysis results. The path can be the same as for the \texttt{permutationResultsDir} parameter.

- \texttt{permutationResultsDir} \\
a character string, the path to the directory that contains the permutation results. The path can be the same as for the \texttt{analysisResultsDir} parameter.

Value

0 indicating that all parameters validations have been successful.

Author(s)

Astrid Deschenes

Examples

\begin{verbatim}
## Load dataset
data(methylInheritanceResults)

## The function returns 0 when all parameters are valid
methylInheritance:::validateExtractInfo(
    allResults = methylInheritanceResults, type = "sites",
    inter = "i2", 2)

## The function raises an error when at least one parameter is not valid
## Not run: methylInheritance:::validateExtractInfo(
##    allResults = methylInheritanceResults, type = "sites",
##    inter = "i2", 12)
## End(Not run)
\end{verbatim}
validateMergePermutationAndObservation

Position

a positive integer, the position in the list where the information will be extracted.

by

a integer, the increment of the number of permutations where the significant level is tested. Default: 100.

Value

0 indicating that all parameters validations have been successful.

Author(s)

Astrid Deschenes, Pascal Belleau

Examples

```r
## Get the name of the directory where files are stored
filesDir <- system.file("extdata", "TEST", package="methylInheritance")

## Merge permutation and observation results
methylInheritance:::validateLoadConvergenceData(analysisResultsDir = filesDir, permutationResults = filesDir, position = 1, by = 1)

## The function raises an error when at least one parameter is not valid
## Not run: methylInheritance:::validateLoadConvergenceData(
##     analysisResultsDir = filesDir, permutationResults = filesDir,
##     position = "hello", by = 1))
## End(Not run)
```

---

**validateMergePermutationAndObservation**

Validation of some parameters of the `mergePermutationAndObservation` function

### Description

Validation of some parameters needed by the public `mergePermutationAndObservation` function.

### Usage

```r
validateMergePermutationAndObservation(permutationResults, observationResults)
```

### Arguments

- **permutationResults**
  
a list with 1 entry called PERMUTATION. The PERMUTATION entry is a list with a number of entries corresponding to the number of permutations that have been processed. Each entry contains the result of one permutation.
observationResults

a list with 1 entry called OBSERVATION. The OBSERVATION entry is a list containing the result obtained with the observed dataset (not shuffled).

Value

0 indicating that all parameters validations have been successful.

Author(s)

Astrid Deschenes

Examples

```r
## Create a observation result
observed <- list()
observed[['OBSERVATION']] <- list()
observed[['OBSERVATION']]$SITES[['i2']] <- list(HYPER = list(11, 10), HYPO = list(13, 12))
observed[['OBSERVATION']]$SITES[['iAll']] <- list(HYPER = list(1), HYPO = list(3))

## Create a permutation result containing only 1 permutation result
## Real permutations results would have more entries
permutated <- list()
permutated[['PERMUTATION']] <- list()
permutated[['PERMUTATION']][[1]]$SITES[['i2']] <- list(HYPER = list(11, 12), HYPO = list(8, 11))
permutated[['PERMUTATION']][[1]]$SITES[['iAll']] <- list(HYPER = list(1), HYPO = list(3))

## Merge permutation and observation results
methylInheritance:::validateMergePermutationAndObservation(
  permutationResults = permutated, observationResults = observed)
```

## The function raises an error when at least one paramater is not valid
## Not run: methylInheritance:::validateMergePermutationAndObservation(
##  permutationResults = permutated, observationResults = NULL)
## End(Not run)

---

**validateRunObservation**

*Validation of some parameters of the runObservation function*

**Description**

Validation of some parameters needed by the public `runObservation` function.
validateRunObservation

Usage

validateRunObservation(
  methylKitData,
  type,
  outputDir,
  nbrCoresDiffMeth,
  minReads,
  minMethDiff,
  qvalue,
  maxPercReads,
  destrand,
  minCovBasesForTiles,
  tileSize,
  stepSize,
  vSeed,
  restartCalculation,
  saveInfoByGeneration
)

Arguments

methylKitData  a list of methylRawList entries or the name of the RDS file containing the list. Each methylRawList contains all the methylRaw entries related to one generation (first entry = first generation, second entry = second generation, etc.). The number of generations must correspond to the number of entries in the methylKitData. At least 2 generations must be present to calculate the conserved elements. More information can be found in the methylKit package.

type  One of the "sites","tiles" or "both" strings. Specifies the type of differentially methylated elements should be returned. For retrieving differentially methylated bases type="sites"; for differentially methylated regions type="tiles". Default: "both".

outputDir  a string, the name of the directory that will contain the results of the permutation. If the directory does not exist, it will be created.

nbrCoresDiffMeth  a positive integer, the number of cores to use for parallel differential methylation calculations. Parameter used for both sites and tiles analysis. The parameter corresponds to the num.cores parameter in the methylKit package.

