

Package: karyotapR (via r-universe)

September 9, 2024

Title DNA Copy Number Analysis for Genome-Wide Tapestri Panels

Version 1.0.1.9000

Description Analysis of DNA copy number in single cells using custom genome-wide targeted DNA sequencing panels for the Mission Bio Tapestri platform. Users can easily parse, manipulate, and visualize datasets produced from the automated 'Tapestri Pipeline', with support for normalization, clustering, and copy number calling. Functions are also available to deconvolute multiplexed samples by genotype and parsing barcoded reads from exogenous lentiviral constructs.

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Encoding UTF-8

Roxygen list(markdown = TRUE)

RoxygenNote 7.2.3

URL <https://github.com/joeymays/karyotapR>,
<http://joeymays.xyz/karyotapR/>

BugReports <https://github.com/joeymays/karyotapR/issues>

Imports circlize, cli, ComplexHeatmap, dbscan, dplyr, fitdistrplus, GenomicRanges, ggplot2, gtools, IRanges, magrittr, methods, purrr, rhdf5, rlang, S4Vectors, stats, SummarizedExperiment, tibble, tidyr, umap, viridisLite

Depends R (>= 3.6), SingleCellExperiment

Suggests Biostrings, knitr, rmarkdown, Rsamtools, testthat (>= 3.0.0)

Config/testthat/edition 3

Config/Needs/website forcats, purrr, ggdist

Repository <https://joeymays.r-universe.dev>

RemoteUrl <https://github.com/joeymays/karyotapR>

RemoteRef HEAD

RemoteSha f675f908c089d5c2828c5fdb0c0ba1ce58a509f9

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| | |
|--------------|--|
| assayBoxPlot | <i>Generate a box plot from assay data</i> |
|--------------|--|

Description

Draws box plot of data from indicated TapestriExperiment assay slot. This is especially useful for visualizing altExp count data, such as counts from probes on chrY or barcode probe counts.

Usage

```
assayBoxPlot(
  TapestriExperiment,
  alt.exp = NULL,
  assay = NULL,
  log.y = TRUE,
  split.features = FALSE,
  split.x.by = NULL,
  split.y.by = NULL
)
```

Arguments

| | |
|--------------------|---|
| TapestriExperiment | TapestriExperiment object |
| alt.exp | Character, altExp to plot. NULL (default) uses the top-level experiment in TapestriExperiment. |
| assay | Character, assay to plot. NULL (default) selects first assay listed TapestriExperiment. |
| log.y | Logical, if TRUE, scales data using <code>log1p()</code> . Default TRUE. |
| split.features | Logical, if TRUE, splits plot by rowData features if slot has more than one row feature/probe. Default FALSE. |
| split.x.by | Character, colData column to use for X-axis categories. Default NULL. |
| split.y.by | Character, colData column to use for Y-axis splitting/faceting. Default NULL. |

Value

ggplot object, box plot

See Also

[ggplot2::geom_boxplot\(\)](#)

Examples

```
tap.object <- newTapestriExperimentExample() # example TapestriExperiment object
assayBoxPlot(tap.object, alt.exp = "chrYCounts", split.features = TRUE, split.x.by = "test.cluster")
```

| | |
|--------------|---------------------------------------|
| assayHeatmap | <i>Generate heatmap of assay data</i> |
|--------------|---------------------------------------|

Description

Creates a heatmap of data from the indicated `TapestriObject` assay slot using the `ComplexHeatmap` package. Heatmaps are generated as transposed (i.e. x-y flipped) representations of the assay matrix. Additional `ComplexHeatmap::Heatmap()` parameters can be passed in to overwrite defaults.

Usage

```
assayHeatmap(
  TapestriExperiment,
  alt.exp = NULL,
  assay = NULL,
  split.col.by = NULL,
  split.row.by = NULL,
  annotate.row.by = NULL,
  color.preset = NULL,
  color.custom = NULL,
  ...
)
```

Arguments

| | | |
|-----------------|---|---------------------------|
| | TapestriExperiment | TapestriExperiment object |
| alt.exp | Character, altExp slot to use. NULL (default) uses top-level/main experiment. | |
| assay | Character, assay slot to use. NULL (default) uses first-indexed assay (usually "counts"). | |
| split.col.by | Character, rowData column to split columns by, i.e. "chr" or "arm". Default NULL. | |
| split.row.by | Character, colData column to split rows by, i.e. "cluster". Default NULL. | |
| annotate.row.by | Character, colData column to use for block annotation. Default NULL. | |
| color.preset | Character, color preset to use for heatmap color, either "copy.number" or "copy.number.denoise" (see Details). Overrides color.custom. NULL (default) uses default ComplexHeatmap coloring. | |
| color.custom | Color mapping function given by <code>circlize::colorRamp2()</code> . color.preset must be NULL. | |
| ... | Additional parameters to pass to <code>ComplexHeatmap::Heatmap()</code> . | |

Value

A ComplexHeatmap object

Options for color.preset**"copy.number":**

Blue-white-red gradient from 0-2-4. 4 to 8+ is red-black gradient.

```
circlize::colorRamp2(c(0,1,2,3,4,8),
c('#2c7bb6', '#abd9e9', '#ffffff', '#fdae61', '#d7191c', "black"))
```

