

Package ‘CytoGLMM’

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Type Package

Title Conditional Differential Analysis for Flow and Mass Cytometry Experiments

Version 1.4.0

Description The CytoGLMM R package implements two multiple regression strategies: A bootstrapped generalized linear model (GLM) and a generalized linear mixed model (GLMM). Most current data analysis tools compare expressions across many computationally discovered cell types. CytoGLMM focuses on just one cell type. Our narrower field of application allows us to define a more specific statistical model with easier to control statistical guarantees. As a result, CytoGLMM finds differential proteins in flow and mass cytometry data while reducing biases arising from marker correlations and safeguarding against false discoveries induced by patient heterogeneity.

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URL <https://christofseiler.github.io/CytoGLMM>,
<https://github.com/ChristofSeiler/CytoGLMM>

BugReports <https://github.com/ChristofSeiler/CytoGLMM/issues>

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LazyData true

Imports stats, methods, BiocParallel, RColorBrewer, cowplot,
doParallel, dplyr, factoextra, flexmix, ggplot2, magrittr,
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Author Christof Seiler [aut, cre] (<<https://orcid.org/0000-0001-8802-3642>>)

Maintainer Christof Seiler <christof.seiler@maastrichtuniversity.nl>

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Description

Logistic mixture regression

Usage

```
cytoflexmix(  
  df_samples_subset,  
  protein_names,  
  condition,  
  group = "donor",  
  cell_n_min = Inf,  
  cell_n_subsample = 0,  
  ks = seq_len(10),  
  num_cores = 1  
)
```

Arguments

df_samples_subset	Data frame or tibble with proteins counts, cell condition, and group information
protein_names	A vector of column names of protein to use in the analysis
condition	The column name of the condition variable
group	The column name of the group variable
cell_n_min	Remove samples that are below this cell counts threshold
cell_n_subsample	Subsample samples to have this maximum cell count
ks	A vector of cluster sizes
num_cores	Number of computing cores

Value

A list of class `cytglm` containing

flexmixfits	list of <code>flexmix</code> objects
df_samples_subset	possibly subsampled df_samples_subset table
protein_names	input protein names
condition	input condition variable
group	input group names
cell_n_min	input cell_n_min
cell_n_subsample	input cell_n_subsample
ks	input ks
num_cores	input num_cores

Examples

```
set.seed(23)
df <- generate_data()
protein_names <- names(df)[3:12]
df <- dplyr::mutate_at(df, protein_names, function(x) asinh(x/5))
mix_fit <- CytoGLMM::cytologlm(df,
                                  protein_names = protein_names,
                                  condition = "condition",
                                  group = "donor",
                                  ks = 2)
mix_fit
```

cytoglm

Fit GLM with bootstrap resampling

Description

Fit GLM with bootstrap resampling

Usage

```
cytoglm(
  df_samples_subset,
  protein_names,
  condition,
  group = "donor",
  covariate_names = NULL,
  cell_n_min = Inf,
  cell_n_subsample = 0,
  num_boot = 100,
  num_cores = 1
)
```

Arguments

df_samples_subset	Data frame or tibble with proteins counts, cell condition, and group information
protein_names	A vector of column names of protein to use in the analysis
condition	The column name of the condition variable
group	The column name of the group variable
covariate_names	The column names of covariates
cell_n_min	Remove samples that are below this cell counts threshold
cell_n_subsample	Subsample samples to have this maximum cell count
num_boot	Number of bootstrap samples
num_cores	Number of computing cores

Value

A list of class `cytogilmm` containing

<code>tb_coef</code>	coefficent table
<code>df_samples_subset</code>	possibly subsampled <code>df_samples_subset</code> table
<code>protein_names</code>	input protein names
<code>condition</code>	input condition variable
<code>group</code>	input group names
<code>covariate_names</code>	input covariates
<code>cell_n_min</code>	input <code>cell_n_min</code>
<code>cell_n_subsample</code>	input <code>cell_n_subsample</code>
<code>unpaired</code>	true if unpaired samples were provided as input
<code>num_boot</code>	input <code>num_boot</code>
<code>num_cores</code>	input <code>num_cores</code>
<code>formula_str</code>	formula use in the regression model

