Package ‘HiCDOC’

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Description HiCDOC normalizes intrachromosomal Hi-C matrices, uses unsupervised learning to predict A/B compartments from multiple replicates, and detects significant compartment changes between experiment conditions. It provides a collection of functions assembled into a pipeline to filter and normalize the data, predict the compartments and visualize the results. It accepts several type of data: tabular `.tsv` files, Cooler `.cool` or `.mcool` files, Juicer `.hic` files or HiC-Pro `.matrix` and `.bed` files.

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**HiCDOC-package**  

**A/B compartment detection and differential analysis**

**Description**

HiCDOC normalizes intrachromosomal Hi-C matrices, uses unsupervised learning to predict A/B compartments from multiple replicates, and detects significant compartment changes between experiment conditions. It provides a collection of functions assembled into a pipeline to filter and normalize the data, predict the compartments and visualize the results. It accepts several type of data: tabular `.tsv` files, Cooler `.cool` or `.mcool` files, Juicer `.hic` files or HiC-Pro `.matrix` and `.bed` files.

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**See Also**

Useful links:  
- [https://github.com/mzytnicki/HiCDOC](https://github.com/mzytnicki/HiCDOC)  
- Report bugs at [https://github.com/mzytnicki/HiCDOC/issues](https://github.com/mzytnicki/HiCDOC/issues)

**detectCompartments**  

*A and B compartments detection and differences across conditions.*

**Description**

Detects compartments for each genomic position in each condition, and computes p-values for compartment differences between conditions.

**Usage**

```r
detectCompartments(  
  object,  
  parallel = FALSE,  
  kMeansDelta = NULL,  
  kMeansIterations = NULL,  
  kMeansRestarts = NULL,  
  PC1CheckThreshold = NULL  
)
```
**Arguments**

- `object`  A `HiCDOCDataSet`.
- `parallel`  Whether or not to parallelize the processing. Defaults to FALSE See ‘Details’.
- `kMeansDelta`  The convergence stop criterion for the clustering. When the centroids’ distances between two iterations is lower than this value, the clustering stops. Defaults to `object$kMeansDelta` which is originally set to `defaultHiCDOCParameters$kMeansDelta` = 0.0001.

- `kMeansIterations`  The maximum number of iterations during clustering. Defaults to `object$kMeansIterations` which is originally set to `defaultHiCDOCParameters$kMeansIterations` = 50.

- `kMeansRestarts`  The amount of times the clustering is restarted. For each restart, the clustering iterates until convergence or reaching the maximum number of iterations. The clustering that minimizes inner-cluster variance is selected. Defaults to `object$kMeansRestarts` which is originally set to `defaultHiCDOCParameters$kMeansRestarts` = 20.

- `PC1CheckThreshold`  The minimum percentage of variance that should be explained by the first principal component of centroids to pass sanity check. Defaults to `object$PC1CheckThreshold` which is originally set to `defaultHiCDOCParameters$PC1CheckThreshold` = 0.75

**Details**

**Genomic positions clustering:** To clusterize genomic positions, the algorithm follows these steps:

1. For each chromosome and condition, get the interaction vectors of each genomic position. Each genomic position can have multiple interaction vectors, corresponding to the multiple replicates in that condition.
2. For each chromosome and condition, use constrained K-means to clusterize the interaction vectors, forcing replicate interaction vectors into the same cluster. The euclidean distance between interaction vectors determines their similarity.
3. For each interaction vector, compute its concordance, which is the confidence in its assigned cluster. Mathematically, it is the log ratio of its distance to each centroid, normalized by the distance between both centroids, and min-maxed to a [-1,1] interval.
4. For each chromosome, compute the distance between all centroids and the centroids of the first condition. The cross-condition clusters whose centroids are closest are given the same cluster label. This results in two clusters per chromosome, spanning all conditions.

**A/B compartments prediction:** To match each cluster with an A or B compartment, the algorithm follows these steps:

1. For each genomic position, compute its self interaction ratio, which is the difference between its self interaction and the median of its other interactions.
2. For each chromosome, for each cluster, get the median self interaction ratio of the genomic positions in that cluster.
3. For each chromosome, the cluster with the smallest median self interaction ratio is matched with compartment A, and the cluster with the greatest median self interaction ratio is matched with compartment B. Compartment A being open, there are more overall interactions between distant genomic positions, so it is assumed that the difference between self interactions and other interactions is lower than in compartment B.

**Significant differences detection:** To find significant compartment differences across conditions, and compute their p-values, the algorithm follows three steps:

1. For each pair of replicates in different conditions, for each genomic position, compute the absolute difference between its concordances.
2. For each pair of conditions, for each genomic position, compute the median of its concordance differences.
3. For each pair of conditions, for each genomic position whose assigned compartment switches, rank its median against the empirical cumulative distribution of medians of all non-switching positions in that condition pair. Adjust the resulting p-value with the Benjamini–Hochberg procedure.