"copy.number.denoise":

Similar to 'copy.number' present, but white range is from 1.5-2.5 to reduce the appearance of noise around diploid cells.

```
circlize::colorRamp2(c(0,1,1.5,2,2.5,3,4,8),
c('#2c7bb6', '#abd9e9', '#ffffff', '#ffffff', '#ffffff', '#fdae61', '#d7191c', "black"))
```

See Also

[Heatmap](#)

Examples

```
tap.object <- newTapestriExperimentExample() # example TapestriExperiment object
assayHeatmap(tap.object,
  assay = "counts", split.row.by = "test.cluster",
  annotate.row.by = "test.cluster", split.col.by = "chr"
)
```

| | |
|----------------|---|
| calcCopyNumber | <i>Calculate relative copy number value for each cell-probe unit using reference sample</i> |
|----------------|---|

Description

calcCopyNumber() transforms the normalized count matrix normcounts of a TapestryExperiment object into copy number values based on a set of reference cell barcodes and given copy number value (e.g. 2 for diploid). This is practically used to set the median copy number of a usually diploid reference cell population to a known copy number value, e.g. 2, and then calculate the copy number for all the cells relative to that reference population. This occurs individually for each probe, such that the result is one copy number value per cell barcode per probe (cell-probe unit). control.copy.number is a data.frame lookup table used to indicate the copy number value and cell barcodes to use as the reference. A template for control.copy.number can be generated using generateControlCopyNumberTemplate(), which will have a row for each chromosome arm represented in TapestryExperiment.

The control.copy.number data.frame should include 3 columns named arm, copy.number, and sample.label. arm is chromosome arm names from chr1p through chrXq, copy.number is the reference copy number value (2 = diploid), and sample.label is the value corresponding to the colData column given in sample.feature to indicate the set of reference cell barcodes to use to set the copy number. This is best used in a workflow where the cells are clustered first into their respective samples, and then one cluster is used as the reference population the other clusters. This also allows for the baseline copy number to be set for each chromosome arm individually in the case where the reference population is not completely diploid.

Usage

```
calcCopyNumber(
  TapestryExperiment,
  control.copy.number,
  sample.feature = "cluster",
  remove.bad.probes = FALSE
)

generateControlCopyNumberTemplate(
  TapestryExperiment,
  copy.number = 2,
  sample.feature.label = NA
)
```

Arguments

TapestryExperiment
TapestryExperiment object.

control.copy.number
data.frame with columns arm, copy.number, and sample.label. See details.

sample.feature Character, colData column to use for subsetting cell.barcodes. Default "cluster".

remove.bad.probes Logical, if TRUE, probes with median normalized counts = 0 are removed from the returned TapestriExperiment. If FALSE (default), probes with median normalized counts = 0 throw error and stop function.

copy.number Numeric, sets all entries of copy.number column in output. Default 2 (diploid).

sample.feature.label Character, sets all entries of sample.label column in output.

Value

TapestriExperiment object with cell-probe copy number values in copyNumber assay slot.
data.frame with 3 columns named arm, copy.number, and sample.label

Functions

- generateControlCopyNumberTemplate(): generates a data.frame template for control.copy.number in calcCopyNumber().

Examples

```
tap.object <- newTapestriExperimentExample() # example TapestriExperiment object
tap.object <- calcNormCounts(tap.object)
control.copy.number <- generateControlCopyNumberTemplate(tap.object,
  copy.number = 2,
  sample.feature.label = "cellline1"
)
tap.object <- calcCopyNumber(tap.object,
  control.copy.number,
  sample.feature = "test.cluster"
)
tap.object <- newTapestriExperimentExample() # example TapestriExperiment object
control.copy.number <- generateControlCopyNumberTemplate(tap.object,
  copy.number = 2,
  sample.feature.label = "cellline1"
)
```

calcGMMCopyNumber *Call copy number for each cell-chromosome using Gaussian mixture models*

Description

Uses control cells to simulate expected smoothed copy number distributions for all chromosomes across each of model.components (copy number level). Then uses the distributions to calculate posterior probabilities for each cell-chromosome belonging to each of copy number level. Each cell-chromosome is assigned the copy number value for which its posterior probability is highest. This is done for both whole chromosomes and chromosome arms.

Usage

```
calcGMMCopyNumber(
  TapestriExperiment,
  cell.barcodes,
  control.copy.number,
  model.components = 1:5,
  model.priors = NULL,
  ...
)
```

Arguments

| | |
|---------------------|--|
| TapestriExperiment | TapestriExperiment object. |
| cell.barcodes | character, vector of cell barcodes to fit GMM. Usually corresponds to diploid control. |
| control.copy.number | data.frame with columns arm and copy.number, indicating of known copy number of cells in cell.barcodes. |
| model.components | numeric, vector of copy number GMM components to calculate, default 1:5 (for copy number = 1, 2, 3, 4, 5). |
| model.priors | numeric, relative prior probabilities for each GMM component. If NULL (default), assumes equal priors. |
| ... | Additional parameters to be passed to internal functions. |

Value

TapestriExperiment object with copy number calls based on the calculated GMMs, saved to gmmCopyNumber slot of smoothedCopyNumberByChr and smoothedCopyNumberByArm altExps. GMM parameters for each feature.id are saved to the metadata slot.