Examples

```
set.seed(23)
df <- generate_data()
protein_names <- names(df)[3:12]
df <- dplyr::mutate_at(df, protein_names, function(x) asinh(x/5))
glm_fit <- CytoGLMM::cytogilmm(df,
                                    protein_names = protein_names,
                                    condition = "condition",
                                    group = "donor",
                                    num_boot = 10) # in practice >=1000
glm_fit
```

Description

Fit GLMM with method of moments

Usage

```
cytoglmm(
  df_samples_subset,
  protein_names,
  condition,
  group = "donor",
  covariate_names = NULL,
  cell_n_min = Inf,
  cell_n_subsample = 0,
  num_cores = 1
)
```

Arguments

<code>df_samples_subset</code>	Data frame or tibble with proteins counts, cell condition, and group information
<code>protein_names</code>	A vector of column names of protein to use in the analysis
<code>condition</code>	The column name of the condition variable
<code>group</code>	The column name of the group variable
<code>covariate_names</code>	The column names of covariates
<code>cell_n_min</code>	Remove samples that are below this cell counts threshold
<code>cell_n_subsample</code>	Subsample samples to have this maximum cell count
<code>num_cores</code>	Number of computing cores

Value

A list of class `cytoglm` containing

<code>glmmfit</code>	<code>mbest</code> object
<code>df_samples_subset</code>	possibly subsampled <code>df_samples_subset</code> table
<code>protein_names</code>	input protein names
<code>condition</code>	input condition variable
<code>group</code>	input group names
<code>covariate_names</code>	input covariates
<code>cell_n_min</code>	input <code>cell_n_min</code>
<code>cell_n_subsample</code>	input <code>cell_n_subsample</code>
<code>num_cores</code>	input <code>num_cores</code>

Examples

```
set.seed(23)
df <- generate_data()
protein_names <- names(df)[3:12]
df <- dplyr::mutate_at(df, protein_names, function(x) asinh(x/5))
glmm_fit <- CytoGLMM::cytoglmm(df,
                                    protein_names = protein_names,
                                    condition = "condition",
                                    group = "donor")
glmm_fit
```

cytogroup

Group-specific fixed effects model

Description

Group-specific fixed effects model

Usage

```
cytogroup(
  df_samples_subset,
  protein_names,
  condition,
  group = "donor",
  cell_n_min = Inf,
  cell_n_subsample = 0
)
```

Arguments

df_samples_subset	Data frame or tibble with proteins counts, cell condition, and group information
protein_names	A vector of column names of protein to use in the analysis
condition	The column name of the condition variable
group	The column name of the group variable
cell_n_min	Remove samples that are below this cell counts threshold
cell_n_subsample	Subsample samples to have this maximum cell count

Value

A list of class `cytoglm` containing

groupfit	<code>speedglm</code> object
df_samples_subset	possibly subsampled df_samples_subset table

```

protein_names    input protein names
condition        input condition variable
group            input group names
cell_n_min       input cell_n_min
cell_n_subsample input cell_n_subsample

```

Examples

```

set.seed(23)
df <- generate_data()
protein_names <- names(df)[3:12]
df <- dplyr::mutate_at(df, protein_names, function(x) asinh(x/5))
group_fit <- CytoGLMM::cytogroup(df,
                                    protein_names = protein_names,
                                    condition = "condition",
                                    group = "donor")
group_fit

```

cytostab

Evaluate parameter stability with respect to gating sheme

Description

Evaluate parameter stability with respect to gating sheme

Usage

```

cytostab(
  df_samples_subset,
  protein_names,
  condition,
  group = "donor",
  cell_n_min = Inf,
  cell_n_subsample = 0
)

```

Arguments

df_samples_subset	Data frame or tibble with proteins counts, cell condition, and group information
protein_names	A vector of column names of protein to use in the analysis
condition	The column name of the condition variable
group	The column name of the group variable
cell_n_min	Remove samples that are below this cell counts threshold
cell_n_subsample	Subsample samples to have this maximum cell count

Value

A data frame

Examples

```
set.seed(23)
df <- generate_data()
protein_names <- names(df)[3:12]
df <- dplyr::mutate_at(df, protein_names, function(x) asinh(x/5))
stab <- CytoGLMM::cytostab(df,
                           protein_names = protein_names,
                           condition = "condition",
                           group = "donor")
stab
```

cyto_check

Check if input to cytoxxx function have errors

Description

Check if input to cytoxxx function have errors

Usage

```
cyto_check(cell_n_subsample, cell_n_min, protein_names)
```

Arguments

cell_n_subsample

Subsample samples to have this maximum cell count

cell_n_min

A vector of column names of protein to use in the analysis

protein_names

A vector of column names of protein to use in the analysis

Value

NULL.