**Parallel processing:** The parallel version of detectCompartments uses the `bpmapply` function. Before to call the function in parallel you should specify the parallel parameters such as:

- On Linux:
  ```r
  multiParam <- BiocParallel::MulticoreParam(workers = 10)
  ```
- On Windows:
  ```r
  multiParam <- BiocParallel::SnowParam(workers = 10)
  ```

And then you can register the parameters to be used by BiocParallel:
```r
BiocParallel::register(multiParam, default = TRUE)
```

You should be aware that using MulticoreParam, reproducibility of the detectCompartments function using a RNGseed may not work. See this issue for more details.

**Value**

A `HiCDOCDataSet`, with compartments, concordances, distances, centroids, and differences.

**Examples**

```r
data(exampleHiCDOCDataSet)
## Run all filtering and normalization steps (not run for timing reasons)
# object <- filterSmallChromosomes(exampleHiCDOCDataSet)
# object <- filterSparseReplicates(object)
# object <- filterWeakPositions(object)
# object <- normalizeTechnicalBiases(object)
# object <- normalizeBiologicalBiases(object)
# object <- normalizeDistanceEffect(object)

# Detect compartments and differences across conditions
object <- detectCompartments(exampleHiCDOCDataSet)
```
Example HiCDOCDataSet

Description
A S4 HiCDOCDataSet object with 4 chromosomes, 3 conditions and 3 replicates.

Usage
data(exampleHiCDOCDataSet)

Format
S4 HiCDOCDataSet object with the following characteristics:
- **chromosomes**: 4 chromosomes: W, X, Y, Z
- **conditions**: 3 conditions: 1, 2, 3
- **replicates**: 3 replicates: R1, R2, R3
- **binSize**: A resolution of 137 bases

Value
A HiCDOCDataSet.

Examples
  data(exampleHiCDOCDataSet)
  exampleHiCDOCDataSet

Example HiCDOCDataSet Processed

Example HiCDOCDataSet, filtered, normalized and with compartments detected.

Description
A S4 HiCDOCDataSet object with 3 chromosomes, 3 conditions and 3 replicates. Can be retrieved by running:

```r
data(exampleHiCDOCDataSet)
set.seed(123)
exampleHiCDOCDataSetProcessed <- HiDOC(exampleHiCDOCDataSet)
```

Usage
data(exampleHiCDOCDataSetProcessed)
filterSmallChromosomes

Format

S4 HiCDOCDataSet object with the following characteristics:

- **chromosomes**: 4 chromosomes: X, Y, Z
- **conditions**: 3 conditions: 1, 2, 3
- **replicates**: 3 replicates: R1, R2, R3
- **binSize**: A resolution of 137 bases

Value

A HiCDOCDataSet, already filtered and normalized.

Examples

data(exampleHiCDOCDataSetProcessed)
exampleHiCDOCDataSetProcessed

filterSmallChromosomes

*Filter small chromosomes.*

Description

Removes chromosomes whose length (in number of positions) is smaller than the threshold.

Usage

filterSmallChromosomes(object, threshold = NULL)

Arguments

- **object**: A HiCDOCDataSet.
- **threshold**: The minimum length (number of positions) for a chromosome to be kept. Defaults to `object$smallChromosomeThreshold` which is originally set to `defaultHiCDOCParameters$smallChromosomeThreshold` = 100.

Value

A filtered HiCDOCDataSet.

See Also

filterSparseReplicates, filterWeakPositions, HiCDOC
Examples

```r
data(exampleHiCDOCDataSet)
object <- exampleHiCDOCDataSet

chromosomes(object)
object <- filterSmallChromosomes(object)
chromosomes(object)
```

---

filterSparseReplicates

`Filter sparse replicates.`

Description

Removes chromosome replicates whose percentage of non-zero interactions is smaller than the threshold.

Usage

```r
filterSparseReplicates(object, threshold = NULL)
```

Arguments

- **object**: A `HiCDOCDataSet`.
- **threshold**: The minimum percentage of non-zero interactions for a chromosome replicate to be kept. If a chromosome replicate’s percentage of non-zero interactions is lower than this value, it is removed. Defaults to `object$smallChromosomeThreshold` which is originally set to `defaultHiCDOCParameters$smallChromosomeThreshold = 30%`.

Value

A filtered `HiCDOCDataSet`.

See Also

`filterSmallChromosomes`, `filterWeakPositions`, `HiCDOC`

Examples

```r
data(exampleHiCDOCDataSet)
object <- exampleHiCDOCDataSet

object <- filterSparseReplicates(object)
```
filterWeakPositions  Filter weak positions.

Description
Removes weak genomic positions whose interactions average is lower than the threshold.

