Package 'SureTypeSCR'

May 28, 2021

| Description |
|--|
| SureTypeSCR is R package for QC of (single cell) SNP arrays and single cell genotype scoring |
| Version 0.99.0 |
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| Suggests testthat,rmarkdown,markdown,knitr |
| Depends R (>= 3.5.0), reticulate, tidyverse, magrittr |
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| |

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Imports ggrepel, RColorBrewer, BiocStyle

 ${\bf bioc Views}\ \ Software, Genotyping Array, Single Cell$

VignetteBuilder knitr

SystemRequirements python (>= 3.6), sklearn, numpy, pandas,

Title Interface to python based package SureTypeSC via reticulate

SureTypeSC, IlluminaBeadArrayFiles

NeedsCompilation no

R topics documented:

| calculate_ma |
|------------------------------|
| callrate |
| configure_iaap |
| create_dataobject_from_frame |
| create_from_frame |
| getGEO_and_folder_in |
| get_multiind_df |
| get_simpleind_df |
| get_threshold |
| idat_to_gtc |
| plot_ma ? |
| plot_pca |
| predict_suretype |
| preprocess_pca |
| scbasic |
| scEls |
| scload |
| scpredict |

2 callrate

| calcı | ulate_ma | Ca | ılcu | lai | te i | M | (le | oge | arı | ith | ımı | ic i | dij | ffei | rei | ιсε | e) . | an | ıd | Α | (le | ge | ari | ith | ım | ic | av | er | ag | e) | of |
|-------|-------------------|----|------|-----|------|---|-----|-----|-----|-----|-----|------|-----|------|-----|-----|------|----|----|---|-----|----|-----|-----|----|----|----|----|----|----|------|
| Index | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | 17 |
| | write_samplesheet | | | | | • | | | | | | | | | • | | | | | | | | | | | | | | | • | . 15 |
| | suretype_model . | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | |
| | set_threshold | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | |
| | scTrain | | | | | | | | | | | | | | | | | | | | | | | | | | | | | | . 13 |

Description

Function applies Logartihmic transformation on signal intensities which is a preliminary step for classification using SureTypeSC.

Usage

```
calculate_ma(df)
```

Arguments

df

data frame from scbasic(.) or compatible

allelic intensities

Value

df with extra columns representing the results of MA transformation (m, a, m_raw and a_raw)

Examples

```
setwd(system.file(package='SureTypeSCR'))
samplesheet=system.file('files/GSE19247_example.csv',package='SureTypeSCR')
manifest=system.file('files/HumanCytoSNP-12v2_H.bpm',package='SureTypeSCR')
cluster=system.file('files/HumanCytoSNP-12v2_H.egt',package='SureTypeSCR')
#Load data
df=scbasic(manifest,cluster,samplesheet)
#calculate MA transform and store in the original dataframe
df
```

callrate

Calculates call rate as proportion of called SNPs in the input dataframe

Description

Calculate call rate as proportion of called SNPs in the input dataframe.

Usage

```
callrate(.data)
```

configure_iaap 3

Arguments

.data

data frame from scbasic or compatible

Value

Table with call rate(s) of the input data frame.

Examples

```
setwd(system.file(package='SureTypeSCR'))
samplesheet=system.file('files/GSE19247_example.csv',package='SureTypeSCR')
manifest=system.file('files/HumanCytoSNP-12v2_H.bpm',package='SureTypeSCR')
cluster=system.file('files/HumanCytoSNP-12v2_H.egt',package='SureTypeSCR')
#Load data
df=scbasic(manifest,cluster,samplesheet)
#Get overall callrate
df
#Get callrate per individual
df
#Get callrate as allelic fractions and pivot to columns
df
```

configure_iaap

Procedure for setting path to IAAP-cli executable

Description

This procedure first checks whether the package home folder contains IAAP-cli executable. If not, then it guides the user to download the IAAP-cli archive and configure path to the archive. The procedure then decompresses the archive and sets up the path.

