Package ‘RchyOptimyx’

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Type Package
Title Optimized Cellular Hierarchies for Flow Cytometry
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Description Constructs a hierarchy of cells using flow cytometry for maximization of an external variable (e.g., a clinical outcome or a cytokine response).
Imports Rgraphviz, sfsmisc, graphics, methods, graph, grDevices, flowType (>= 2.0.0)
Depends R (>= 2.10)
Suggests flowCore
biocViews FlowCytometry
License Artistic-2.0
LazyLoad yes
NeedsCompilation yes

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Description

RchyOptimyx is a computational tool which determines the minimal sets of markers that can identify a target population within a given purity standard. This can enable panel redesign using smaller subsets of markers, with consequent savings on reagents, potential for adaptation to older instruments, or the addition of other markers of interest to the panel design.

Details

| Package:  | RchyOptimyx          |
| Type:     | Package              |
| Version:  | 0.99.1               |
| Date:     | 2011-10-12           |
| License:  | Artistic-2.0         |
| LazyLoad: | yes                  |

See the `RchyOptimyx` function.

Author(s)

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References


See Also

RchyOptimyx

Examples

data(HIVData)

res<-RchyOptimyx(PhenoCodes, -log10(LogRankPvals), '2000102001', 5,FALSE)
plot(res,phenotypeCodes=PhenoCodes,phenotypeScores=-log10(LogRankPvals),marker.names=marker.names)

HIVData

Description

A dataset of two sets of scores (particularly, correlation with protection against HIV and overlap with the Naive T-cell population) assigned to immunophenotypes measured by flow cytometry. 10 markers were measured: KI-67, CD28, CD45RO, CD8, CD4, CD57, CCR5, CD27, CCR7, and CD127.
merge

Usage
data(HIVData)

Details

This dataset consists of 2 numeric (LogRankPvals and OverlapScores) and 2 character (Pheno.Codes and marker.names).

Pheno.Codes vector is described in the RchyOptimyx function’s manual.

LogRankPvals is a vector of log-rank test P-values to determine the correlation between HIV’s progression and each of the measured immunophenotypes in 466 HIV positive patients (lower values represent a stronger correlation). For more details see the Aghaeepour et.al. manuscript below.

Ganesan et. al. define Naive T-cells as CD28+CD45RO-CD57-CCR5-CD27+ CCR7+ within the CD3+CD14- compartment. The OverlapScores vector has the proportion of Naive T-cells (as defined above) to the total number of cells in any given immunophenotype (a higher value represents a larger overlap).

marker.names is the name of all markers involved in the analysis.

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References


merge  Merges two OptimizedHierarchy objects.

Description

Merges two OptimizedHierarchy objects.

Usage

## S4 method for signature 'OptimizedHierarchy,OptimizedHierarchy'
merge(x, y)

Arguments

x  First OptimizedHierarchy.

y  Second OptimizedHierarchy.
Value

OptH  An OptimizedHierarchy object

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plot

Plots an optimized cellular hierarchy

Description

Plots an optimized cellular hierarchy

Usage

```r
## S4 method for signature 'OptimizedHierarchy,ANY'
plot(x, phenotypeScores, phenotypeCodes, marker.names,
     partitions.per.marker = NULL,
     uniformColors = FALSE, ylab = NULL, xlab = NULL, colors = c("blue",
     "cyan", "yellow", "red"), edgeWeights = TRUE, edgeLabels = TRUE,
     nodeLabels = TRUE, min.score = NA, max.score = NA, cell.proportions =
     NULL, min.proportion = NA, max.proportion = NA, proportion.colors =
     c("black", "white"), node.lwd = 5, root.name = "All Cells",
     legend.size=1.25, plot.legend=TRUE, textcol=par(fg'))
```

Arguments

- **x**: An OptimizedHierarchy object as generated by RchyOptimy.
- **phenotypeScores**: The score vector used for determining the colors of the nodes.
- **phenotypeCodes**: A vector of strings of length N (the number of markers) for each phenotype measured. For every phenotype, the character corresponding to a given marker can be 0, 1, 2, etc for neutral, negative, positive, bright, etc. See the provided vignette for more details and examples.
- **marker.names**: A vector of characters representing the names of all markers involved in the analysis.
- **partitions.per.marker**: A vector of integers, one per marker, describing the number of partitions per marker. Default is 2 (positive and negative) for all markers.
- **uniformColors**: Boolean variable. If TRUE, the graph will not have any colors.
- **ylab**: y-axis label of the density plot on the right.
- **xlab**: x-axis label of the density plot of the phenotype cell proportion in the bottom.
- **colors**: Color vector indicating colors to be used in the right pallet.
- **edgeWeights**: Boolean value indicating weather edgeWeights should be plotted or not.
- **edgeLabels**: Boolean value indicating weather edgeLabels should be plotted or not.
- **nodeLabels**: Boolean value indicating weather nodeLabels should be plotted or not.
**RchyOptimyx**