Usage
filterWeakPositions(object, threshold = NULL)

Arguments
- object: A HiCDOCDataSet.
- threshold: The minimum average interaction for a position to be kept. If a position’s average interaction with the entire chromosome is lower than this value in any of the replicates, it is removed from all replicates and conditions. Defaults to `object$smallChromosomeThreshold` which is originally set to `defaultHiCDOCParameters$smallChromosomeThreshold = 1`.

Details
Detects weak genomic positions in each replicate, and removes them from all replicates to guarantee comparability across conditions when calling `detectCompartments`.

Value
A filtered HiCDOCDataSet.

See Also
- `filterSmallChromosomes`, `filterSparseReplicates`, `HiCDOC`

Examples
```r
data(exampleHiCDOCDataSet)
object <- exampleHiCDOCDataSet
object <- filterWeakPositions(object)
```
HiCDOC

Default pipeline to run on the HiCDOC analysis.

Description

Runs the default filtering, normalization, and computational steps on a HiCDOCDataSet. To learn more about HiCDOC, browse the vignette: browseVignettes(package = "HiCDOC").

Usage

HiCDOC(object, parallel = FALSE)

Arguments

object A HiCDOCDataSet.
parallel Whether or not to parallelize each step. Defaults to FALSE.

Details

HiCDOC pipeline: The HiCDOC pipeline has seven steps:

Three filtering steps: • filterSmallChromosomes to filter out small chromosomes
• filterWeakPositions to filter out weak positions with very few interactions
• filterSparseReplicates to filter out sparse replicates with many null interactions

Three normalization steps: • normalizeTechnicalBiases to normalize technical biases in each replicate
• normalizeBiologicalBiases to normalize biological biases in each replicate
• normalizeDistanceEffect to normalize the distance effect in each chromosome

One computational step: • detectCompartments to detect compartments in each condition and find significant changes between conditions.

Parallel processing: The parallel version of HiCDOC uses the BiocParallel package. Before to call the function in parallel you should specify the parallel parameters such as:

• On Linux:
  multiParam <- BiocParallel::MulticoreParam(workers = 10)
• On Windows:
  multiParam <- BiocParallel::SnowParam(workers = 10)

And then you can register the parameters to be used by BiocParallel:
BiocParallel::register(multiParam, default = TRUE)

You should be aware that using MulticoreParam, reproducibility of the detectCompartments function using a RNGseed may not work. See this issue for more details.

Value

A HiCDOCDataSet with all slots filled.
HiCDOCDataSet-class

See Also

HiCDOCDataSet, filterSmallChromosomes, filterWeakPositions, filterSparseReplicates,
normalizeTechnicalBiases, normalizeBiologicalBiases, normalizeDistanceEffect, detectCompartments

Examples

data(exampleHiCDOCDataSet)
# Default HiCDOC pipeline
# Not printing loess warnings for example purpose.
# Results should be inspected if there is any.
suppressWarnings(
  object <- HiCDOC(exampleHiCDOCDataSet)
)

# Equivalent to
if(FALSE){
  object <- filterSmallChromosomes(exampleHiCDOCDataSet)
  object <- filterSparseReplicates(object)
  object <- filterWeakPositions(object)
  object <- normalizeTechnicalBiases(object)
  object <- normalizeBiologicalBiases(object)
  object <- normalizeDistanceEffect(object)
  object <- detectCompartments(object)
}

HiCDOCDataSet-class

HiCDOCDataSet S4 class.

Description

Data structure for a Hi-C experiment.

Details

An instance of HiCDOCDataSet describes a Hi-C experiment with slots for path(s) to input file(s), interactions, pipeline parameters defaulting to defaultHiCDOCParameters, and computation results. It can be constructed from 4 different types of data:

- Tabular files: see HiCDOCDataSetFromTabular
- (m)Cool files: see HiCDOCDataSetFromCool
- HiC files: see HiCDOCDataSetFromHiC
- HiC-Pro matrices and bed files: see HiCDOCDataSetFromHiCPro An example HiCDOCDataSet is also available, see exampleHiCDOCDataSet. The HiCDOCDataSet object can be explored using the appropriate accessors.
**Accessors**

The accessors for a HiCDOCDataset object are the following:

- **chromosomes** to retrieve the vector of chromosome names.
- **sampleConditions** to retrieve the vector of condition names, one for each sample.
- **sampleReplicates** to retrieve the vector of replicate names, one for each sample.

After the detection of compartments you can use this accessors:

- **compartments** returns a GenomicRange of the compartment of every position in every condition.
- **concordances** returns a GenomicRange of the significant compartment differences between conditions, and their p-values.
- **differences** returns a GenomicRange of the concordance (confidence in assigned compartment) of every position in every replicate.

See the HiCDOCDataset-methods man page for more details on methods and accessors.

**See Also**

HiCDOC, exampleHiCDOCDataset, HiCDOCDatasetFromTabular, HiCDOCDatasetFromCool, HiCDOCDatasetFromHiC, HiCDOCDatasetFromHiCPro

---

**HiCDOCDataset-methods**  
*Methods to access a HiCDOCDataset components.*

---

**Description**

Retrieve information and results from a HiCDOCDataset.