Examples

```
tap.object <- newTapestriExperimentExample() # example TapestriExperiment object
tap.object <- calcNormCounts(tap.object)
control.copy.number <- generateControlCopyNumberTemplate(tap.object,
  copy.number = 2,
  sample.feature.label = "cellline1"
)
tap.object <- calcCopyNumber(tap.object,
  control.copy.number,
  sample.feature = "test.cluster"
)
tap.object <- calcSmoothCopyNumber(tap.object)
tap.object <- calcGMMCopyNumber(tap.object,
  cell.barcodes = colnames(tap.object),
  control.copy.number = control.copy.number,
  model.components = 1:5
```

)

| | |
|----------------|-----------------------------|
| calcNormCounts | <i>Normalize raw counts</i> |
|----------------|-----------------------------|

Description

Normalizes raw counts from counts slot in TapestriExperiment and returns the object with normalized counts in the normcounts slot. Also calculates the standard deviation for each probe using normalized counts and adds it to rowData.

Usage

```
calcNormCounts(TapestriExperiment, method = "mb", scaling.factor = NULL)
```

Arguments

| | |
|--------------------|---|
| TapestriExperiment | TapestriExperiment object. |
| method | Character, normalization method. Default "mb". |
| scaling.factor | Numeric, optional number to scale normalized counts if method == "libNorm". Default NULL. |

Details

"mb" method performs the same normalization scheme as in Mission Bio's mosaic package for python: Counts for each barcode are normalized relative to their barcode's mean and probe counts are normalized relative to their probe's median. "libNorm" method performs library size normalization, returning the proportion of counts of each probe within a cell. The proportion is multiplied by scaling.factor if provided.

Value

TapestriExperiment object with normalized counts added to normcounts slot.

Examples

```
tap.object <- newTapestriExperimentExample() # example TapestriExperiment object
tap.object <- calcNormCounts(tap.object)
```

calcSmoothCopyNumber *Smooth copy number values across chromosomes and chromosome arms*

Description

calcSmoothCopyNumber() takes copyNumber slot values for probes on a chromosome and smooths them by median (default) for each chromosome and chromosome arm, resulting in one copy number value per chromosome and chromosome arm for each cell barcode. Cell-chromosome values are then discretized into integers by conventional rounding ($1.5 \leq x < 2.5$ rounds to 2). Smoothed copy number and discretized smoothed copy number values are stored as smoothedCopyNumber and discreteCopyNumber assays, in altExp slots smoothedCopyNumberByChr for chromosome-level smoothing, and smoothedCopyNumberByArm for chromosome arm-level smoothing.

Usage

```
calcSmoothCopyNumber(TapestriExperiment, method = "median")
```

Arguments

TapestriExperiment TapestriExperiment object.

method Character, smoothing method: median (default) or mean.

Value

TapestriExperiment with smoothedCopyNumber and discreteCopyNumber assays in altExp slots smoothedCopyNumberByChr and smoothedCopyNumberByArm.

Examples

```
tap.object <- newTapestriExperimentExample() # example TapestriExperiment object
tap.object <- calcNormCounts(tap.object)
control.copy.number <- generateControlCopyNumberTemplate(tap.object,
  copy.number = 2,
  sample.feature.label = "cellline1"
)
tap.object <- calcCopyNumber(tap.object,
  control.copy.number,
  sample.feature = "test.cluster"
)
tap.object <- calcSmoothCopyNumber(tap.object)
```

callSampleLabels *Call sample labels based on feature counts*

Description

callSampleLabels() assigns labels (stored as colData column) to cells using feature count data in colData. This is most useful for assigning barcode labels based on barcoded reads (see [countBar-codedReads](#)). For method = max, labels are dictated by whichever input . features column has the highest number of counts. By default, ties are broken by choosing whichever label has the lowest index position (ties.method = "first"). Samples with 0 counts for all input . features columns are labeled according to neg . label. If only one feature column is used, labels are assigned to cells with counts > min . count . threshold, and neg . label otherwise.

Usage

```
callSampleLabels(
  TapestriExperiment,
  input.features,
  output.feature = "sample.call",
  return.table = FALSE,
  neg.label = NA,
  method = "max",
  ties.method = "first",
  min.count.threshold = 1
)
```

Arguments

TapestriExperiment A TapestriExperiment object.

input.features Character vector, column names in colData to evaluate.

output.feature Character, column name to use for the call output. Default "sample.call".

return.table Logical, if TRUE, returns a data.frame of the sample.calls. If FALSE (default), returns updated TapestriExperiment object.

neg.label Character, label for samples with no counts. Default NA.

method Character, call method. Only "max" currently supported, calls based on whichever input . features column has the most counts.

ties.method Character, passed to max . col () indicating how to break ties. Default "first".

min.count.threshold Numeric, minimum number of counts per cell to use for call. Default 1.

Value

A TapestriExperiment object with sample calls added to colData column sample.name. If return.table == TRUE, a data.frame of sample calls.

