Package ‘cytometree’

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Type Package

Title Automated Cytometry Gating and Annotation

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LinkingTo Rcpp, RcppArmadillo

Description Given the hypothesis of a bi-modal distribution of cells for each marker, the algorithm constructs a binary tree, the nodes of which are subpopulations of cells. At each node, observed cells and markers are modeled by both a family of normal distributions and a family of bi-modal normal mixture distributions. Splitting is done according to a normalized difference of AIC between the two families. Method is detailed in: Commenges, Alkhassim, Gottardo, Hejblum & Thiebaut (2018) <doi: 10.1002/cyto.a.23601>.

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LazyData true

Depends R (>= 3.1.0), Rcpp (>= 0.12.11)

Imports ggplot2, graphics, igraph, mclust, methods, stats, cowplot, GoFKernel

RoxygenNote 7.0.2

BugReports https://github.com/sistm/Cytometree/issues

Encoding UTF-8

Suggests knitr, formatR, rmarkdown, viridis

VignetteBuilder knitr

NeedsCompilation yes

Repository CRAN

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cytometree-package

R topics documented:

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| cytomtree-package | Binary tree algorithm for cytometry data analysis. |

Description

The algorithm is based on the construction of a binary tree, the nodes of which are subpopulations of cells. At each node, observed cells and markers are modeled by both a family of normal distributions and a family of bi-modal normal mixture distributions. Splitting is done according to a normalized difference of AIC between the two families. Given the unsupervised nature of the binary tree, some of the available markers may not be used to find the different cell populations present in a given sample. To recover a complete annotation, we defined, as a post processing procedure, an annotation method which allows the user to distinguish two or three expression levels per marker.

Details

| Package: | cytometree |
| Type:     | Package    |
| Version:  | 2.0.2      |
| Date:     | 2019-12-04 |
| License:  | LGPL-3     |

The main function in this package is CytomeTree.

Author(s)

Chariff Alkhassim, Boris Hejblum, Anthony Devaux, Van Hung Huynh Tran — Maintainer: Boris Hejblum
Annotation

References


Annotation

Annotates cell populations found using CytomeTree.

Description

Annotates cell populations found using CytomeTree.

Usage

Annotation(
  CytomeTreeObj,
  K2markers = NULL,
  K3markers = NULL,
  plot = TRUE,
  t = 0.2,
  remove_outliers_inplot = TRUE,
  center_fun = c("median", "mean")
)

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>CytomeTreeObj</td>
<td>An object of class CytomeTree.</td>
</tr>
<tr>
<td>K2markers</td>
<td>A vector of class character where the names of the markers for which 2 levels of expression are sought can be specified. Default is NULL i.e. unsupervised.</td>
</tr>
<tr>
<td>K3markers</td>
<td>A vector of class character where the names of the markers for which 3 levels of expression are sought can be specified. Default is NULL i.e. unsupervised.</td>
</tr>
<tr>
<td>plot</td>
<td>A logical value indicating whether or not to plot the partitioning in 1, 2 or 3 groups for each marker. Default is TRUE.</td>
</tr>
<tr>
<td>t</td>
<td>A real positive-or-null number used for comparison with the normalized AIC computed to compare the fits of the marginal distributions obtained by one normal distribution and by a mixture of two or three normal. For markers used in the tree, the algorithm compares the fits obtained by a mixture of two and three normal distributions. Default value is .2. A higher value leads to a smaller number of expression levels per marker.</td>
</tr>
<tr>
<td>remove_outliers_inplot</td>
<td>a logical flag indicating whether the y-axis should be scaled by removing outliers or not. Default is TRUE.</td>
</tr>
<tr>
<td>center_fun</td>
<td>a character string either 'median' or 'mean' indicating based on which summary the populations should be ordered. Default is 'median', which is more robust to outliers and long tail distributions.</td>
</tr>
</tbody>
</table>
Details

The algorithm is set to find the partitioning in 1, 2 or 3 groups of cell populations found using Cy-
tomeTree. In an unsupervised mode, it minimizes the within-leaves sum of squares of the observed
values on each marker and computes the normalized AIC to compare the fits of the marginal distrib-
utions obtained by one normal distribution and by a mixture of two or three normal. For markers
used in the tree, the algorithm compares the fits obtained by a mixture of two and three normal
distributions.

Value

A data.frame containing the annotation of each cell population.