**Usage**

```r
## S4 method for signature 'HiCDOCDataset'
chromosomes(object)

## S4 method for signature 'HiCDOCDataset'
sampleConditions(object)

## S4 method for signature 'HiCDOCDataset'
sampleReplicates(object)

## S4 method for signature 'HiCDOCDataset'
compartments(object, passChecks = TRUE)

## S4 method for signature 'HiCDOCDataset'
differences(object, threshold = NULL)
```
## S4 method for signature 'HiCDOCDataSet'
concordances(object, passChecks = TRUE)

## S4 method for signature 'HiCDOCDataSet'
show(object)

### Arguments

- **object**: a HiCDOCDataSet object
- **passChecks**: logical. Display only the concordances/compartments for the chromosomes passing sanity checks.
- **threshold**: a numeric value between 0 and 1. If no threshold, all the differences will be printed even the non significant ones. Otherwise the differences printed are filtered to show the ones with an adjusted p-value <= threshold.

### Value

A character vector (for chromosomes, sampleConditions, sampleReplicates), or a GRanges object (for compartments, concordances, differences).

### Functions

- **chromosomes()**: Retrieves the vector of chromosome names.
- **sampleConditions()**: Retrieves the vector of condition names, one for each sample.
- **sampleReplicates()**: Retrieves the vector of replicate names, one for each sample.
- **compartments()**: Retrieves a GenomicRange of the compartment of every position in every condition.
- **differences()**: Retrieves a GenomicRange of the significant compartment differences between conditions, and their p-values.
- **concordances()**: Retrieves a GenomicRange of the concordance (confidence in assigned compartment) of every position in every replicate.

### Examples

```r
# Load an example dataset already processed
# (i.e. after the detection of compartments)
data(exampleHiCDOCDataSetProcessed)
exa..."
HiCDOCDataSet-parameters

Access the parameters of a HiCDOCDataSet.

Description
Retrieves or sets parameters used for filtering, normalization, and prediction of compartments.

Usage

```r
defaultHiCDOCParameters

## S4 method for signature 'HiCDOCDataSet'
parameters(object)

## S4 replacement method for signature 'HiCDOCDataSet'
parameters(object) <- value
```

Arguments

- `object` A `HiCDOCDataSet`.
- `value` a named list containing the names and values of the parameters to change (see Details).

Format

An object of class `list` of length 9.

Details

A `HiCDOCDataSet`'s parameters are automatically set to default values retrieved from `defaultHiCDOCParameters`. They are accessed by filtering, normalization, and compartment detection functions. If those functions are called with custom arguments, the object's parameters are updated to record the actual parameters used. If the object's parameters are customized before calling the functions, the custom parameters will be used.

See `filterSmallChromosomes`, `filterSparseReplicates`, `filterWeakPositions`, `normalizeDistanceEffect`, and `detectCompartments`, for details on how these parameters are used.

All parameters are listed here:

- `smallChromosomeThreshold` The minimum length (number of positions) for a chromosome to be kept when filtering with `filterSmallChromosomes`. Defaults to `defaultHiCDOCParameters$smallChromosomeThreshold = 100`.
- `sparseReplicateThreshold` The minimum percentage of non-zero interactions for a chromosome replicate to be kept when filtering with `filterSparseReplicates`. If a chromosome replicate's percentage of non-zero interactions is lower than this value, it is removed. Defaults to `defaultHiCDOCParameters$smallChromosomeThreshold = 30`.
weakPositionThreshold  The minimum average interaction for a position to be kept when filtering with `filterWeakPositions`. If a position’s average interaction with the entire chromosome is lower than this value in any of the replicates, it is removed from all replicates and conditions. Defaults to `defaultHiCDOCParameters$smallChromosomeThreshold = 1`.

cyclicLoessSpan  The span for cyclic loess normalization used in `normalizeTechnicalBiases`. This value is passed to `multiHiCcompare::cyclic_loess`. Defaults to NA indicating that span will be automatically calculated using generalized cross validation. For large dataset, it is highly recommended to set this value to reduce computing time and necessary memory.

loessSampleSize  The number of positions used as a sample to estimate the effect of distance on proportion of interactions when normalizing with `normalizeDistanceEffect`. Defaults to `defaultHiCDOCParameters$loessSampleSize = 20000`.

kMeansDelta  The convergence stop criterion for the clustering when detecting compartments with `detectCompartments`. When the centroids’ distances between two iterations is lower than this value, the clustering stops. Defaults to `defaultHiCDOCParameters$kMeansDelta = 0.0001`.

kMeansIterations  The maximum number of iterations during clustering when detecting compartments with `detectCompartments`. Defaults to `defaultHiCDOCParameters$kMeansIterations = 50`.

kMeansRestarts  The amount of times the clustering is restarted when detecting compartments with `detectCompartments`. For each restart, the clustering iterates until convergence or reaching the maximum number of iterations. The clustering that minimizes inner-cluster variance is selected. Defaults to `defaultHiCDOCParameters$kMeansRestarts = 20`.

PC1CheckThreshold  The minimum percentage of variance that should be explained by the first principal component of centroids to pass sanity check. Defaults to `defaultHiCDOCParameters$PC1CheckThreshold = 0.75`.

Examples