<table>
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<th>Variable</th>
<th>Description</th>
</tr>
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<tbody>
<tr>
<td>min.score</td>
<td>Double value indicating the minimum value of scores. Default value will use minimum node scores of the given OptimizedHierarchy object.</td>
</tr>
<tr>
<td>max.score</td>
<td>Double value indicating the maximum value of scores. Default value will use maximum node scores of the given OptimizedHierarchy object.</td>
</tr>
<tr>
<td>cell.proportions</td>
<td>The score vector used to determine the color of node borders to illustrate phenotype cell proportions.</td>
</tr>
<tr>
<td>min.proportion</td>
<td>Double value indicating the minimum value of the cell proportions. Default value will use minimum cell proportion of the given OptimizedHierarchy object.</td>
</tr>
<tr>
<td>max.proportion</td>
<td>Double value indicating the maximum value of the cell proportions. Default value will use minimum cell proportion of the given OptimizedHierarchy object.</td>
</tr>
<tr>
<td>proportion.colors</td>
<td>Color vector indicating colors to be used in the bottom pallet.</td>
</tr>
<tr>
<td>node.lwd</td>
<td>Integer value indicating node border width.</td>
</tr>
<tr>
<td>root.name</td>
<td>Character value indicating the root node’s name.</td>
</tr>
<tr>
<td>legend.size</td>
<td>Size of the color legend in inches.</td>
</tr>
<tr>
<td>plot.legend</td>
<td>Boolean indicating whether the color legend should be plotted.</td>
</tr>
<tr>
<td>textcol</td>
<td>The color of the axis label(s) for the color legend(s).</td>
</tr>
</tbody>
</table>

**Author(s)**

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**See Also**

*RchyOptimyx-methods*

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**RchyOptimyx**

*Gating Hierarchy Optimization for Flow Cytometry*

**Description**

RchyOptimyx is a computational tool which determines the minimal sets of markers that can identify a target population within a given purity standard. This can enable panel redesign using smaller subsets of markers, with consequent savings on reagents, potential for adaptation to older instruments, or the addition of other markers of interest to the panel design.

**Usage**

```r
## S4 method for signature 'character,numeric,character,numeric,logical'
RchyOptimyx(pheno.codes, phenotypeScores, startPhenotype, pathCount=1, trimPaths=FALSE, trim.tolerance=0)
```
Arguments

pheno.codes  A vector of strings of length N (the number of markers) for each phenotype measured. For every phenotype, the character corresponding to a given marker can be 0, 1, 2, etc for neutral, negative, positive, bright, etc. See the provided vignette (inst/doc/RchyOptimyx.pdf) for more details and examples.

phenotypeScores  A vector of the scores assigned to every phenotype. The optimization algorithm will try to maximize this score. See the provided vignette (inst/doc/RchyOptimyx.pdf) for more details and examples.

startPhenotype  The terminal phenotype which includes every marker that must be considered in the analysis. This variable is a vector of the length of the number of markers. Every element of the vector can be 0, 1, or 2 for negative, neutral, or positive respectively (see details).

pathCount  The number of paths that must be generated.

trimPaths  A boolean indicating whether the paths should be trimmed or not. If set to TRUE, the paths will be terminated as soon as adding trim.tolerance extra number of edges to the path doesn’t result in an improvement in the score.

trim.tolerance  An integer indicating the number of levels before the path is trimmed if the score is decreasing by adding extra edges.

trim.level  An integer indicating from which level the tree is going to be trimmed. This parameter has no effect if set to zero.

Details

Every marker can be positive, negative, or neutral. Neutral markers are not included in the measurement of the respective phenotype.

A path is trimmed in one of the following conditions: 1. If the trimPaths parameter is TRUE, the path is trimmed as soon as it decreases for equal or more levels that trim.tolerance parameter. If the path is decreasing while reaching the trim.level, the path is trimmed from the point that it started to decrease. 2. If the path reaches the trim.level.

Value

OptH  An OptimizedHierarchy object

Author(s)

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References


Examples

data(HIVData)

res<-RchyOptimyx(PhenoCodes, -log10(LogRankPvals), '2000102001', 5,FALSE)
plot(res,phenotypeCodes=PhenoCodes,phenotypeScores=-log10(LogRankPvals),marker.names=marker.names)

summary-methods

Description

~~ Methods for function summary ~~

Methods

signature(object = "OptimizedHierarchy") Prints basic characteristics of an OptimizedHierarchy object.

Author(s)

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See Also

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