Package ‘multistateQTL’

May 30, 2024

Type  Package
Date  2024-03-22
Title  Toolkit for the analysis of multi-state QTL data
Version  1.0.0
Description  A collection of tools for doing various analyses of multi-state QTL data, with a focus on visualization and interpretation. The package ‘multistateQTL’ contains functions which can remove or impute missing data, identify significant associations, as well as categorise features into global, multi-state or unique. The analysis results are stored in a 'QTLExperiment' object, which is based on the 'SummarisedExperiment' framework.
License  GPL-3
URL  https://github.com/dunstone-a/multistateQTL
BugReports  https://github.com/dunstone-a/multistateQTL/issues
Encoding  UTF-8
Depends  QTLExperiment, SummarizedExperiment, ComplexHeatmap, data.table, collapse
Imports  methods, S4Vectors, grid, dplyr, tidyr, matrixStats, stats, fitdistrplus, viridis, ggplot2, circlize, mashr, grDevices
Suggests  testthat, BiocStyle, knitr, covr, rmarkdown
biocViews  FunctionalGenomics, GeneExpression, Sequencing, Visualization, SNP, Software
VignetteBuilder  knitr
RoxygenNote  7.3.1
git_url  https://git.bioconductor.org/packages/multistateQTL
git_branch  RELEASE_3_19
git_last_commit  7c5eeae
git_last_commit_date  2024-04-30
Repository  Bioconductor 3.19
Date/Publication  2024-05-29
multistateQTL-package

multistateQTL-package

multistateQTL: Toolkit for the analysis of multi-state QTL data

Description

A collection of tools for doing various analyses of multi-state QTL data, with a focus on visualization and interpretation. The package 'multistateQTL' contains functions which can remove or impute missing data, identify significant associations, as well as categorise features into global, multi-state or unique. The analysis results are stored in a 'QTLExperiment' object, which is based on the 'SummarisedExperiment' framework.

Author(s)

Maintainer: Amelia Dunstone <amelia.dunstone@svi.edu.au> (ORCID) [contributor]

Authors:

- Christina Del Azodi <cazodi@svi.edu.au>

Other contributors:

- Davis McCarthy <dmccarthy@svi.edu.au> [contributor]
callSignificance

See Also

Useful links:
- https://github.com/dunstone-a/multistateQTL
- Report bugs at https://github.com/dunstone-a/multistateQTL/issues

Description

Call associations as significant in each state.

Usage

callSignificance(object, ...)
callSignificance(object, ...) <- value

## S4 method for signature 'QTLExperiment'
callSignificance(
  object,
  thresh = 0.05,
  secondThresh = thresh,
  feature = .feature_id,
  assay = "pvalues",
  mode = "simple",
  p.adjust.method = "fdr",
  ...
)

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>object</td>
<td>A QTLExperiment object.</td>
</tr>
<tr>
<td>...</td>
<td>arguments passed to callSignificance</td>
</tr>
<tr>
<td>value</td>
<td>Value to place in getSignificance</td>
</tr>
<tr>
<td>thresh</td>
<td>Significance threshold.</td>
</tr>
<tr>
<td>secondThresh</td>
<td>Significance threshold for associations with significance in one state.</td>
</tr>
<tr>
<td>feature</td>
<td>rowData column name with feature identifier.</td>
</tr>
<tr>
<td>assay</td>
<td>assay name with significance score.</td>
</tr>
<tr>
<td>mode</td>
<td>Method to determine significance threshold per state. Options are ‘simple’, ‘feature-wise-FDR’, and ‘global-FDR’.</td>
</tr>
<tr>
<td>p.adjust.method</td>
<td>Method of multiple-test correction if mode != simple</td>
</tr>
</tbody>
</table>
**getComplete**

**Details**

This function adds a new assay to multistateQTL object with TRUE/FALSE significance calls for each test for each state.

**Value**

A ‘QTLExperiment’ object with a new assay called ‘significant’ and with a column called nSignificant added to the colData.

**Author(s)**

Christina B Azodi

**See Also**

wilcox.test, on which this function is based.