Usage

```
configure_iaap()
```

Value

returns path to IAAP-cli executable

4 create_from_frame

```
create_dataobject_from_frame
```

Internal function for converting basic data frame from SureTypeSCR to SuretypeSC's python dataobject

Description

The function returns SureTypeSC's dataobject that contains multindexed pandas data frame and various metadata. This structre is used in some of the SureTypeSC's methods that are called from SureTypeSCR.

Usage

```
create_dataobject_from_frame(df_single)
```

Arguments

df_single

data frame used in SureTypeSCR

Value

SuretypeSC's data object (instance of class Data) containing multiindexed data frame and metadata.

create_from_frame

Internal function for converting between python structures

Description

This functions is used internally in scbasic and converts python data frame in to SureTypeSC's data object.

Usage

```
create_from_frame(df)
```

Arguments

df

genotyping pandas dataframe from scbasic function

Value

Instance of class Data from Python SureTypeSC library with multiindex.

getGEO_and_folder_in

Description

Downloads metadata (and data) for GEO records specified in x.

Usage

```
getGEO_and_folder_in(x,download=TRUE)
```

Arguments

x GEO record ID

download boolean flag indicating whether to download the data (TRUE) or just the metadata

(FALSE)

Value

Data frame with metadata

Examples

```
#library(GEOquery)

#gse <- getGEO("GSE19247",GSEMatrix = TRUE)

#samplelist_sperm=as.data.frame(gse$`GSE19247-GPL6985_series_matrix.txt.gz`) %>%

# filter((str_detect(cell.type.ch1,'sperm')) & str_detect(cell.amplication.ch1,'MDA')) %>%

# rownames()

#metadf_sperm=map_if(samplelist_sperm,function(x) !is.null(x),function(x) getGEO_and_folder_in(x,download=T #metadf_sperm_merged = Reduce(function(...) merge(..., all=T), metadf_sperm)
```

get_multiind_df

Internal function that creates data frame with multiindex

Description

Internal function for index conversion. Function converts R data frame to a Python pandas dataframe with multiindex, that is compatible with Python methods implemented in SureTypeSC.

Usage

```
get_multiind_df(df)
```

Arguments

df

R data frame as returned by function scbasic(.) or compatible

Value

returns data frame with multiindex

6 get_threshold

get_simpleind_df

Internal function that creates data frame with simple column index

Description

Internal function for index conversion. Function converts data frame's column multiindex into a simple form.

Usage

```
get_simpleind_df(df)
```

Arguments

df

data frame with column multiindex

Value

returns data frame with simple index

get_threshold

Determine threshold used on the data in data frame

Description

Determine score cutoff that was applied on the data - that is all genotypes below the returned value are no-calls.

Usage

```
get_threshold(.data,col='score')
```

Arguments

.data basic data frame compatible with output from scbasic(.)

col column name with score - effective columns GenCall's score (scode), Sure-

TypeSC's score (rfgda_score) or single layer SureTypeSC's score (rf_score)

Value

returns float

idat_to_gtc 7

idat_to_gtc

Convert raw itensity IDAT files to GTC using IAAP-cli

Description

The raw itensity data (red and green channel) are processed into GTC using Illumina's IAAP-cli. Function runs configure_iaap() before instantiating IAAP-cli to make sure IAAP-cli is properly configured.

Usage

```
idat_to_gtc(idat_inputfolder,gtc_output,manifest,cluster)
```

Arguments

```
idat_inputfolder
```

input folder with stored idat files

gtc_output output folder for the GTC files

manifest path to manifest file cluster path to cluster file

Examples

```
#library(GEOquery)
#gse <- getGEO("GSE19247",GSEMatrix = TRUE)

#samplelist_sperm=as.data.frame(gse$`GSE19247-GPL6985_series_matrix.txt.gz`) %>%
# filter((str_detect(cell.type.ch1,'sperm')) & str_detect(cell.amplication.ch1,'MDA')) %>%
# rownames()

#metadf_sperm=map_if(samplelist_sperm,function(x) !is.null(x),function(x) SureTypeSCR::getGEO_and_folder_in #metadf_sperm_merged = Reduce(function(...) merge(..., all=T), metadf_sperm)

#manifest=system.file("files/HumanCytoSNP-12v2_H.bpm",package="SureTypeSCR")
#cluster=system.file("files/HumanCytoSNP-12v2_H.egt",package="SureTypeSCR")

#for (ar in unique(metadf_sperm_merged$arrayid))
#{
# idat_to_gtc(ar,'GTC',manifest,cluster)
```

plot_ma

#}

Create MA plot per individual.