Author(s)

Chariff Alkhassim, Boris Hejblum

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**bootstrapCI**

*Bootstrapped Confidence Interval.*

**Description**

Bootstrapped Confidence Interval.

**Usage**

`bootstrapCI(stat, n, alpha)`

**Arguments**

- `stat`  A numeric vector of statistics for which to compute a confidence interval.
- `n` An integer giving the number of bootstrap samples.
- `alpha` A real number comprised in ]0, 1[ - desired confidence level.

**Author(s)**

Chariff Alkhassim
CytofTree

Binary tree algorithm for mass cytometry data analysis.

Description

Binary tree algorithm for mass cytometry data analysis.

Usage

CytofTree(
  M,
  minleaf = 1,
  t = 0.1,
  verbose = TRUE,
  force_first_markers = NULL,
  transformation = c("asinh", "biexp", "log10", "none"),
  num_col = 1:ncol(M)
)

Arguments

M A matrix of size n x p containing mass cytometry measures of n cells on p markers.
minleaf An integer indicating the minimum number of cells per population. Default is 1.
t A real positive-or-null number used for comparison with the normalized AIC computed at each node of the tree. A higher value limits the height of the tree.
verbose A logical controlling if a text progress bar is displayed during the execution of the algorithm. By default is TRUE.
force_first_markers a vector of index to split the data on first. This argument is used in the semi-supervised setting, forcing the algorithm to consider those markers first, in the order they appear in this force_first_markers vector, and forcing the split at every node. Default is NULL, in which case the clustering algorithm is unsupervised.
transformation A string indicating the transformation used among asinh biexp, log10 and none. Default is asinh transformation.
num_col An integer vector of index indicating the columns to be transform. Default is 1:ncol(M) to transform all the data.

Details

First of all, data can be transformed using different transformations. The algorithm is based on the construction of a binary tree, the nodes of which are subpopulations of cells. At each node, observed cells and markers are modeled by both a family of normal distributions and a family of bi-modal normal mixture distributions. Splitting is done according to a normalized difference of AIC between the two families.
Value

An object of class 'cytomeTree' providing a partitioning of the set of n cells.

- annotation A data.frame containing the annotation of each cell population underlying the tree pattern.
- labels The partitioning of the set of n cells.
- M The transformed matrix of mass cytometry.
- mark_tree A two level list containing markers used for node splitting.
- transformation Transformation used
- num_col Indexes of columns transformed

Author(s)

Anthony Devaux, Boris Hejblum

Examples

data(IMdata)

# dimension of data
dim(IMdata)

# given the size of the dataset, the code below can take several minutes to run

if(interactive()){
  # Don't transform Time et Cell_length column
  num_col <- 3:ncol(IMdata)

  # Build Cytoftree binary tree
  tree <- CytofTree(M = IMdata, minleaf = 1, t = 0.1, transformation = "asinh", num_col = num_col)

  # Annotation
  annot <- Annotation(tree, plot = FALSE, K2markers = colnames(IMdata))

  # Provide subpopulations
  annot$combinations
}

CytomeTree  Binary tree algorithm for cytometry data analysis.

Description

Binary tree algorithm for cytometry data analysis.
Usage

CytomeTree(M, minleaf = 1, t = 0.1, verbose = TRUE, force_first_markers = NULL)

Arguments

M A matrix of size n x p containing cytometry measures of n cells on p markers.

minleaf An integer indicating the minimum number of cells per population. Default is 1.

t A real positive-or-null number used for comparison with the normalized AIC computed at each node of the tree. A higher value limits the height of the tree.

verbose A logical controlling if a text progress bar is displayed during the execution of the algorithm. By default is TRUE.

force_first_markers a vector of index to split the data on first. This argument is used in the semi-supervised setting, forcing the algorithm to consider those markers first, in the order they appear in this force_first_markers vector, and forcing the split at every node. Default is NULL, in which case the clustering algorithm is unsupervised.

Details

The algorithm is based on the construction of a binary tree, the nodes of which are subpopulations of cells. At each node, observed cells and markers are modeled by both a family of normal distributions and a family of bi-modal normal mixture distributions. Splitting is done according to a normalized difference of AIC between the two families.

Value

An object of class ‘cytomeTree’ providing a partitioning of the set of n cells.

• annotation A data.frame containing the annotation of each cell population underlying the tree pattern.
• labels The partitioning of the set of n cells.
• M The input matrix.
• mark_tree A two level list containing markers used for node splitting.