Examples

```
tap.object <- newTapestriExperimentExample() # example TapestriExperiment object
colData(tap.object)$gRNA1 <- 2 # example barcode counts
colData(tap.object)$gRNA2 <- 10 # example barcode counts
tap.object <- callSampleLabels(tap.object,
  input.features = c("gRNA1", "gRNA2"),
  output.feature = "sample.grna"
)
```

 corner

Print the top-left corner of a matrix

Description

Outputs up to 5 rows and columns of the input matrix object (with rownames and colnames) to get a quick look without filling the console.

Usage

```
corner(input.mat)
```

Arguments

`input.mat` A matrix-like object.

Value

A matrix-like object matching input class, subset to a maximum of 5 rows and columns.

Examples

```
tap.object <- newTapestriExperimentExample() # example TapestriExperiment object
corner(assay(tap.object, "counts"))
```

 countBarcodedReads

Get read counts from barcoded reads

Description

`countBarcodedReads()` and `countBarcodedReadsFromContig()` match exogenous DNA barcode sequences to their associated cell barcodes and saves them to the `colData` (cell barcode metadata) of `TapestriExperiment`. `countBarcodedReads()` is a shortcut for `countBarcodedReadsFromContig()`, allowing the user to specify 'gRNA' or 'barcode' to use the `grnaCounts` or `barcodeCounts` altExp slots. The entries in the barcode lookup table do not have to be present in the sample, allowing users to keep one master table/file of available barcode sequences for use in all experiments. The `Rsamtools` and `Biostrings` packages must be installed to use these functions.

Usage

```

countBarcodedReads(
  TapestriExperiment,
  bam.file,
  barcode.lookup,
  probe,
  return.table = FALSE,
  max.mismatch = 2,
  with.indels = FALSE,
  ...
)

countBarcodedReadsFromContig(
  bam.file,
  barcode.lookup,
  contig,
  cell.barcode.tag = "RG",
  max.mismatch = 2,
  with.indels = FALSE
)

```

Arguments

| | |
|--------------------|--|
| TapestriExperiment | TapestriExperiment object |
| bam.file | File path of BAM file. .bai BAM index file must be in the same location (can be generated using Rsamtools::indexBam()). |
| barcode.lookup | data.frame where the first column is the barcode identifier/name and the second column is the DNA sequence. Headers are ignored. |
| probe | Character, either "gRNA" or "barcode" to parse counts from grnaCounts or barcodeCounts altExp slots, respectively. |
| return.table | Logical, if TRUE, returns table of read counts per barcode. If FALSE, returns TapestriExperiment. Default FALSE. |
| max.mismatch | Numeric, the maximum and minimum number of mismatching letters allowed. Default 2. |
| with.indels | If TRUE, then indels are allowed. Default FALSE. |
| ... | Arguments to pass on to countBarcodedReadsFromContig(). |
| contig | Character, contig or chromosome name to search for barcodes in. Can be a vector of more than one contig to expand search space. |
| cell.barcode.tag | Character of length 2, indicates cell barcode field in BAM, specified by Tapestri pipeline (currently "RG"). Default "RG". |

Value

TapestriExperiment with barcoded read counts added to colData.
 A data.frame of read counts for each specified barcode.

See Also

[Rsamtools::indexBam\(\)](#)
[Biostrings::matchPattern\(\)](#)

Examples

```
## Not run:
counts <- countBarcodedReads(
  TapestriExperiment,
  bam.file, barcode.lookup, "gRNA"
)

## End(Not run)
## Not run:
counts <- countBarcodedReadsFromContig(bam.file, barcode.lookup, "virus_ref2")

## End(Not run)
```

createTapestriExperiment

Create TapestriExperiment object from Tapestri Pipeline output

Description

createTapestriExperiment() constructs a TapestriExperiment container object from data stored in the .h5 file output by the Tapestri Pipeline. Read count matrix (probe x cell barcode) is stored in the "counts" assay slot of the top-level experiment. Allele frequency matrix (variant x cell barcode) is stored in the "alleleFrequency" assay slot of the "alleleFrequency" altExp (alternative experiment) slot. panel.id is an optional shortcut to set special probe identities for specific custom panels.

Usage

```
createTapestriExperiment(
  h5.filename,
  panel.id = NULL,
  get.cytobands = TRUE,
  genome = "hg19",
  move.non.genome.probes = TRUE,
  filter.variants = TRUE,
  verbose = TRUE
)
```

Arguments

h5.filename File path for .h5 file from Tapestri Pipeline output.

| | |
|-------------------------------------|--|
| <code>panel.id</code> | Character, Tapestri panel ID, either CO261, CO293, CO610, or NULL. Initializes <code>barcodeProbe</code> and <code>grnaProbe</code> slots. Default NULL. |
| <code>get.cytobands</code> | Logical, if TRUE (default), retrieve and add chromosome cytobands and chromosome arms to <code>rowData</code> (probe metadata). |
| <code>genome</code> | Character, reference genome for pulling cytoband coordinates and chromosome arm labels (see <code>getCytobands()</code>). Only "hg19" (default) is currently supported. |
| <code>move.non.genome.probes</code> | Logical, if TRUE (default), move counts and metadata from non-genomic probes to <code>altExp</code> slots (see <code>moveNonGenomeProbes()</code>). |
| <code>filter.variants</code> | Logical, if TRUE (default), only stores variants that have passed Tapestri Pipeline filters. |
| <code>verbose</code> | Logical, if TRUE (default), metadata is output in message text. |

Value

TapestriExperiment object containing data from Tapestri Pipeline output.