Description

Plot XY plot of logarithmic difference (M) and logarithmic average (A) of intensity signals-

8 plot_pca

Usage

```
plot_ma(.data,norm=TRUE,smooth=FALSE,nocalls=FALSE,n=1)
```

Arguments

| .data | data frame returned by scbasic or a compatible function |
|---------|--|
| norm | boolean flag indicating whether to use normalized (TRUE) or raw (FALSE) intensities |
| smooth | boolean flag indicating whether to apply smoothing spline per genotype (AA/BB/AB) cluster (TRUE or FALSE, set to FALSE by default. |
| nocalls | boolean flag indicating whether to plot SNPs that did not call (TRUE or FALSE, set to FALSE by default. |
| n | fraction of data to use from .data (real number in range 01), set to 1 by default meaning all points will by plotted. |

Value

ggplot object with MA plots facetted by individual

Examples

```
setwd(system.file(package='SureTypeSCR'))
samplesheet=system.file('files/GSE19247_example.csv',package='SureTypeSCR')
manifest=system.file('files/HumanCytoSNP-12v2_H.bpm',package='SureTypeSCR')
cluster=system.file('files/HumanCytoSNP-12v2_H.egt',package='SureTypeSCR')
#Load data
df=scbasic(manifest,cluster,samplesheet)
#Visualise MA plot
df
```

plot_pca

Run and visualise PCA

Description

Function creates feature matrix from non-zero SNPs and feature types indicated in features. The feature matrix used for calculating PCA has n rows and k columns, while n corresponds to number of individuals in the data frame and k is defined as length of features x number of non-zero SNPs across all samples.

Usage

```
\verb|plot_pca(.data,by\_chrom=TRUE,features=c('x','y'),importances=TRUE,metadata=NULL,labels=TRUE)||
```

predict_suretype 9

Arguments

| .data | dataframe from schasic or compatible |
|-------------|--|
| by_chrom | boolean flag indicating whether to run PCA by chromosome (TRUE or FALSE) |
| features | vector of feature types to use for PCA (columns from .data , i.e. $c('x', 'y')$) |
| importances | boolean flag indicating whether to render dimension importances (TRUE or $FALSE$) |
| metadata | data frame with metadata - currently grouping by family is supported, given that metadata contains familyid and individual columns and metadata will be merged to internal data frame by matching individual |
| labels | boolean flag indicating whether to render sample names using package ggrepel to prevent overplotting (TRUE or FALSE) |

Value

ggplot object with PCA plot

Examples

```
setwd(system.file(package='SureTypeSCR'))
samplesheet=system.file('files/GSE19247_example.csv',package='SureTypeSCR')
manifest=system.file('files/HumanCytoSNP-12v2_H.bpm',package='SureTypeSCR')
cluster=system.file('files/HumanCytoSNP-12v2_H.egt',package='SureTypeSCR')
#Load data
df=scbasic(manifest,cluster,samplesheet)
#create ggplot object with PCA
df
```

predict_suretype Run classification model using combination of Random Forest and Gaussian Discriminant Analysis

Description

Cascade classification model with Random Forest in the first layer and Gaussian Discriminant Analysis in the second layer. This function analyzes the whole dataset comprised in .data in one batch. To minimize bias in the model, it is recommended to run suretype_model(.) and build a model per sample basic instead of the whole dataset.