**Examples**

```r
qtle <- mockQTL()
assays(qtle)
qtle <- callSignificance(qtle)
# There is now an assay called 'significant'
assays(qtle)

# Use feature-wise FDR correction -------------------------------------------
qtle_feat <- callSignificance(qtle, thresh=0.1, mode="feature-wise-FDR")
```

---

**filterQTLBasedMissing**

**Filter QTLExperiment based on missing data**

**Description**

Method to filter QTLExperiment objects to remove tests with greater than the permitted rate of missing values.

**Usage**

callComplete(qtle, n = 1, verbose = FALSE)

**Arguments**

- **qtle** A ‘QTLExperiment‘ object
- **n** Number (or percent if n < 1) of states requiring non-null values
- **verbose** logical. Whether to print progress messages.
**getSignificant**

**Value**

a subset of the ‘QTLExperiment’ object, with only tests with fewer NAs than specified by n.

**Examples**

```r
# Create a QTLExperiment object with NA values ------------------------------
sim <- qtleSimulate(
  nStates=10, nFeatures=100, nTests=1000,
  global=0.2, multi=0.4, unique=0.2, k=2)

# Randomly remove 1000 elements from the betas matrix.
na_pattern <- sample(seq(1, ncol(sim)*nrow(sim)), 1000)
sim_na <- sim
assay(sim_na, "betas")[na_pattern] <- NA

# Original object has more rows than the output of getComplete()
dim(sim)
dim(sim_na)

dim(sim_complete) <- getComplete(sim_na)
dim(sim_complete)
```

---

**getSignificant**  
Filter to only QTLs significant in at least one state

**Description**

Filter to only QTLs significant in at least one state

**Usage**

`getSignificant(qtle, n = 1, assaySig = "significant", verbose = FALSE)`

**Arguments**

- `qtle`  
  ‘QTLExperiment’ object
- `n`  
  Number (or percent if n < 1) of states with significant association
- `assaySig`  
  The assay containing TRUE/FALSE significance calls for each QTL test.
- `verbose`  
  logical. Whether to print progress messages.

**Value**

a subset of the ‘QTLExperiment’ object, where all rows are significant in at least one state.
Examples

```r
qtle <- mockQTLE()
qtle <- callSignificance(qtle)
dim(qtle)
qtle_sig <- getSignificant(qtle)

# There are fewer rows because we have removed tests which are not significant
# in any state.
dim(qtle_sig)
```

---

getTopHits

Filter QTLExperiment to keep only top hits

Description

Method to return a subset of a QTLExperiment object containing only the tests that are top hits. Top hits are defined as the test for each feature with the most significant test statistic. Returns an array of the top QTL for each feature across all states.

Usage

```r
getTopHits(
  qtle,
  mode = c("global", "state"),
  assay = "pvalues",
  assaySig = "significant",
  verbose = FALSE
)
```

Arguments

- `qtle`: A ‘QTLExperiment’ object.
- `mode`: global/state to specify if the top hit per feature is desired from across all states or for each state.
- `assay`: The assay containing the test statistic to minimize.
- `assaySig`: The assay containing TRUE/FALSE significance calls for each QTL test.
- `verbose`: logical. Whether to print progress messages.

Value

A subset of the ‘QTLExperiment’ object, with only tests that are the top hits for each feature (‘mode=global’) or for each feature for each state (‘mode=state’).
Examples

```r
sumstats <- mockSummaryStats(nStates=10, nQTL=100, names=TRUE)
qtle <- QTLExperiment(
  assay=list(
    betas=sumstats$betas,
    errors=sumstats$errors,
    pvalues=sumstats$pvalues,
    lfsrs=sumstats$lfsrs)
)

# Add 'significant' assay to object
qtle <- callSignificance(qtle)

# Filter to the top tests for each feature
qtle_glob <- getTopHits(qtle, assay="lfsrs", mode="global", verbose = TRUE)
# There are 3 rows corresponding to the three features.
table(feature_id(qtle_glob))

# At most one QTL is retained for each combination of feature_id and state_id
qtle_feat <- getTopHits(qtle, assay="lfsrs", mode="state", verbose = TRUE)
table(feature_id(qtle_feat))
```

---

**plotCompareStates**  
*Compare QTL between two states*

**Description**

Convenience method for comparing the assay value, specified by assay, between two states.