```r
data(exampleHiCDOCDataSet)

# Retrieve parameters
parameters(exampleHiCDOCDataSet)

# Set parameters
parameters(exampleHiCDOCDataSet) <- list("smallChromosomeThreshold" = 50)
parameters(exampleHiCDOCDataSet) <- list(
  "weakPositionThreshold" = 10,
  "kMeansRestarts" = 30
)
```

HiCDOCDataSetFromCool  The `HiCDOCDataSet` constructor from Cool files.

Description

Constructs a `HiCDOCDataSet` from a set of .cool or .mcool files.
HiCDOCDataSetFromCool(paths, replicates, conditions, binSize = NA)

Arguments

paths A vector of paths to .cool or .mcool files.
replicates A vector of replicate names repeated along the conditions.
conditions A vector of condition names repeated along the replicates.
binSize The resolution (span of each position in number of bases). Optionally provided to select the appropriate resolution in .mcool files. Defaults to NULL.

Value

A HiCDOCDataSet.

Examples

## Not run:
# Path to each file
paths = c(
  'path/to/condition-1.replicate-1.cool',
  'path/to/condition-1.replicate-2.cool',
  'path/to/condition-2.replicate-1.cool',
  'path/to/condition-2.replicate-2.cool',
  'path/to/condition-3.replicate-1.cool'
)

# Replicate and condition of each file. Can be names instead of numbers.
replicates <- c(1, 2, 1, 2, 1)
conditions <- c(1, 1, 2, 2, 3)

# Resolution to select in .mcool files
binSize = 500000

# Instantiation of data set
object <- HiCDOCDataSetFromCool(
  paths,
  replicates = replicates,
  conditions = conditions,
  binSize = binSize # Specified for .mcool files.
)

## End(Not run)
HiCDOCDataSetFromHiC

HiCDOCDataSet constructor from HiC files.

Description

Constructs a HiCDOCDataSet from a set of .hic files.

Usage

HiCDOCDataSetFromHiC(paths, replicates, conditions, binSize)

Arguments

paths A vector of paths to .hic files.
replicates A vector of replicate names repeated along the conditions.
conditions A vector of condition names repeated along the replicates.
binSize The resolution (span of each position in number of bases) to select within the .hic files.

Value

A HiCDOCDataSet.

Examples

## Not run:

```r
# Path to each file
paths = c(
  'path/to/condition-1.replicate-1.hic',
  'path/to/condition-1.replicate-2.hic',
  'path/to/condition-2.replicate-1.hic',
  'path/to/condition-2.replicate-2.hic',
  'path/to/condition-3.replicate-1.hic'
)

# Replicate and condition of each file. Can be names instead of numbers.
replicates <- c(1, 2, 1, 2, 1)
conditions <- c(1, 1, 2, 2, 3)

# Resolution to select
binSize <- 500000

# Instantiation of data set
hic.experiment <- HiCDOCDataSetFromHiC(paths,
  replicates = replicates,
  conditions = conditions,
  binSize = binSize
)```

HiCDOCDataSetFromHiCPro

**HiCDOCDataSet** constructor from HiC-Pro files.

---

### Description

Constructs a **HiCDOCDataSet** from a set of HiC-Pro generated files.

### Usage

```r
HiCDOCDataSetFromHiCPro(matrixPaths, bedPaths, replicates, conditions)
```

### Arguments

- **matrixPaths**: A vector of paths to HiC-Pro matrix files.
- **bedPaths**: A vector of paths to HiC-Pro bed files.
- **replicates**: A vector of replicate names repeated along the conditions.
- **conditions**: A vector of condition names repeated along the replicates.

### Value

A **HiCDOCDataSet**.

### Examples

```r
## Not run:
# Path to each matrix file
matrixPaths = c(
  'path/to/condition-1.replicate-1.matrix',
  'path/to/condition-1.replicate-2.matrix',
  'path/to/condition-2.replicate-1.matrix',
  'path/to/condition-2.replicate-2.matrix',
  'path/to/condition-3.replicate-1.matrix'
)

# Path to each bed file
bedPaths = c(
  'path/to/condition-1.replicate-1.bed',
  'path/to/condition-1.replicate-2.bed',
  'path/to/condition-2.replicate-1.bed',
  'path/to/condition-2.replicate-2.bed',
  'path/to/condition-3.replicate-1.bed'
)
```
# Replicate and condition of each file. Can be names instead of numbers.
replicates <- c(1, 2, 1, 2, 1)
conditions <- c(1, 1, 2, 2, 3)

# Instantiation of data set
hic.experiment <- HiCDOCDataSetFromHiCPro(
  matrixPaths = matrixPaths,
  bedPaths = bedPaths,
  replicates = replicates,
  conditions = conditions
)

## End(Not run)

HiCDOCDataSetFromTabular

**HiCDOCDataSet** constructor from a tabular file.

**Description**

Constructs a **HiCDOCDataSet** from a tabular file.