Usage

```
predict_suretype(.data, rf_clf)
```

Arguments

| .data | Input dataframe coming from scbasic() or compatible |
|--------|---|
| rf_clf | Instance of classifier loaded using scload currently embodied in RF |

10 preprocess_pca

Value

data frame decorated with SureTypeSC genotyping score

Examples

```
samplesheet=system.file('files/GSE19247_example.csv',package='SureTypeSCR')
manifest=system.file('files/HumanCytoSNP-12v2_H.bpm',package='SureTypeSCR')
cluster=system.file('files/HumanCytoSNP-12v2_H.egt',package='SureTypeSCR')
clf=system.file('files/rf.clf',package='SureTypeSCR')

#Load data
df=scbasic(manifest,cluster,samplesheet)

# The Random Forest classifier
clf_instance <- scload(clf)

#assign prediction results back to the original dataframe using margittr %<>% operator
df

preprocess_pca

Internal function that calculates matrix for principal component anal-
ysis
```

Description

Internal function that preprocesses feature matrix for PCA. Output of this function is used in plot_pca(.)

Usage

```
preprocess_pca(.data,.group,features=c('x','y'))
```

Arguments

.data output data frame from scbasic(.) or compatible

.group formal parameter effective when run under group_by(.) and nest(.) from

tidyverse

features list of columns from the .data data frame that will be used for creation of the

feature matrix

Value

```
matrix formatted for PCA analysis using stats::prcomp(.)
```

scbasic 11

scbasic

Load data in GTC format into data frame

Description

Function instantiates Illumina BeadArray library to load data from GTC files into data frame using information from manifest and cluster file.

Usage

```
scbasic(bpm,egt,samplesheet)
```

Arguments

bpm pathname to manifest file egt pathname to cluster file samplesheet pathname to samplesheet

Value

data frame stacked by individual

Examples

```
setwd(system.file(package='SureTypeSCR'))
samplesheet=system.file('files/GSE19247_example.csv',package='SureTypeSCR')
manifest=system.file('files/HumanCytoSNP-12v2_H.bpm',package='SureTypeSCR')
cluster=system.file('files/HumanCytoSNP-12v2_H.egt',package='SureTypeSCR')
#Load data
df=scbasic(manifest,cluster,samplesheet)
```

scEls

mediate access to python modules

Description

mediate access to python modules

Usage

```
scEls()
```

Value

list of (S3) "python.builtin.module"

Note

Returns a list containing objects sc and pd that refer to Python modules SureTypeSC and pandas, respectively.

12 scload

Examples

```
els = scEls()
els
##$sc
##Module(SureTypeSC)
##$pd
##Module(pandas)
```

scload

Load classifier from file

Description

Load pre-trained classifier from file.

Usage

```
scload(filename)
```

Arguments

filename

path to a file with classifier

Value

instance of a classifier

Examples

```
#gtc_path=system.file('files/GTCs',package='SureTypeSCR')
samplesheet=system.file('files/GSE19247_example.csv',package='SureTypeSCR')
manifest=system.file('files/HumanCytoSNP-12v2_H.bpm',package='SureTypeSCR')
cluster=system.file('files/HumanCytoSNP-12v2_H.egt',package='SureTypeSCR')
clf=system.file('files/rf.clf',package='SureTypeSCR')

#Load data
df=scbasic(manifest,cluster,samplesheet)

rf=scload(clf)
df
```

scpredict 13

| scpredict | <pre>Internal function used for prediction with suretype_model(.)</pre> | |
|-----------|---|--|
| | | |

Description

Internal function that evaluates score given an instance of classifier. The function is used in suretype_model(.)

Usage

```
scpredict(.data,clf,clftype='rf')
```

Arguments

.data data frame containing m and a features

clf instance of a classifier clftype The type of classifier

Value

data frame with predicted score using given classifier in clf

scTrain

Internal function that fits parameters of GDA to given data

Description

Internal function that creates GDA classifier based on the results of first layer (RF).

Usage

```
scTrain(trainingdata,clfname='gda')
```

Arguments

trainingdata intermediate results of RF classification

clfname name of the classifier to be fitted parameters for, currently 'Gaussian discrimi-

nant analysis' (GDA) is supported and change of this parameter will only have

effect on column name in the resulting data frame.

Value

instance of a GDA classifier

14 suretype_model

| set | thresho | ıα |
|-----|---------|----|

Set threshold on a classification method.