`generate_data`*Generate dataset for vignettes and simulation studies***Description**

Generate dataset for vignettes and simulation studies

Usage

```
generate_data()
```

Value

`tibble` data frame

Examples

```
set.seed(23)
df <- generate_data()
str(df)
df
```

`glmm_moment`*Generalized linear mixed model with maximum likelihood***Description**

Generalized linear mixed model with maximum likelihood

Usage

```
glmm_moment(
  df_samples,
  protein_names,
  response,
  group = "donor",
  covariate_names = NULL,
  num_cores = 1
)
```

Arguments

df_samples	Data frame or tibble with proteins counts, cell condition, and group information
protein_names	A vector of column names of protein to use in the analysis
response	The column name of the condition variable
group	The column name of the group variable
covariate_names	The column names of covariates
num_cores	Number of computing cores

Value

`mbest` object

is_unpaired	<i>Check if samples match or paired on condition</i>
-------------	--

Description

Check if samples match or paired on condition

Usage

```
is_unpaired(df_samples_subset, condition, group)
```

Arguments

df_samples_subset	Data frame or tibble with proteins counts, cell condition, and group information
condition	The column name of the condition variable
group	The column name of the group variable

Value

A boolean

plot.cytoflexmix *Plot all components of mixture regression*

Description

Plot all components of mixture regression

Usage

```
## S3 method for class 'cytoflexmix'
plot(x, k = NULL, separate = FALSE, ...)
```

Arguments

x	A cytoflexmix class
k	Number of clusters
separate	create two separate ggplot2 objects
...	Other parameters

Value

[ggplot2](#) object

Examples

```
set.seed(23)
df <- generate_data()
protein_names <- names(df)[3:12]
df <- dplyr::mutate_at(df, protein_names, function(x) asinh(x/5))
mix_fit <- CytoGLMM:cytoflexmix(df,
                                   protein_names = protein_names,
                                   condition = "condition",
                                   group = "donor",
                                   ks = 2)
plot(mix_fit)
```

plot.cytoglm *Plot bootstraped coefficients*

Description

Plot bootstraped coefficients

Usage

```
## S3 method for class 'cytoglm'
plot(x, order = FALSE, separate = FALSE, ...)
```

Arguments

x	A <code>cytogi^m</code> class
order	Order the markers according to the magnitude of the coefficients
separate	create two separate <code>ggplot2</code> objects
...	Other parameters

Value

`ggplot2` object

Examples

```
set.seed(23)
df <- generate_data()
protein_names <- names(df)[3:12]
df <- dplyr::mutate_at(df, protein_names, function(x) asinh(x/5))
glm_fit <- CytoGLMM::cytogim(df,
                                    protein_names = protein_names,
                                    condition = "condition",
                                    group = "donor",
                                    num_boot = 10) # in practice >=1000
plot(glm_fit)
```

`plot.cytoglm`

Plot fixed coefficients of random effects model

Description

Plot fixed coefficients of random effects model

Usage

```
## S3 method for class 'cytogim'
plot(x, order = FALSE, separate = FALSE, ...)
```

Arguments

x	A <code>cytogi^m</code> class
order	Order the markers according to the magnitude of the coefficients
separate	create two separate <code>ggplot2</code> objects
...	Other parameters

Value

`ggplot2` object

Examples

plot.cytogroup

Plot fixed coefficients of group-specific fixed effects model

Description

Plot fixed coefficients of group-specific fixed effects model

Usage

```
## S3 method for class 'cytogroup'  
plot(x, order = FALSE, separate = FALSE, ...)
```

Arguments

x	A <code>cytoglmM</code> class
order	Order the markers according to the magnitude of the coefficients
separate	create two separate <code>ggplot2</code> objects
...	Other parameters

Value

ggplot2 object

Examples

plot_coeff	<i>Helper function to plot regression coefficient</i>
------------	---