**Usage**

HiCDOCDataSetFromTabular(path, sep = '\t')

**Arguments**

- **path** A path to a tabular file.
- **sep** The separator of the tabular file. Default to tabulation.

**Details**

Accepts a tabular file with chromosome, position 1, position 2, and multiple replicate columns listing interaction counts. Null interactions do not have to be listed. Values must be separated by tabulations. The header must be chromosome position 1 position 2 x.y x.y x.y ... with x replaced by condition names and y replaced by replicate names.

**Value**

A **HiCDOCDataSet**.

**Examples**

```r
path <- system.file("extdata", "liver_18_10M_500000.tsv", package = "HiCDOC")
object <- HiCDOCDataSetFromTabular(path, sep = '\t')
```
normalizeBiologicalBiases

Normalize biological biases.

Description

Normalizes biological biases such as GC content and repeated regions. Uses the Knight-Ruiz balancing algorithm to transform interaction matrices into doubly stochastic matrices, with sum of each row and sum of each column equal to 1.

Usage

normalizeBiologicalBiases(object, parallel = FALSE)

Arguments

- **object**: A `HiCDOCDataSet`.
- **parallel**: Should the normalization be run in parallel mode? Default to FALSE.

Value

A `HiCDOCDataSet` with normalized interactions.

See Also

`filterSparseReplicates`, `filterWeakPositions`, `normalizeTechnicalBiases`, `normalizeDistanceEffect`, `HiCDOC`

Examples

```r
data(exampleHiCDOCDataSet)
object <- exampleHiCDOCDataSet
object <- filterSparseReplicates(object)
object <- filterWeakPositions(object)
object <- normalizeBiologicalBiases(object)
```

normalizeDistanceEffect

Normalize distance effect.

Description

Normalizes interactions by their "expected" value relative to the distance that separates their positions. The "expected" values are estimated with a loess regression on the proportion of interactions for each distance.
normalizeTechnicalBiases

Usage

normalizeDistanceEffect(object, loessSampleSize = NULL, parallel = FALSE)

Arguments

object A HiCDOCDataSet.
loessSampleSize The number of positions used as a sample to estimate the effect of distance on proportion of interactions. Defaults to object$loessSampleSize which is originally set to defaultHiCDOCParameters$loessSampleSize = 20000.
parallel Should the normalization be run in parallel mode? Default to FALSE.

Value

A HiCDOCDataSet with normalized interactions.

See Also

normalizeTechnicalBiases, normalizeBiologicalBiases, HiCDOC

Examples

data(exampleHiCDOCDataSet)
object <- normalizeDistanceEffect(exampleHiCDOCDataSet)

normalizeTechnicalBiases

Normalize technical biases.

Description

Normalizes technical biases such as sequencing depth by using a cyclic loess to recursively normalize each pair of interaction matrices. Depends on multiHiCcompare.

Usage

normalizeTechnicalBiases(object, parallel = FALSE, cyclicLoessSpan = NULL)

Arguments

object A HiCDOCDataSet.
parallel Logical. Whether or not to parallelize the processing. Defaults to FALSE
normalizeTechnicalBiases

cyclicLoessSpan
A numeric value in between 0 and 1. The span for cyclic loess normalization. This value is passed to multiHiCcompare::cyclic_loess. Defaults to NULL. NULL indicates that the value of parameters(object)$cyclicLoessSpan will be used. If this value is NA, the span will be automatically calculated using generalized cross validation. **For large dataset, it is highly recommended to set this value to reduce computing time and necessary memory.**

Details

**Parallel processing:** If parallel = TRUE, the function cyclic_loess is launched in parallel mode, using bplapply function. Before to call the function in parallel you should specify the parallel parameters such as:

- On Linux:
  ```
  multiParam <- BiocParallel::MulticoreParam(workers = 10)
  ```
- On Windows:
  ```
  multiParam <- BiocParallel::SnowParam(workers = 10)
  ```

And then you can register the parameters to be used by BiocParallel:
```
BiocParallel::register(multiParam, default = TRUE)
```

Value

A HiCDOCDataSet with normalized interactions.