Panel ID Shortcuts

`panel.id` is an optional shortcut to set the `barcodeProbe` and `grnaProbe` slots in `TapestriExperiment` for specific custom Tapestri panels.

CO261:

- `barcodeProbe` = "not specified"
- `grnaProbe` = "not specified"

CO293:

- `barcodeProbe` = "AMPL205334"
- `grnaProbe` = "AMPL205666"

CO610:

- `barcodeProbe` = "CO610_AMP351"
- `grnaProbe` = "CO610_AMP350"

Automatic Operations**Raw Data:**

Read count and allele frequency matrices are imported to their appropriate slots as described above. `filter.variants == TRUE` (default) only loads allele frequency variants that have passed internal filters in the Tapestri Pipeline. This greatly reduces the number of variants from tens of thousands to hundreds of likely more consequential variants, saving RAM and reducing operation time.

Metadata:

Several metadata sets are copied or generated and then stored in the appropriate `TapestriExperiment` slot during construction.

- Probe panel metadata stored in the .h5 file are copied to rowData.
- Basic QC stats (e.g. total number of reads per probe) are added to rowData.
- Basic QC stats (e.g. total number of reads per cell barcode) are added to colData.
- Experiment-level metadata is stored in metadata.

Optional Operations

Two additional major operations are called by default during `TapestriExperiment` construction for convenience. `get.cytobands == TRUE` (default) calls `getCytobands()`, which retrieves the chromosome arm and cytoband for each probe based on stored positional data and saves them in rowData. Some downstream smoothing and plotting functions may fail if chromosome arms are not present in rowData, so this generally should always be run. `move.non.genome.probes` calls `moveNonGenomeProbes()`, which moves probes corresponding to the specified tags to altExp (alternative experiment) slots in the `TapestriExperiment` object. The exception is probes on chromosome Y; CNVs of chrY are more rare, so we move it to an altExp for separate analysis. Probes corresponding to the barcodeProbe and grnaProbe slots, which are specified by the `panel.id` shortcut or manually (see [Custom Slot Getters and Setters](#)), are automatically moved to altExp by this operation as well. If such probes are not present, the function will only generate a warning message, so it is always safe (and recommended) to run by default. Any remaining probes that are not targeting a human chromosome and are not specified by the shortcut tags are moved to the `otherProbeCounts` slot.

See Also

`moveNonGenomeProbes()`, `getCytobands()`, which are run as part of this function by default.

Examples

```
## Not run:
tapExperiment <- createTapestriExperiment("myh5file.h5", "C0293")

## End(Not run)
```

Custom Slot Getters and Setters

Getter and Setter functions for TapestriExperiment slots

Description

Get and set custom slots in `TapestriExperiment`. Slots include `barcodeProbe` for a sample barcode probe ID and `grnaProbe` for a gRNA-associated probe ID. These are used as shortcuts for `moveNonGenomeProbes()` and `countBarcodedReads()`. `gmmParams` holds parameters and metadata for GMM copy number calling models.

Usage

```

barcodeProbe(x)

## S4 method for signature 'TapestriExperiment'
barcodeProbe(x)

barcodeProbe(x) <- value

## S4 replacement method for signature 'TapestriExperiment'
barcodeProbe(x) <- value

grnaProbe(x)

## S4 method for signature 'TapestriExperiment'
grnaProbe(x)

grnaProbe(x) <- value

## S4 replacement method for signature 'TapestriExperiment'
grnaProbe(x) <- value

gmmParams(x)

## S4 method for signature 'TapestriExperiment'
gmmParams(x)

```

Arguments

| | |
|--------------------|---------------------------------------|
| x | A TapestriExperiment object |
| value | Character, probe ID to assign to slot |
| TapestriExperiment | A TapestriExperiment object |

Value

For the getter methods `barcodeProbe`, `grnaProbe`, and `gmmParams`, the value of the given slot is returned. For the setter methods `barcodeProbe` and `grnaProbe`, a `TapestriExperiment` object is returned with modifications made to the given slot.

Functions

- `barcodeProbe(TapestriExperiment)`: `barcodeProbe` getter
- `barcodeProbe(TapestriExperiment) <- value`: `barcodeProbe` setter
- `grnaProbe(TapestriExperiment)`: `grnaProbe` getter
- `grnaProbe(TapestriExperiment) <- value`: `grnaProbe` setter
- `gmmParams(TapestriExperiment)`: `gmmParams` getter

Examples

```
tap.object <- newTapestriExperimentExample() # example TapestriExperiment object
barcodeProbe(tap.object) <- "Probe01"
barcodeProbe(tap.object)

grnaProbe(tap.object) <- "Probe02"
grnaProbe(tap.object)

gmmParams(tap.object)
```

`getChrOrder`*Get chromosome order from a string of chromosome/contig names*

Description

`getChrOrder()` takes a string of chromosome or contig names and returns the indices of the string in typical chromosome order, i.e. 1 through 22, X, Y. Contig names that do not match 1:22, X, or Y are sorted numerically and alphabetically (with numbers coming first), and added to the end of the order. The output string can then be used to sort the input string into typical chromosome order.