Author(s)

Chariff Alkhassim, Boris Hejblum

Examples

head(DLBCL)

# number of cell event
N <- nrow(DLBCL)

# Cell events
cellevents <- DLBCL[, c("FL1", "FL2", "FL4")]

# Manual partitioning of the set N (from FlowCAP-I)
manual_labels <- DLBCL[, "label"]

# Build the binary tree
Tree <- CytomeTree(cellevents, minleaf = 1, t=.1)

# Retreive the resulting partition of the set N
Tree_Partition <- Tree$labels

# Plot node distributions
par(mfrow=c(1, 2))
plot_nodes(Tree)

# Choose a node to plot
plot_nodes(Tree,"FL4.1")

# Plot a graph of the tree
par(mfrow=c(1,1))
plot_graph(Tree,edge.arrow.size=.3, Vcex = .5, vertex.size = 30)

# Run the annotation algorithm
Annot <- Annotation(Tree,plot=FALSE)
Annot$combinations

# Compare to the annotation gotten from the tree
Tree$annotation

# Example of sought phenotypes
# Variable in which sought phenotypes can be entered in the form of matrices.
phenotypes <- list()

# Sought phenotypes:
## FL2+ FL4-.
phenotypes[[1]] <- rbind(c("FL2", 1), c("FL4", 0))

## FL2- FL4+.
phenotypes[[2]] <- rbind(c("FL2", 0), c("FL4", 1))

## FL2+ FL4+.
phenotypes[[3]] <- rbind(c("FL2", 1), c("FL4", 1))

# Retreive cell populations found using Annotation.
PhenoInfos <- RetrievePops(Annot, phenotypes)
PhenoInfos$phenotypesinfo

# F-measure ignoring cells labeled 0 as in FlowCAP-I.
# Use FmeasureC() in any other case.
FmeasureC_no0(ref=manual_labels, pred=Tree_Partition)

if(interactive()){
  # Scatterplots.
  library(ggplot2)
  # Ignoring cells labeled 0 as in FlowCAP-I.
  rm_zeros <- which(!manual_labels)
  # Building the data frame to scatter plot the data.
  FL1 <- cellevents[-c(rm_zeros),"FL1"]
  FL2 <- cellevents[-c(rm_zeros),"FL2"]
  FL4 <- cellevents[-c(rm_zeros),"FL4"]
  n <- length(FL1)
  Labels <- c(manual_labels[-c(rm_zeros)]%%2+1, Tree_Partition[-c(rm_zeros)])
  Labels <- as.factor(Labels)
  method <- as.factor(c(rep("FlowCap-I",n),rep("CytomeTree",n)))
  scatter_df <- data.frame("FL2" = FL2, "FL4" = FL4, "labels" = Labels, "method" = method)
  p <- ggplot2::ggplot(scatter_df, ggplot2::aes_string(x = "FL2", y = "FL4", colour = "labels")) +
    ggplot2::geom_point(alpha = 1,cex = 1) +
    ggplot2::scale_colour_manual(values = c("green","red","blue")) +
    ggplot2::facet_wrap(~ method) +
    ggplot2::theme_bw() +
    ggplot2::theme(legend.position="bottom")
  p
}

---

**DLBCL**  
*Diffuse large B-cell lymphoma data set from the FlowCAP-I challenge.*

**Description**


**Usage**

data(DLBCL)

**Format**

A data frame with 5524 cell events and 3 markers.
FmeasureC

**Description**
C++ implementation of the F-measure computation

**Usage**
```
FmeasureC(pred, ref)
```

**Arguments**
- `pred` vector of a predicted partition
- `ref` vector of a reference partition

**Author(s)**
Boris Hejblum

FmeasureC_no0

**Description**
Aghaeepour in FlowCAP 1 ignore the reference class labeled "0"

**Usage**
```
FmeasureC_no0(pred, ref)
```

**Arguments**
- `pred` vector of a predicted partition
- `ref` vector of a reference partition

**Author(s)**
Boris Hejblum

Source

http://flowcap.flowsite.org/
HIPC T cell data set from HIPC program, patient 12828. The data was analyzed and gated by Stanford.

Description

HIPC T cell data set from HIPC program, patient 12828. The data was analyzed and gated by Stanford.

Usage

data(HIPC)

Format

A data frame with 33992 cell events and 6 markers.