See Also

filterSparseReplicates, filterWeakPositions, normalizeBiologicalBiases, normalizeDistanceEffect, HiDOC

Examples

```r
data(exampleHiCDOCDataSet)
object <- filterSmallChromosomes(exampleHiCDOCDataSet)
object <- filterSparseReplicates(object)
object <- filterWeakPositions(object)
# Not printing loess warnings for example purpose. # Results should be inspected if there is any.
suppressWarnings(
  object <- normalizeTechnicalBiases(object)
)
```
plotCentroids

Description
Plots the result of the PCA on the compartments’ centroids.

Usage
plotCentroids(object, chromosome, size = 2, checks = TRUE)

Arguments
- object: A HiCDOCDataSet.
- chromosome: A chromosome name or index in chromosomes(object).
- size: Size of each point. Defaults to 2.
- checks: Whether or not to add sanity checks messages on centroids. Default to TRUE.

Value
A ggplot.

Examples
data(exampleHiCDOCDataSetProcessed)
plotCentroids(exampleHiCDOCDataSetProcessed, chromosome = 1)

plotCompartmentChanges

Description
Plots the predicted compartments, along with their concordance in each replicate, and significant changes between experiment conditions.

Usage
plotCompartmentChanges(
  object,
  chromosome,
  threshold = 0.05,
  xlim = NULL,
  points = FALSE,
  checks = TRUE,
  colour = "gray90"
)


plotCompartments

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>object</td>
<td>A HiCDOCDataSet.</td>
</tr>
<tr>
<td>chromosome</td>
<td>A chromosome name or index in chromosomes(object).</td>
</tr>
<tr>
<td>threshold</td>
<td>Significance threshold for the compartment changes. Defaults to 0.05.</td>
</tr>
<tr>
<td>xlim</td>
<td>A vector of the minimum and maximum positions to display. If NULL, displays all positions. Defaults to NULL.</td>
</tr>
<tr>
<td>points</td>
<td>Whether or not to add points to the concordances. Defaults to FALSE.</td>
</tr>
<tr>
<td>checks</td>
<td>Whether or not to add sanity checks messages. Default to TRUE.</td>
</tr>
<tr>
<td>colour</td>
<td>Border color for the compartments. Default to ‘gray90’. ‘NA’ means no border.</td>
</tr>
</tbody>
</table>

Value

A ggplot.

Examples

data(exampleHiCDOCDataSetProcessed)
plotCompartmentChanges(exampleHiCDOCDataSetProcessed, chromosome = 1)

plotCompartments  Plot A/B compartments.

Description

Plots the predicted compartments in each experiment condition.

Usage

plotCompartments(object, chromosome, xlim = NULL, colour = "gray90")

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>object</td>
<td>A HiCDOCDataSet.</td>
</tr>
<tr>
<td>chromosome</td>
<td>A chromosome name or index in chromosomes(object).</td>
</tr>
<tr>
<td>xlim</td>
<td>A vector of the minimum and maximum positions to display. If NULL, displays all positions. Defaults to NULL.</td>
</tr>
<tr>
<td>colour</td>
<td>Border color for the compartments. Default to ‘gray90’. ‘NA’ means no border.</td>
</tr>
</tbody>
</table>

Value

A ggplot.

Examples

data(exampleHiCDOCDataSetProcessed)
plotCompartments(exampleHiCDOCDataSetProcessed, chromosome = 1)
plotConcordanceDifferences

*Plot the distribution of concordance differences.*

**Description**

Plots the distribution of concordance differences, which are the differences between concordances of each pair of replicates from different conditions. A concordance can be understood as a confidence in a genomic position’s assigned compartment. Mathematically, it is the log ratio of a genomic position’s distance to each compartment’s centroid, normalized by the distance between both centroids, and min-maxed to a [-1,1] interval.

**Usage**

plotConcordanceDifferences(object)

**Arguments**

object A HiCDOCDataSet.

**Value**

A ggplot.

**Examples**

data(exampleHiCDOCDataSetProcessed)
plotConcordanceDifferences(exampleHiCDOCDataSetProcessed)

---

plotConcordances *Plot concordances.*

**Description**

Plots the concordances of each replicate in each experiment condition. A concordance can be understood as a confidence in a genomic position’s assigned compartment. Mathematically, it is the log ratio of a genomic position’s distance to each compartment’s centroid, normalized by the distance between both centroids, and min-maxed to a [-1,1] interval.
plotConcordances(
    object,
    chromosome,
    xlim = NULL,
    threshold = 0.05,
    points = FALSE
)

Arguments

object A \texttt{HiCDOCDataSet}.
chromosome A chromosome name or index in \texttt{chromosomes(object)}.
xlim A vector of the minimum and maximum positions to display. If NULL, displays all positions. Defaults to NULL.
threshold Significance threshold for the compartment changes. Defaults to 0.05.
points Whether or not to add points to the concordances. Defaults to FALSE.

Value

A \texttt{ggplot}.