**Usage**

```r
plotCompareStates(
  object,
  x,
  y,
  assay = "betas",
  FUN = identity,
  assaySig = "significant",
  alpha = 0.2,
  colBoth = "#4477AA",
  colDiverging = "#EE6677",
  colNeither = "gray50",
  colX = "#CCBB44",
  colY = "#AA3377"
)
```
plotPairwiseSharing

Heatmap of pairwise QTL sharing with state-level annotations

**Arguments**

- **object**: an QTLExperiment object.
- **x**: Name of state for x-axis
- **y**: Name of state for y-axis
- **assay**: Name of assay to plot.
- **FUN**: Function to be applied to fillBy assay before plotting (e.g. identity, abs, log10).
- **assaySig**: name of assay with TRUE/FALSE significance calls.
- **alpha**: Transparency.
- **colBoth**: Color for tests significant in both states.
- **colDiverging**: Color for tests significant in both states, with diverging effect sizes.
- **colNeither**: Color for null tests.
- **colX**: Color for tests significant in the x-axis state only.
- **colY**: Color for tests significant in the y-axis state only.

**Value**

Returns a list containing the counts for each color category and the plot object.

**Examples**

```r
sim <- qtleSimulate(
  nStates=10, nFeatures=100, nTests=1000,
  global=0.2, multi=0.4, unique=0.2, k=2)
sim <- callSignificance(sim, mode="simple", assay="lfsrs",
  thresh=0.0001, secondThresh=0.0002)
sim_sig <- getSignificant(sim)
sim_top <- getTopHits(sim_sig, assay="lfsrs", mode="state")
sim_top <- runPairwiseSharing(sim_top)
sim_top <- runTestMetrics(sim_top)
plotCompareStates(sim_top, x="S01", y="S02")
```

**Description**

Methods to plot a heatmap of the pairwise sharing of QTL as calculated by `runPairwiseSharing`. 
plotPairwiseSharing

Usage

plotPairwiseSharing(
  object,
  slot = "pairwiseSharing",
  annotateRowsBy = NULL,
  annotateColsBy = NULL,
  annotateCells = FALSE,
  colourRange = NULL,
  name = "colnames",
  distMethod = "pearson",
  size = 8,
  ...
)

Arguments

  object  A QTLExperiment object
  slot    Name of slot in metadata list with Pairwise Sharing matrix.
  annotateRowsBy character or array of characters specifying the column(s) in colData to be plotted
                as row annotations.
  annotateColsBy character or array of characters specifying the column(s) in colData to be plotted
                as column annotations.
  annotateCells Logical to annotate cells with their values.
  colourRange Optional range for the color legend
  name      character specifying the column in colData to use to label rows and columns. Default is
            colnames(qtle).
  distMethod Distance method used for hierarchical clustering. Valid values are the supported
                methods in dist() function.
  size      numeric scalar giving default font size for plotting theme.
  ...       further arguments passed to Rtsne

Value

Returns a ComplexHeatmap object.

Examples

sim <- qtleSimulate(
  nStates=10, nFeatures=100, nTests=1000,
  global=0.2, multi=0.4, unique=0.2, k=2)
sim <- callSignificance(sim, mode="simple", assay="lfsrs",
  thresh=0.0001, secondThresh=0.0002)
sim_sig <- getSignificant(sim)
sim_top <- getTopHits(sim_sig, assay="lfsrs", mode="state")
sim_top <- runPairwiseSharing(sim_top)
plotPairwiseSharing(sim_top)
# Plot with complex column annotations
plotPairwiseSharing(sim_top, annotateColsBy = c("nSignificant", "multistateGroup"))

---

### Description

Convenience method for plotting values from any assay specified by fillBy across states and tests.

### Usage

```r
plotQTLClusters(
  object,
  fillBy = "betas",
  FUN = identity,
  minSig = 1,
  annotateColsBy = NULL,
  annotateRowsBy = NULL,
  show_row_names = FALSE,
  state_id = "state_id",
  columnOrder = NULL,
  rowOrder = NULL,
  row_km = 0,
  column_km = 0
)
```