minReads  a positive integer Bases and regions having lower coverage than this count are discarded. The parameter correspond to the lo.count parameter in the methylKit package.

minMethDiff  a positive double between [0,100], the absolute value of methylation percentage change between cases and controls. The parameter correspond to the difference parameter in the methylKit package.

qvalue  a positive double between [0,1], the cutoff for qvalue of differential methylation statistic.
validateRunObservation

maxPercReads  a double between [0,100], the percentile of read counts that is going to be used as upper cutoff. Bases ore regions having higher coverage than this percentile are discarded. Parameter used for both CpG sites and tiles analysis. The parameter correspond to the hi.perc parameter in the methylKit package.

destrand  a logical, when TRUE will merge reads on both strands of a CpG dinucleotide to provide better coverage. Only advised when looking at CpG methylation. Parameter used for both CpG sites and tiles analysis.

minCovBasesForTiles  a non-negative integer, the minimum number of bases to be covered in a given tiling window. The parameter corresponds to the cov.bases parameter in the package methylKit. Only used when doingTiles = TRUE. Default: 0.

tileSize  a positive integer, the size of the tiling window. The parameter corresponds to the win.size parameter in the methylKit package. Only used when doingTiles = TRUE.

stepSize  a positive integer, the step size of tiling windows. The parameter corresponds to the stepSize parameter in the methylKit package. Only used when doingTiles = TRUE.

vSeed  a integer, a seed used when reproducible results are needed. When a value inferior or equal to zero is given, a random integer is used.

restartCalculation  a logical, when TRUE, only permutations that don’t have an associated RDS result file are run. Useful to restart a permutation analysis that has been interrupted.

saveInfoByGeneration  a logical, when TRUE, the information about differentially methylated sites and tiles for each generation is saved in a RDS file. The information is saved in a different file for each permutation. The files are only saved when the outputDir is not NULL.

Value

0 indicating that all parameters validations have been successful.

Author(s)

Astrid Deschenes

Examples

```r
## Load dataset
data(samplesForTransgenerationalAnalysis)

## The function returns 0 when all parameters are valid
methylInheritance:::validateRunObservation(  
  methylKitData = samplesForTransgenerationalAnalysis, type = "sites",  
  outputDir = "test", nbrCoresDiffMeth = 1, minReads = 10,  
  minMethDiff = 25, qvalue = 0.01,  
  maxPercReads = 99.9, destrand = TRUE, minCovBasesForTiles = 10,
```
validateRunPermutation

Parameters validation for the runPermutation function

Description

Validation of all parameters needed by the public runPermutation function.

Usage

validateRunPermutation(
  methylKitData,
  type,
  outputDir,
  runObservedAnalysis,
  nbrPermutations,
  nbrCores,
  nbrCoresDiffMeth,
  minReads,
  minMethDiff,
  qvalue,
  maxPercReads,
  destrand,
  minCovBasesForTiles,
  tileSize,
  stepSize,
  vSeed,
  restartCalculation,
  saveInfoByGeneration
)
Arguments

methylKitData  a list of methylRawList entries or the name of the RDS file containing the
list. Each methylRawList entry must contain all the methylRaw entries rel-
ted to one generation (first entry = first generation, second entry = second
generation, etc.). The number of generations must correspond to the number of
entries in the methylKitData. At least 2 generations must be present to do a
permutation analysis. More information can be found in the methylKit package.

type  One of the "sites","tiles" or "both" strings. Specifies the type of differentially
methylated elements should be returned. For retrieving differentially methylated
bases type="sites"; for differentially methylated regions type="tiles". Default:
"both".

outputDir  a string, the name of the directory that will contain the results of the permutation.
If the directory does not exist, it will be created.

runObservedAnalysis  a logical, when runObservedAnalysis = TRUE, a CpG analysis on the ob-
served dataset is done.

nbrPermutations  a positive integer, the total number of permutations that is going to be done.

nbrCores  a positive integer, the number of cores to use when processing the analysis.

nbrCoresDiffMeth  a positive integer, the number of cores to use for parallel differential methyla-
tion calculations. Parameter used for both sites and tiles analysis. The parameter
 corresponds to the num.cores parameter in the methylKit package.

minReads  a positive integer. Bases and regions having lower coverage than this count
are discarded. The parameter corresponds to the lo.count parameter in the
methylKit package.

minMethDiff  a positive double between [0,100], the absolute value of methylation percent-
age change between cases and controls. The parameter corresponds to the
difference parameter in the methylKit package.

qvalue  a positive double between [0,1], the cutoff for qvalue of differential methyla-
tion statistic. TODO

maxPercReads  a double between [0,100], the percentile of read counts that is going to be used
as upper cutoff. Bases or regions having higher coverage than this percentile
are discarded. Parameter used for both CpG sites and tiles analysis. The param-
eter correspond to the hi.perc parameter in the methylKit package.