Examples

data(exampleHiCDOCDataSetProcessed)
plotConcordances(exampleHiCDOCDataSetProcessed, chromosome = 1)

plotDistanceEffect

\textit{Plot the distance effect.}

Description

Plots the distance effect on proportion of interactions. Each point is a cell in the interaction matrix, such that the x-axis is the distance with respect to the diagonal, the y-axis is number of counts. Dots are binned.

Usage

plotDistanceEffect(
    object,
    chromosome = NULL,
    transformX = "identity",
    transformY = "identity"
)
plotInteractions

Arguments

<table>
<thead>
<tr>
<th>argument</th>
<th>description</th>
</tr>
</thead>
<tbody>
<tr>
<td>object</td>
<td>A HiCDOCDataSet.</td>
</tr>
<tr>
<td>chromosome</td>
<td>Name (character) or index of the chromosome, if the plot should be restricted to only one chromosome. Default to NULL.</td>
</tr>
<tr>
<td>transformX</td>
<td>Transformation of the X axis. Default to &quot;identity&quot;. See scale_x_continuous for other accepted values.</td>
</tr>
<tr>
<td>transformY</td>
<td>Transformation of the Y axis. Default to &quot;identity&quot;. See scale_y_continuous for other accepted values.</td>
</tr>
</tbody>
</table>

Value

A ggplot.

Examples

data(exampleHiCDOCDataSet)
plotDistanceEffect(exampleHiCDOCDataSet)

Description

Plots the interaction matrices as heatmaps.

Usage

plotInteractions(
  object,
  chromosome,
  transform = NULL,
  colours = c(low = "#2c7bb6", mid = "#fffbfb", high = "#d7191c"),
  midpoint = 0
)

Arguments

<table>
<thead>
<tr>
<th>argument</th>
<th>description</th>
</tr>
</thead>
<tbody>
<tr>
<td>object</td>
<td>A HiCDOCDataSet.</td>
</tr>
<tr>
<td>chromosome</td>
<td>A chromosome name or index in chromosomes(object).</td>
</tr>
<tr>
<td>transform</td>
<td>Transformation of the color scale. Default to NULL (no transformation). See scale_fill_gradient2 for other accepted values.</td>
</tr>
<tr>
<td>colours</td>
<td>A character vector colours of length 3 to use for the gradient. See scale_fill_gradient2 for more info. Defaults to c(&quot;low&quot;=#2c7bb6&quot;, &quot;mid&quot;=#fffbfb&quot;, &quot;high&quot;=#d7191c&quot;).</td>
</tr>
<tr>
<td>midpoint</td>
<td>midpoint value to be passed to scale_fill_gradient2. Default to 0.</td>
</tr>
</tbody>
</table>
plotSelfInteractionRatios

Value
A ggplot.

Examples

```r
data(exampleHiCDOCDataSet)
plotInteractions(exampleHiCDOCDataSet, chromosome = 1)
```

---

```
plotSelfInteractionRatios

Plot boxplots of self interaction ratios.
```

Description

Plots the boxplots of self interaction ratios, which are the differences between self interaction and median of other interactions for each genomic position. Since the A compartment is open with more interactions overall, it is assumed that self interaction ratios in compartment A are smaller than in compartment B.

Usage

```r
plotSelfInteractionRatios(object, chromosome, checks = TRUE)
```

Arguments

- `object`: A `HiCDOCDataSet`.
- `chromosome`: A chromosome name or index in `chromosomes(object)`. A `HiCDOCDataSet`.
- `checks`: Logical. Should sanity checks messages be printed on plot? Default to TRUE.

Value

A ggplot.

Examples

```r
data(exampleHiCDOCDataSetProcessed)
plotSelfInteractionRatios(exampleHiCDOCDataSetProcessed, chromosome = 1)
```
**reduceHiCDOCDataSet**

Reduce a HiCDOCDataSet.

### Description

Reduces a HiCDOCDataSet by keeping only given chromosomes, conditions, or replicates.

### Usage

```r
reduceHiCDOCDataSet(
  object,
  chromosomes = NULL,
  conditions = NULL,
  replicates = NULL,
  dropLevels = TRUE
)
```

### Arguments

- **object**
  - A HiCDOCDataSet.
- **chromosomes**
  - The chromosome names or indices in chromosomes(object) to keep. Defaults to NULL.
- **conditions**
  - The condition names in sampleConditions(object) to keep. Defaults to NULL.
- **replicates**
  - The replicate names in sampleReplicates(object) to keep. Defaults to NULL.
- **dropLevels**
  - Whether or not to also remove unused factor levels after filtering. Should be set to FALSE if the reduced objects are meant to be re-combined later. Defaults to TRUE.

### Value

A reduced HiCDOCDataSet.

### Examples

```r
data(exampleHiCDOCDataSet)
reduced <- reduceHiCDOCDataSet(exampleHiCDOCDataSet, chromosomes = c(1, 2))
```
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