### Arguments

- **object**
  - an `QTLExperiment` object.
- **fillBy**
  - name of assay to use for main heatmap object.
- **FUN**
  - Function to be applied to fillBy assay before plotting (e.g. identity, abs, log10).
- **minSig**
  - minimum number of significant states for QTL to be included.
- **annotateColsBy**
  - character or array of characters specifying the column(s) in colData to be plotted to annotate states.
- **annotateRowsBy**
  - character or array of characters specifying the column(s) in rowData to be plotted to annotate QTL tests.
- **show_row_names**
  - Logical to plot row (i.e. test) names.
- **state_id**
  - colData column to use to label states.
- **columnOrder**
  - Null for clustering or array to overwrite state order.
- **rowOrder**
  - Null for clustering or array to overwrite test order.
- **row_km**
  - Set k for k-means clustering of tests.
- **column_km**
  - Set k for k-means clustering of states.
plotSimulationParams

Value

Returns a ComplexHeatmap object.

Examples

```r
sim <- qtleSimulate(
  nStates=10, nFeatures=100, nTests=1000,
  global=0.2, multi=0.4, unique=0.2, k=2)
sim <- callSignificance(sim, mode="simple", assay="lfsrs",
  thresh=0.0001, secondThresh=0.0002)

sim_sig <- getSignificant(sim)
sim_top <- getTopHits(sim_sig, assay="lfsrs", mode="state")
sim_top <- runTestMetrics(sim_top)
sim_top <- runPairwiseSharing(sim_top)
sim_top_ms <- subset(sim_top, qtl_type_simple == "multistate")

plotQTLClusters(sim_top)

# Plot with annotations for multi state group
plotQTLClusters(sim_top_ms, annotateColsBy = c("multistateGroup"),
  annotateRowsBy = c("qtl_type", "mean_beta", "QTL"))
```

plotSimulationParams  Distribution of estimated simulation parameters

Description

Distribution of estimated simulation parameters

Usage

`plotSimulationParams(params, n = 1e+05)`

Arguments

- `params`: Estimated simulation parameters from qtleEstimate.
- `n`: Number of random values to sample for plots.

Value

A ggplot2 object
Examples

```r
qtle <- mockQTLTE()
params <- qtleEstimate(qtle, threshSig = 0.05, threshNull = 0.5)
plotSimulationParams(params=params)
```

---

**plotUpSet**

*Upset plot of QTL sharing between states with state-level annotations*

**Description**

Convenient method to plot an UpSet plot showing the number of QTL that are significant in sets of states.

**Usage**

```r
plotUpSet(
    object, 
    assay = "significant", 
    name = "colnames", 
    minShared = 10, 
    minDegree = 2, 
    maxDegree = NULL, 
    annotateColsBy = NULL, 
    comb_order = "comb_size", 
    set_order = order(ss), 
    ... 
)
```

**Arguments**

- **object**: an QTLExperiment object
- **assay**: Name of assay to use to assess significance.
- **name**: character specifying the column in colData to use to label rows and columns. Default is colnames(qtle).
- **minShared**: minimum number of shared QTL for set to be included.
- **minDegree**: minimum degree of sharing for set to be included.
- **maxDegree**: maximum degree of sharing for set to be included.
- **annotateColsBy**: character or array of characters specifying the column(s) in colData to be plotted as annotations.
- **comb_order**: characters specifying how sets should be ordered. Options include the set size (set_size), combination size (comb_size), degree (deg).
- **set_order**: Array specifying order of states.
- **...**: further arguments passed to `Rtsne`
qtleEstimate

Value

Returns a ComplexHeatmap object.

Examples

```r
sim <- qtleSimulate(
    nStates=10, nFeatures=100, nTests=1000,
    global=0.2, multi=0.4, unique=0.2, k=2)
sim <- callSignificance(sim, mode="simple", assay="lfsrs",
    thresh=0.0001, secondThresh=0.0002)
sim_sig <- getSignificant(sim)
sim_top <- getTopHits(sim_sig, assay="lfsrs", mode="state")
sim_top <- runPairwiseSharing(sim_top)
plotUpSet(sim_top)

# Upset plot with complex row annotations
plotUpSet(sim_top, annotateColsBy = c("nSignificant", "multistateGroup"))
```

qtleEstimate

| qtleEstimate | Estimate parameters from real data for simulating multi-state QTL summary statistics |

Description

Estimate parameters from real data for simulating multi-state QTL summary statistics

Usage

```r
qtleEstimate(
    data,
    assay = "pvalues",
    threshSig = 0.001,
    threshNull = 0.1,
    verbose = TRUE
)
```

Arguments

data | A ‘QTLExperiment’ object or named list containing "betas" and "errors" matrices.

assay | Assay containing test statistic information to use.

threshSig | Max threshold (pval/lfsr) for calling tests as significant.

threshNull | Min threshold (pval/lfsr) for calling tests as null.

verbose | Logical.
Details
The simulation consists of user-defined numbers of equal numbers of four different types of effects: null, equal among conditions, present only in the first condition, independent across conditions.