Usage

```
getChrOrder(chr.vector)
```

Arguments

`chr.vector` Character vector of chromosome or contig names.

Value

A numerical vector of the input vectors indices in chromosome order.

Examples

```
chr.order <- getChrOrder(c(1, "virus", 5, "X", 22, "plasmid", "Y"))
ordered.vector <- c(1, "virus", 5, "X", 22, "plasmid", "Y")[chr.order]
```

| | |
|--------------|---|
| getCytobands | <i>Add chromosome cytobands and chromosome arms to TapestriExperiment</i> |
|--------------|---|

Description

getCytobands() retrieves the chromosome arm and cytoband for each probe based on stored positional data and saves them in rowData. This is run automatically as part of [createTapestriExperiment\(\)](#). Note: Some downstream smoothing and plotting functions may fail if chromosome arms are not present in rowData.

Usage

```
getCytobands(TapestriExperiment, genome = "hg19", verbose = TRUE)
```

Arguments

| | |
|--------------------|---|
| TapestriExperiment | TapestriExperiment object. |
| genome | Character, reference genome to use. Only hg19 is currently supported. |
| verbose | Logical, if TRUE (default), progress is output as message text. |

Value

TapestriExperiment object with rowData updated to include chromosome arms and cytobands.

Examples

```
tap.object <- newTapestriExperimentExample() # example TapestriExperiment object
tap.object <- getCytobands(tap.object, genome = "hg19")
```

| | |
|------------------|---|
| getGMMBoundaries | <i>Calculate decision boundaries between components of copy number GMMs</i> |
|------------------|---|

Description

Calculate decision boundaries between components of copy number GMMs

Usage

```
getGMMBoundaries(TapestriExperiment, chromosome.scope = "chr")
```

Arguments

`TapestriExperiment`
TapestriExperiment object.

`chromosome.scope`
"chr" or "arm", for using models for either whole chromosomes or chromosome arms. Default "chr".

Value

tibble containing boundary values of GMMs for each `feature.id`.

Examples

```
tap.object <- newTapestriExperimentExample() # example TapestriExperiment object
tap.object <- calcNormCounts(tap.object)
control.copy.number <- generateControlCopyNumberTemplate(tap.object,
  copy.number = 2,
  sample.feature.label = "cellline1"
)
tap.object <- calcCopyNumber(tap.object,
  control.copy.number,
  sample.feature = "test.cluster"
)
tap.object <- calcSmoothCopyNumber(tap.object)
tap.object <- calcGMMCopyNumber(tap.object,
  cell.barcodes = colnames(tap.object),
  control.copy.number = control.copy.number,
  model.components = 1:5
)

boundaries <- getGMMBoundaries(tap.object,
  chromosome.scope = "chr"
)
```

getTidyData

Get tidy-style data from TapestriExperiment objects

Description

`getTidyData()` pulls data from the indicated assay and/or `altExp` slot(s), and rearranges it into tidy format. `colData` (cell metadata) from the top-level/main experiment is included. `rowData` (probe metadata) from the indicated assay and/or `altExp` slot(s) is included. Attempts are made to sort by "chr" and "start.pos" columns if they are present to simplify plotting and other downstream operations.

Usage

```
getTidyData(
  TapestriExperiment,
  alt.exp = NULL,
  assay = NULL,
  feature.id.as.factor = TRUE
)
```

Arguments

TapestriExperiment
TapestriExperiment object.

alt.exp
Character, altExp slot to use. NULL (default) uses top-level/main experiment.

assay
Character, assay slot to use. NULL (default) uses first-indexed assay (often "counts").

feature.id.as.factor
Logical, if TRUE (default), the feature.id column is returned as a factor.

Value

A tibble of tidy data with corresponding metadata from colData and rowData.

Examples

```
tap.object <- newTapestriExperimentExample() # example TapestriExperiment object
tidy.data <- getTidyData(tap.object, alt.exp = "alleleFrequency")
```

moveNonGenomeProbes *Move non-genome probes counts and metadata to altExp slots*

Description

moveNonGenomeProbes() takes the probe IDs corresponding to grnaProbe and barcodeProbe slots of the TapestriExperiment object, as well as probes on chrY, and moves them to their own altExp slots in the object. This allows those counts and associated metadata to be manipulated separately without interfering with the probes used for CNV measurements which target the endogenous genome. [SingleCellExperiment::splitAltExps\(\)](#) can be used for manual specification of probes to move to altExp slots if the shortcut slots are not used.

Usage

```
moveNonGenomeProbes(TapestriExperiment)
```

Arguments

TapestriExperiment
TapestriExperiment object.