Description

Helper function to plot regression coefficient

Usage

```
plot_coeff(  
  tb,  
  title_str,  
  title_str_right,  
  xlab_str,  
  redline = 0,  
  order = FALSE,  
  separate = FALSE  
)
```

Arguments

tb	A data frame
title_str	Title string for summary plot
title_str_right	Title for bootstrap sample plot
xlab_str	Label on x-axis
redline	Point on x-axis to draw the red line
order	Order the markers according to the magnitude of the coefficients
separate	Plot both summary and bootstrap samples

Value

[ggplot2](#) object or list of two objects if separate is true

plot_heatmap	<i>Heatmap of median marker expression</i>
--------------	--

Description

Heatmap of median marker expression

Usage

```
plot_heatmap(
  df_samples,
  sample_info_names,
  protein_names,
  arrange_by_1,
  arrange_by_2 = "",
  cluster_cols = FALSE,
  fun = median
)
```

Arguments

<code>df_samples</code>	Data frame or tibble with proteins counts, cell condition, and group information
<code>sample_info_names</code>	Column names that contain information about the cell, e.g. donor, condition, file name, or cell type
<code>protein_names</code>	A vector of column names of protein to use in the analysis
<code>arrange_by_1</code>	Column name
<code>arrange_by_2</code>	Column name
<code>cluster_cols</code>	Apply hierarchical cluster to columns
<code>fun</code>	Summary statistics of marker expression

Value

[phewmat](#) object

Examples

```
set.seed(23)
df <- generate_data()
protein_names <- names(df)[3:12]
df <- dplyr::mutate_at(df, protein_names, function(x) asinh(x/5))
CytoGLMM::plot_heatmap(df,
  protein_names = protein_names,
  sample_info_names = c("donor", "condition"),
  arrange_by_1 = "condition")
```

plot_lda

LDA on marker expression

Description

LDA on marker expression

Usage

```
plot_lda(
  df_samples,
  protein_names,
  group,
  cor_scaling_factor = 1,
  arrow_color = "black",
  marker_color = "black",
  marker_size = 5
)
```

Arguments

<code>df_samples</code>	Data frame or tibble with proteins counts, cell condition, and group information
<code>protein_names</code>	A vector of column names of protein to use in the analysis
<code>group</code>	The column name of the group variable
<code>cor_scaling_factor</code>	Scaling factor of circle of correlations
<code>arrow_color</code>	Color of correlation circle
<code>marker_color</code>	Colors of marker names
<code>marker_size</code>	Size of markerr names

Value

`ggplot2` object

Examples

```
set.seed(23)
df <- generate_data()
protein_names <- names(df)[3:12]
df <- dplyr::mutate_at(df, protein_names, function(x) asinh(x/5))
df$condition <- rep(c("A", "B", "C", "D"), each = length(df$condition)/4)
CytoGLMM::plot_lda(df,
  protein_names = protein_names,
  group = "condition",
  cor_scaling_factor = 2)
```

plot_mds

MDS on median marker expression

Description

MDS on median marker expression

Usage

```
plot_mds(
  df_samples,
  protein_names,
  sample_info_names,
  color,
  sample_label = ""
)
```

Arguments

`df_samples` Data frame or tibble with proteins counts, cell condition, and group information
`protein_names` A vector of column names of protein to use in the analysis
`sample_info_names` Column names that contain information about the cell, e.g. donor, condition, file name, or cell type
`color` Column name
`sample_label` Column name

Value

[cowplot](#) object

Examples

```
set.seed(23)
df <- generate_data()
protein_names <- names(df)[3:12]
df <- dplyr::mutate_at(df, protein_names, function(x) asinh(x/5))
CytoGLMM::plot_mds(df,
  protein_names = protein_names,
  sample_info_names = c("donor", "condition"),
  color = "condition")
```

`plot_model_selection` *Plot model selection to choose number optimal number of clusters*