Source


IMdata

Influenza vaccine response dataset

Description

A dataset containing 10,000 cells and 39 markers of mass cytometry subsampled from the sample SUB116516.478 from the study SDY478 by Mark Davis retrieved from ImmuneSpace

Usage

data(IMdata)

Format

A data frame with 10,000 rows and 39 variables:

Source

https://www.immunespace.org/project/Studies/SDY478/begin.view?
plot_cytopop

Plot the cell count for each population using CytomeTree.

Description
Plot the cell count for each population using CytomeTree.

Usage
plot_cytopop(
  AnnotObj,
  nbpop = 10,
  mincount = 1,
  maxcount = NULL,
  y_axis = c("abs_count", "prop")
)

Arguments
AnnotObj: An object of class Annotation.

nbpop: Number indicating the maximum of population plotted. Default is 10

mincount: Number indicating the minimum of cell count for the populations. Default is 1.

maxcount: Number indicating the maximum of cell count for the populations. Default is NULL i.e no maximum selected.

y_axis: a character string either "abs_count" or "prop" indicating whether the absolute cell count or the relative populations proportions should be plotted. Default is "abs_count".

Author(s)
Anthony Devaux, Boris Hejblum

Examples
# Run CytomeTree
data(DLBCL)
cellevents <- DLBCL[,c("FL1", "FL2", "FL4")]
Tree <- CytomeTree(cellevents, minleaf = 1, t=.1)
Annot <- Annotation(Tree,plot=FALSE)

# Plot the cell count
plot_cytopop(Annot)
**plot_graph**

*Plot the binary tree built using CytomeTree.*

**Description**

Plot the binary tree built using CytomeTree.

**Usage**

```r
plot_graph(CytomeTreeObj, Ecex = 1, Ecolor = 8, Vcex = 0.8, Vcolor = 0, ...)```

**Arguments**

- **CytomeTreeObj**
  An object of class CytomeTree.
- **Ecex**
  Number indicating the amount by which text on the edges should be scaled. Default is 1.
- **Ecolor**
  An integer or a string of character to color edges of the graph. Default is 8.
- **Vcex**
  Number indicating the amount by which text in the vertices should be scaled. Default is 0.8.
- **Vcolor**
  A vector of class numeric or character to color vertices of the graph. Default is 0.
- **...**
  Additional arguments to be passed to `plot_graph`.

**Author(s)**

Chariff Alkhassim

---

**plot_nodes**

*Plot the distribution of the observed cells at each node of the binary tree built using CytomeTree.*

**Description**

Plot the distribution of the observed cells at each node of the binary tree built using CytomeTree.

**Usage**

```r
plot_nodes(
  CytomeTreeObj,
  nodes = NULL,
  nodesPerCol = NULL,
  nodesPerRow = NULL,
  ...
)
```
RetrievePops

Arguments

CytomeTreeObj  An object of class CytomeTree.

nodes  A list of character elements containing the name of the nodes for which the
distribution is to be plotted. Default is NULL, and plots the distribution for each
node.

nodesPerCol  an integer specifying the number of plots to be displayed per column when plot-
ting multiple nodes at once. Default is NULL.

nodesPerRow  an integer specifying the number of plots to be displayed per row when plotting
multiple nodes at once. Default is NULL.

...  further arguments to be passed to plot_grid.

Details

if both nodesPerCol and nodesPerRow are NULL then all the nodes are plotted on a single page.

Value

a list of ggplot2 plot objects, containing each node plot.

Author(s)

Chariff Alkhassim, Boris Hejblum

Examples

data(DLBCL)
myct <- CytomeTree(DLBCL[, c("FL1", "FL2", "FL4")], minleaf = 1, t=.1)
plot_nodes(myclt)

---

RetrievePops  Retrieve cell populations found using Annotation.

Description

Retrieve cell populations found using Annotation.

Usage

RetrievePops(AnnotationObj, phenotypes)
Arguments

AnnotationObj An object of class Annotation.

phenotypes A list containing at least one element of class matrix describing a sought phenotype. Each matrix should have two columns where the name of a used marker is associated to a value chosen between 0, 1 and 2. 0 translates to -, 1 to + and 2 to ++.

Value

A list of two elements.

- phenotypesinfo A list containing informations about sought populations.
- Mergedleaves The partitioning of the set of n cells with potentially merged leaves.

Author(s)

Chariff Alkhassim, Boris Hejblum
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