destrand  a logical, when TRUE will merge reads on both strands of a CpG dinucleotide
 to provide better coverage. Only advised when looking at CpG methylation. Parameter used for both CpG sites and tiles analysis.

minCovBasesForTiles  a non-negative integer, the minimum number of bases to be covered in a given
 tiling window. The parameter corresponds to the cov.bases parameter in the
 package methylKit. Only used when doingTiles = TRUE. Default: 0.

tileSize  a positive integer, the size of the tiling window. The parameter corresponds to
the win.size parameter in the methylKit package. Only used when doingTiles
 = TRUE.
**validateRunPermutation**

- **stepSize**: A positive integer, the step size of tiling windows. The parameter corresponds to the `stepSize` parameter in the `methylKit` package. Only used when `doingTiles = TRUE`.
- **vSeed**: An integer, a seed used when reproducible results are needed. When a value inferior or equal to zero is given, a random integer is used.
- **restartCalculation**: A logical, when `TRUE`, only permutations that don’t have an associated RDS result file are run. Useful to restart a permutation analysis that has been interrupted.
- **saveInfoByGeneration**: A logical, when `TRUE`, the information about differentially methylated sites and tiles for each generation is saved in a RDS file. The information is saved in a different file for each permutation. The files are only saved when the `outputDir` is not NULL.

**Value**

0 indicating that all parameters validations have been successful.

**Author(s)**

Astrid Deschenes

**Examples**

```r
## Load dataset
data(samplesForTransgenerationalAnalysis)

## The function returns 0 when all parameters are valid
methylInheritance:::validateRunPermutation(
  methylKitData = samplesForTransgenerationalAnalysis, type = "sites",
  outputDir = "test", runObservedAnalysis = TRUE,
  nbrPermutations = 10000, nbrCores = 1,
  nbrCoresDiffMeth = 1, minReads = 10, minMethDiff = 25, qvalue = 0.01,
  maxPercReads = 99.9, destrand = TRUE, minCovBasesForTiles = 10,
  tileSize = 1000, stepSize = 500, vSeed = 12, restartCalculation = FALSE,
  saveInfoByGeneration = FALSE)

## The function raises an error when at least one parameter is not valid
## Not run: methylInheritance:::validateRunPermutation(
## methylKitData = "HI", type = "tiles", outputDir = "test",
## runObservedAnalysis = FALSE, nbrPermutations = 10000, nbrCores = 1,
## nbrCoresDiffMeth = 1, minReads = 10, minMethDiff = 25, qvalue = 0.01,
## maxPercReads = 99.9, destrand = TRUE, minCovBasesForTiles = 10,
## tileSize = 1000, stepSize = 500, vSeed = 12, restartCalculation = FALSE,
## saveInfoByGeneration = FALSE)
```

## End(Not run)
Index

* datasets
  - demoForTransgenerationalAnalysis, 6
  - methylInheritanceResults, 18
  - samplesForTransgenerationalAnalysis, 35

* internal
  - createDataStructure, 4
  - createOutputDir, 5
  - formatInputMethylData, 9
  - getGRangesFromMethylDiff, 10
  - interGeneration, 11
  - isInterGenerationResults, 12
  - readInterGenerationResults, 25
  - runOnePermutationOnAllGenerations, 28
  - saveInterGenerationResults, 36
  - validateExtractInfo, 37
  - validateLoadConvergenceData, 38
  - validateMergePermutationAndObservation, 39
  - validateRunObservation, 40
  - validateRunPermutation, 43

* package
  - methylInheritance-package, 2

  - calculateSignificantLevel, 3
  - createDataStructure, 4
  - createOutputDir, 5

  - demoForTransgenerationalAnalysis, 6

  - extractInfo, 8, 22, 37

  - formatInputMethylData, 9

  - getGRangesFromMethylDiff, 10

  - interGeneration, 11
  - isInterGenerationResults, 12

  - loadAllRDSResults, 13
  - loadConvergenceData, 14, 38

  - mergePermutationAndObservation, 14, 15, 28, 34, 39

  - methylInheritance
    - (methylInheritance-package), 2

  - methylInheritance-package, 2

  - methylInheritanceResults, 18

  - plotConvergenceGraph, 22

  - plotGraph, 23

  - print.methylInheritanceAllResults, 24

  - readInterGenerationResults, 25

  - runObservation, 3, 7, 26, 40

  - runOnePermutationOnAllGenerations, 28

  - runPermutation, 3, 7, 31, 35, 43

  - samplesForTransgenerationalAnalysis, 35

  - saveInterGenerationResults, 36

  - validateExtractInfo, 37

  - validateLoadConvergenceData, 38

  - validateMergePermutationAndObservation, 39

  - validateRunObservation, 40

  - validateRunPermutation, 43