Description

Plot model selection to choose number optimal number of clusters

Usage

```
plot_model_selection(fit, k = NULL)
```

Arguments

fit	A cytoflexmix class
k	Number of clusters

Value

[cowplot](#) object

Examples

```
set.seed(23)
df <- generate_data()
protein_names <- names(df)[3:12]
df <- dplyr::mutate_at(df, protein_names, function(x) asinh(x/5))
mix_fit <- CytoGLMM::cytoflexmix(df,
                                    protein_names = protein_names,
                                    condition = "condition",
                                    group = "donor",
                                    ks = 1:2)
plot_model_selection(mix_fit)
```

plot_prcomp

Plot PCA of subsampled data using ggplot

Description

Plot PCA of subsampled data using ggplot

Usage

```
plot_prcomp(
  df_samples,
  protein_names,
  color_var = "treatment",
  subsample_size = 10000,
  repel = TRUE
)
```

Arguments

df_samples	Data frame or tibble with proteins counts, cell condition, and group information
protein_names	A vector of column names of protein to use in the analysis
color_var	A column name
subsample_size	Subsample per color_var variable
repel	Repel labels

Value

`cowplot` object

Examples

```
set.seed(23)
df <- generate_data()
protein_names <- names(df)[3:12]
df <- dplyr::mutate_at(df, protein_names, function(x) asinh(x/5))
CytoGLMM::plot_prcomp(df,
                      protein_names = protein_names,
                      color_var = "condition")
```

print.cytoglm

Extract and print bootstrap GLM fit

Description

Extract and print bootstrap GLM fit

Usage

```
## S3 method for class 'cytoglm'
print(x, ...)
```

Arguments

x	A <code>cytoglm</code> class
...	Other parameters

Value

NULL.

Examples

```
set.seed(23)
df <- generate_data()
protein_names <- names(df)[3:12]
df <- dplyr::mutate_at(df, protein_names, function(x) asinh(x/5))
glm_fit <- CytoGLMM::cytoglm(df,
                               protein_names = protein_names,
                               condition = "condition",
                               group = "donor",
                               num_boot = 10) # in practice >=1000
print(glm_fit)
```

print.cytoglm	<i>Extract and print GLMM fit</i>
---------------	-----------------------------------

Description

Extract and print GLMM fit

Usage

```
## S3 method for class 'cytoglm'
print(x, ...)
```

Arguments

x	A cytoglm class
...	Other parameters

Value

NULL.

Examples

```
set.seed(23)
df <- generate_data()
protein_names <- names(df)[3:12]
df <- dplyr::mutate_at(df, protein_names, function(x) asinh(x/5))
glmm_fit <- CytoGLMM::cytoglm(df,
                                protein_names = protein_names,
                                condition = "condition",
                                group = "donor")
print(glmm_fit)
```

remove_samples	<i>Remove samples based on low cell counts</i>
----------------	--

Description

Remove samples based on low cell counts

Usage

```
remove_samples(df_samples_subset, condition, group, unpaired, cell_n_min)
```

Arguments

<code>df_samples_subset</code>	Data frame or tibble with proteins counts, cell condition, and group information
<code>condition</code>	The column name of the condition variable
<code>group</code>	The column name of the group variable
<code>unpaired</code>	true if unpaired samples were provided as input
<code>cell_n_min</code>	Remove samples that are below this cell counts threshold

Value

`NULL.`

`summary.cytoglm`

Extract and calculate p-values of bootstrap GLM fit

Description

Extract and calculate p-values of bootstrap GLM fit

Usage

```
## S3 method for class 'cytoglm'
summary(object, method = "BH", ...)
```

Arguments

<code>object</code>	A <code>cytoglm</code> class
<code>method</code>	Multiple comparison adjustment method
<code>...</code>	Other parameters

Value

`tibble` data frame

Examples

```
set.seed(23)
df <- generate_data()
protein_names <- names(df)[3:12]
df <- dplyr::mutate_at(df, protein_names, function(x) asinh(x/5))
glm_fit <- CytoGLMM::cytoglm(df,
                               protein_names = protein_names,
                               condition = "condition",
                               group = "donor",
                               num_boot = 10) # in practice >=1000
summary(glm_fit)
```

`summary.cytoglm` *Extract and calculate p-values of GLMM fit*

Description

Extract and calculate p-values of GLMM fit

Usage

```
## S3 method for class 'cytoglmm'  
summary(object, method = "BH", ...)
```

Arguments

object	A <code>cytobamm</code> class
method	Multiple comparison adjustment method
...	Other parameters

Value

tibble data frame

Examples

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