Description

Function truncates the genotypes (column gtype) given a threshold (cutoff) an classification method. This causes that all genotypes below threshold defined in threshol will be called as NC (no-call)

Usage

```
set_threshold(.data,clfcol,threshold)
```

Arguments

.data dataframe from scbasic or compatible

clfcol column with classification score

threshold threshold from 0 to 1

Value

data frame with gtype set to NC for classification score below chosen threshold

Examples

```
setwd(system.file(package='SureTypeSCR'))
samplesheet=system.file('files/GSE19247_example.csv',package='SureTypeSCR')
manifest=system.file('files/HumanCytoSNP-12v2_H.bpm',package='SureTypeSCR')
cluster=system.file('files/HumanCytoSNP-12v2_H.egt',package='SureTypeSCR')
#Load data
df=scbasic(manifest,cluster,samplesheet)
# changing Gencall score threshold
df
# with higher threshold we expect lower call rate
df
```

suretype_model

Create model and perform classification with RF-GDA in one individual

Description

Function is almost always used with in connection with group_by(invidiual) and nest() to create a classification model per group (typically one individual)

Usage

```
suretype_model(df,individual, clf,.sclist=NULL)))
```

write_samplesheet 15

Arguments

df Basic dataframe with precalculated M and A features

individual sample name to be classified

clf path to the classifier - currently support for Random Forest

.sclist samples to be processed stored in list(), default is NULLmeaning all samples in

df will be classified

Value

returns data frame with two columns corresponding to the classification results: rfgda_score and rf_score, while rfgda_score is score calculated by cascade RF-GDA algorithm and rf_score is score calculated by RF.

Examples

```
setwd(system.file(package='SureTypeSCR'))
samplesheet=system.file('files/GSE19247_example.csv',package='SureTypeSCR')
manifest=system.file('files/HumanCytoSNP-12v2_H.bpm',package='SureTypeSCR')
cluster=system.file('files/HumanCytoSNP-12v2_H.egt',package='SureTypeSCR')
clf=system.file("files/rf.clf",package="SureTypeSCR")

df_model = df
calculate_ma()
group_by(individual)
nest()
mutate(model=map(data , function(df) suretype_model(df,individual, clf,.sclist=list('gsm477563')))
```

write_samplesheet

Function creates samplesheet compatible with SureTypeSCR

Description

Function is used in case data was converted from idat and sample sheet was not available with the data.

Usage

write_samplesheet(filename,array_positions,gtc_output,manifest,experiment_name='Experiment1',pro

Arguments

filename name/path of the newly created sample sheet

array_positions

data frame storing metadata about sample id, array id and position, following

columns are required: sampleid, arrayid, position

gtc_output path where GTC files are stored

manifest name of the manifest file

experiment_name

experiment name

investigator_name

investigator name

16 write_samplesheet

Value

writes external file to filename

Examples

```
# library(GEOquery)

#gse <- getGEO("GSE19247",GSEMatrix = TRUE)
#manifest=system.file("files/HumanCytoSNP-12v2_H.bpm",package="SureTypeSCR")

#samplelist_sperm=as.data.frame(gse$`GSE19247-GPL6985_series_matrix.txt.gz`) %>%
# filter((str_detect(cell.type.ch1,'sperm')) & str_detect(cell.amplication.ch1,'MDA')) %>%
# rownames()

#metadf_sperm=map_if(samplelist_sperm,function(x) !is.null(x),function(x) getGEO_and_folder_in(x,download=T #metadf_sperm_merged = Reduce(function(...) merge(..., all=T), metadf_sperm)

#SureTypeSCR::write_samplesheet('samplesheet.csv',metadf_sperm_merged,'GTC',manifest)
```

Index

```
calculate_ma, 2
callrate, 2
configure_iaap, 3
create_dataobject_from_frame, 4
\verb|create_from_frame|, 4|
get_multiind_df, 5
\verb"get_simpleind_df", 6"
{\tt get\_threshold}, {\tt 6}
getGEO_and_folder_in, 5
idat_to_gtc, 7
plot_ma, 7
plot_pca, 8
predict_suretype, 9
preprocess_pca, 10
scbasic, 11
scEls, 11
scload, 12
scpredict, 13
scTrain, 13
set_threshold, 14
suretype\_model, 14
write\_samplesheet, 15
```