Details

moveNonGenomeProbes() moves probes corresponding to the specified tags to altExp (alternative experiment) slots in the TapestriExperiment object. These probes should be those which do not correspond to a chromosome and therefore would not be used to call copy number variants. The exception is probes on chromosome Y; CNVs of chrY are more rare, so we move it to an altExp for separate analysis. Probes corresponding to the barcodeProbe and grnaProbe slots, which are specified by the panel.id shortcut or manually (see [Custom Slot Getters and Setters](#)), are automatically moved to altExp by this operation as well. If such probes are not present, the function will only generate a warning message, so it is always safe (and recommended) to run by default. Any remaining probes that are not targeting a human chromosome and are not specified by the shortcut tags are moved to the otherProbeCounts slot. This function is run automatically by default and with default behavior as part of [createTapestriExperiment\(\)](#).

Value

TapestriExperiment with altExp slots filled with counts and metadata for non-genomic probes.

See Also

[SingleCellExperiment::splitAltExps\(\)](#) for manual specification of probes to move to altExp slots.

Examples

```
tap.object <- newTapestriExperimentExample() # example TapestriExperiment
tap.object <- moveNonGenomeProbes(tap.object)
```

```
newTapestriExperimentExample
      Create Example TapestriExperiment
```

Description

Creates a TapestriExperiment object for demonstration purposes, which includes 240 probes across the genome, and 300 cells of 3 types. Raw counts are generated randomly. Type 1 has 75 cells, all XY, all diploid. Type 2 has 100 cells, all XX, with 3 copies of chr 7, otherwise diploid. Type 3 has 125 cells, all XY, with 1 copy of chr 1p, otherwise diploid.

Usage

```
newTapestriExperimentExample()
```

Value

TapestriExperiment object with demo data.

Examples

```
tapExperiment <- newTapestriExperimentExample()
```

PCAKneePlot *Plot of PCA proportion of variance explained*

Description

Draws "knee plot" of PCA proportion of variance explained to determine which principal components (PCs) to include for downstream applications e.g. clustering. Variance explained for each PC is indicated by the line. Cumulative variance explained is indicated by the bars.

Usage

```
PCAKneePlot(TapestriExperiment, alt.exp = "alleleFrequency", n.pcs = 10)
```

Arguments

| | |
|--------------------|---|
| TapestriExperiment | TapestriExperiment object |
| alt.exp | Character, altExp to use, NULL uses top-level/main experiment. Default "alleleFrequency". |
| n.pcs | Numeric, number of PCs to plot, starting at 1. Default 10. |

Value

ggplot2 object, combined line plot and bar graph

Examples

```
tap.object <- newTapestriExperimentExample() # example TapestriExperiment object
tap.object <- runPCA(tap.object, alt.exp = "alleleFrequency")
PCAKneePlot(tap.object, n.pcs = 5)
```

plotCopyNumberGMM *Plot copy number GMM components*

Description

Plots the probability densities of GMM components for given chromosome or chromosome arm, store in a TapestriExperiment. [calcGMMCopyNumber\(\)](#) must be run first.

Usage

```
plotCopyNumberGMM(
  TapestriExperiment,
  feature.id = 1,
  chromosome.scope = "chr",
  draw.boundaries = FALSE
)
```

Arguments

`TapestriExperiment`
TapestriExperiment object.

`feature.id` chromosome or chromosome arm to plot.

`chromosome.scope`
"chr" or "arm", for plotting models for either whole chromosomes or chromosome arms.

`draw.boundaries`
logical, if TRUE, draw decision boundaries between each Gaussian component.

Value

ggplot object, density plot

Examples

```
tap.object <- newTapestriExperimentExample() # example TapestriExperiment object
tap.object <- calcNormCounts(tap.object)
control.copy.number <- generateControlCopyNumberTemplate(tap.object,
  copy.number = 2,
  sample.feature.label = "cellline1"
)
tap.object <- calcCopyNumber(tap.object,
  control.copy.number,
  sample.feature = "test.cluster"
)
tap.object <- calcSmoothCopyNumber(tap.object)
tap.object <- calcGMMCopyNumber(tap.object,
  cell.barcodes = colnames(tap.object),
  control.copy.number = control.copy.number,
  model.components = 1:5
)

tap.object <- plotCopyNumberGMM(tap.object,
  feature.id = 7,
  chromosome.scope = "chr",
  draw.boundaries = TRUE
)
```

`reducedDimPlot`*Scatter plot for dimensional reduction results*

Description

Plots a scatter plot of the indicated dimensional reduction results.

Usage

```
reducedDimPlot(
  TapestriExperiment,
  alt.exp = "alleleFrequency",
  dim.reduction,
  dim.x = 1,
  dim.y = 2,
  group.label = NULL
)
```

Arguments

| | |
|--------------------|---|
| TapestriExperiment | TapestriExperiment object |
| alt.exp | Character, altExp to use, NULL uses top-level/main experiment. Default "alleleFrequency". |
| dim.reduction | Character, dimension reduction to plot, either "PCA" or "UMAP". |
| dim.x | Numeric, index of dimensional reduction data to plot on X axis. Default 1. |
| dim.y | Numeric, index of dimensional reduction data to plot on Y axis. Default 2. |
| group.label | Character, colData column for grouping samples by color. Default NULL. |

Value

ggplot2 object, scatter plot

Examples

```
tap.object <- newTapestriExperimentExample() # example TapestriExperiment object
tap.object <- runPCA(tap.object, alt.exp = "alleleFrequency")
reducedDimPlot(tap.object, dim.reduction = "pca")
```

runClustering

Cluster 2D data

Description

Clusters data using dbSCAN method and saves cluster assignments for each cell barcode to colData. Generally used to assign clusters to UMAP projection after PCA and UMAP dimensional reduction.