Value
A list with parameter estimates for the QTLExperiment object.

Examples
```r
qtle <- mockQTLEx()
qtleEstimate(qtle)
```

---

qtleParams

**Default qtle simulation parameters**

Description
Returns a list of the default values used for parameters when simulating multistate QTL data. Parameters include:

- `betas.sig.shape`
- `betas.sig.rate`
- `cv.sig.shape`
- `cv.sig.rate`
- `betas.null.shape`
- `betas.null.rate`
- `cv.null.shape`
- `cv.null.rate`

Usage
```r
qtleParams()
```

Details
The default parameters returned by this function were generated using expression QTL (eQTL) summary statistics from the Genotype-Tissue Expression (GTEx) Project (Version 8) for the ten tissues with the largest sample sizes for eQTL mapping. The eQTL tests were filtered to include only eQTLs on chromosome 1 that were available in all 10 tissues.

Value
A list with the default parameter values which can be used when simulating multistate QTL data.
**Examples**

```r
qtleParams()
```

**Description**

Create simulated multistate QTL data for testing purposes

**Usage**

```r
qtleSimulate(
  params = qtleParams(),
  nTests = 100,
  nFeatures = NULL,
  nStates = 5,
  global = 0.5,
  multi = 0,
  unique = 0,
  k = 2,
  betaSd = 0.1,
  lfsr = TRUE,
  verbose = TRUE
)
```

**Arguments**

- `params`: list of parameters required to simulate betas and beta errors. Generated by `qtleEstimate()` or `qtleParams()`.
- `nTests`: number of QTL tests
- `nFeatures`: number of QTL features to simulate tests for, NULL mean `nFeatures = nTests`.
- `nStates`: number of states
- `global`: percent of QTL tests with significant effects shared across all states
- `multi`: percent of QTL tests with significant effects shared across a subset of states.
- `unique`: percent of QTL tests with significant effects in only one state
- `k`: number of multi-state clusters or an array with the cluster assignments.
- `betaSd`: The desired standard deviation or an array of standard deviations equal to the length of states for sampling beta values for each state.
- `lfsr`: Logical to calculate lfsr using mashr_1by1.
- `verbose`: Logical.
Details

The simulation consists of user defined number of equal numbers of four different types of effects: null, equal among conditions, present only in first condition, independent across conditions

Value

A simulated ‘QTLExperiment’ object.

Examples

```r
qtleSimulate(nTests=100, nStates=5, global=0.1, multi=0.2, unique=0.05)
```

## replaceNAs

*Return multistateQTL object with NAs filled in*

Description

A convenience function for imputing or filling in NAs in a ‘QTLExperiment’ object.

Usage

```r
replaceNAs(
  object,
  methods = list(betas = 0, errors = "mean", pvalues = 1, lfsrs = 1),
  verbose = FALSE
)
```

Arguments

- **object**: A ‘QTLExperiment’ object
- **methods**: A named list with the method for each assay. Available methods are to replace with a given integer or with the row mean or median.
- **verbose**: logical. Whether to print progress messages.

Value

A ‘QTLExperiment’ object with the same dimensions as the original object, but with the NA values replaced according to the input specifications.
Examples

```r
# Create a QTLExperiment object with NA values
qtle <- mockQTLE()

# Randomly remove 1000 elements from the betas matrix.
na_pattern <- sample(seq(1, ncol(qtle)*nrow(qtle)), 1000)
qtle_na <- qtle
assay(qtle_na, "betas")[na_pattern] <- NA

# There are some NA values in the "betas" assay
any(is.na(betas(qtle_na)))

qtle_complete <- replaceNAs(qtle_na)

# Now we don't have any NA values
any(is.na(betas(qtle_complete)))

## Specify a specific method to impute NAs
qtie_median <- replaceNAs(
  qtle_na,
  methods=list(betas = 0, errors = "median", pvalues = 1),
  verbose=TRUE)
```