Usage

```
runClustering(
  TapestriExperiment,
  alt.exp = "alleleFrequency",
  dim.reduction = "UMAP",
  eps = 0.8,
```



```

    dim.1 = 1,
    dim.2 = 2,
    ...
  )

```

Arguments

| | |
|--------------------|---|
| TapestriExperiment | TapestriExperiment object |
| alt.exp | Character, altExp slot to use. NULL uses top-level/main experiment. Default "alleleFrequency". |
| dim.reduction | Character, reduced dimension data to use. Default "UMAP". |
| eps | Numeric, dbSCAN eps parameter. Lower to increase cluster granularity. See dbSCAN::dbSCAN() . Default 0.8. |
| dim.1 | Numeric, index of data dimension to use. Default 1. |
| dim.2 | Numeric, index of data dimension to use. Default 2. |
| ... | Additional parameters to pass to dbSCAN::dbSCAN() . |

Value

TapestriExperiment object with updated colData containing cluster assignments.

See Also

[dbSCAN::dbSCAN\(\)](#)

Examples

```

tap.object <- newTapestriExperimentExample() # example TapestriExperiment object
tap.object <- runPCA(tap.object, alt.exp = "alleleFrequency")
tap.object <- runUMAP(tap.object, pca.dims = 1:3)
tap.object <- runClustering(tap.object, dim.reduction = "UMAP", eps = 0.8)

```

runPCA

Cluster assay data by Principal Components Analysis

Description

Analyzes assay data by Principal Components Analysis (PCA) and saves results to reducedDims slot of TapestriObject.

Usage

```
runPCA(  
  TapestriExperiment,  
  alt.exp = "alleleFrequency",  
  assay = NULL,  
  sd.min.threshold = NULL,  
  center = TRUE,  
  scale. = TRUE  
)
```

Arguments

| | |
|--------------------|--|
| TapestriExperiment | TapestriExperiment object |
| alt.exp | Character, altExp to use, NULL uses top-level/main experiment. Default "alleleFrequency". |
| assay | Character, assay to use. NULL (default) uses first-indexed assay. |
| sd.min.threshold | Numeric, minimum threshold for allelefreq.sd. Increase to run PCA on fewer, more variable dimensions. Set to NULL if not using for alleleFrequency slot. Default NULL. |
| center | Logical, if TRUE (default), variables are shifted to be zero centered. See stats::prcomp() . |
| scale. | Logical, if TRUE (default), variables are scaled to have unit variance prior to PCA. See stats::prcomp() . |

Value

TapestriExperiment with PCA results saved to reducedDims slot of altExp, and proportion of variance explained by each PC saved to metadata slot of altExp.

See Also

[stats::prcomp\(\)](#) for PCA method details.

Examples

```
tap.object <- newTapestriExperimentExample() # example TapestriExperiment  
tap.object <- runPCA(tap.object, alt.exp = "alleleFrequency")
```

| | |
|---------|------------------------------------|
| runUMAP | <i>Cluster matrix data by UMAP</i> |
|---------|------------------------------------|

Description

Analyzes matrix data by UMAP and saves results to reducedDims slot of TapestryObject.

Usage

```
runUMAP(  
  TapestryExperiment,  
  alt.exp = "alleleFrequency",  
  assay = NULL,  
  use.pca.dims = TRUE,  
  pca.dims = NULL,  
  ...  
)
```

Arguments

| | |
|--------------------|--|
| TapestryExperiment | TapestryExperiment object |
| alt.exp | Character, altExp to use, NULL uses top-level/main experiment. Default "alleleFrequency". |
| assay | Character, assay to use. NULL (default) uses first-indexed assay. Not used when use.pca.dims = TRUE. |
| use.pca.dims | Logical, if TRUE, uses experiment PCA, otherwise uses assay data. Default TRUE. |
| pca.dims | Numeric, indices of PCs to use in UMAP. Default NULL. |
| ... | Additional parameters to pass to <code>umap::umap()</code> , e.g. for configuration (see <code>umap::umap.defaults()</code>). |

Value

TapestryExperiment with UMAP embeddings saved to reducedDims slot of altExp.

Examples

```
tap.object <- newTapestryExperimentExample() # example TapestryExperiment object  
tap.object <- runPCA(tap.object, alt.exp = "alleleFrequency")  
tap.object <- runUMAP(tap.object, pca.dims = 1:3)
```

TapestriExperiment-class

TapestriExperiment Class Definition

Description

TapestriExperiment Class Definition

Usage

```
## S4 method for signature 'TapestriExperiment'  
show(object)
```

Arguments

| | |
|--------------------|-----------------------------|
| object | An R object |
| TapestriExperiment | A TapestriExperiment object |

Value

TapestriExperiment object

Methods (by generic)

- show(TapestriExperiment): Show method for TapestriExperiment

Slots

barcodeProbe character.
grnaProbe character.
gmmParams list.

Examples

```
tapExpObject <- new("TapestriExperiment")
```

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