---

**runPairwiseSharing**  
*Compute the proportion of (significant) QTL shared by pairs of conditions*

**Description**
Compute the proportion of (significant) QTL shared by pairs of conditions

**Usage**

```r
runPairwiseSharing(
  qtle,
  assay = "betas",
  assaySig = "significant",
  factor = 0.5,
  FUN = identity,
  ...
)
```

**Arguments**

- **qtle**: A 'QTLExperiment' object.
- **assay**: The assay containing the metric used to determine sharing (i.e. the metric to be within a factor X to be considered shared).
assaySig  The assay containing significance information.
factor    a number in [0,1] the factor within which effects are considered to be shared
FUN      a function to be applied to the estimated effect sizes before assessing sharing. Default 'FUN=identity', 'FUN=abs' ignores the sign of the effects when assessing sharing.
...      Additional parameters to pass on to internal functions.

Details
For each pair of states, the effects that are significant (as determined by 'callSignificance') in at least one of the two states are identified. Then the fraction of those with an estimated effect size (i.e. betas) within a factor ‘factor’ of one another is computed and returned.

Value
The ‘QTLExperiment’ object with a matrix called pairwiseSharing added to the metadata.

Examples
```r
m <- mockQTLE()
m <- callSignificance(m, assay="pvalues")
runPairwiseSharing(m) # sharing by magnitude (same sign)
runPairwiseSharing(m, factor=0) # sharing by sign
runPairwiseSharing(m, FUN=abs) # sharing by magnitude when sign is ignored
```

Description
Takes the results from ‘callSignificance()’ and from the assay ‘betas’ to categorize each QTL test using two classification strategies:
Strategy 1 (qtl_type): (1) global-shared, (2) global-diverging, (3) multi-state-shared, (4) multi-state-diverging, or (5) unique.
Strategy 2 (qtl_type_simple): (1) global, (2) multi-state, or (3) unique.

Usage
```r
runTestMetrics(
  qtle,
  assay = "betas",
  assaySig = "significant",
  globalBuffer = 0,
  ...
)
```
simPerformance

**Arguments**

- `qtle` QTLExperiment qtle
- `assay` Name of assay containing QTL effect size estimate (e.g. betas)
- `assaySig` Name of assay with TRUE/FALSE significance calls
- `globalBuffer` Number of states that can be not-significant and the QTL will still be called as global, for example, if globalBuffer=1, then a QTL will be considered global if it is significant in all or all but 1 state.
- ... arguments passed to runTestMetrics

**Details**

If a test is significant in more than one sign across different states, returns TRUE in rowData(qtle)$diverging

**Value**

The `QTLExperiment` object with the following columns added to the rowData: nSignificant, effect_sd, qtl_type, qtl_type_simple

**Examples**

```r
m <- mockQTLE()
m <- callSignificance(m)
m <- runTestMetrics(m)
```

---

**Description**

Performance metrics for multistateQTL simulations

**Usage**

```r
simPerformance(qtle, assay = "significant")
```

**Arguments**

- `qtle` A ‘multistateQTL’ object.
- `assay` Name of the ‘multistateQTL’ assay containing the significance calls

**Value**

description
Examples

```r
sim <- qtleSimulate()
sim <- callSignificance(sim, assay="lfsrs", thresh=0.1)
simPerformance(sim)
```
Index

* internal
  multistateQTL-package, 2

  callSignificance, 3
  callSignificance, QTLExperiment-method
    (callSignificance), 3
  callSignificance<-(callSignificance), 3

  getComplete, 4
  getSignificant, 5
  getTopHits, 6

  multistateQTL (multistateQTL-package), 2
  multistateQTL-package, 2

  plotCompareStates, 7
  plotPairwiseSharing, 8
  plotQTLClusters, 10
  plotSimulationParams, 11
  plotUpSet, 12

  qtleEstimate, 13
  qtleParams, 14
  qtleSimulate, 15
  QTLExperiment, 3, 4, 6

  replaceNAs, 16
  Rtsne, 9, 12
  runPairwiseSharing, 17
  runTestMetrics, 18

  simPerformance, 19

  wilcox.test, 4