# Package 'OncoSubtype'

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Learning Method Version 1.0.0 Author Dadong Zhang <dadong.zhang.shared@gmail.com> Maintainer Dadong Zhang <dadong.zhang.shared@gmail.com> Description Provide functionality for cancer subtyping using nearest centroids or machine learning methods based on TCGA data. License GPL-3 **Encoding UTF-8** LazyData true RoxygenNote 7.3.1 URL https://github.com/DadongZ/OncoSubtype BugReports https://github.com/DadongZ/OncoSubtype/issues **Suggests** knitr, rmarkdown, testthat (>= 3.0.0) VignetteBuilder knitr LazyDataCompression xz **Imports** caret, randomForest, methods, e1071, pheatmap, tibble, dplyr, limma, rlang, Rdpack RdMacros Rdpack **Depends** SummarizedExperiment, R (>= 3.63), Config/testthat/edition 3

Title Predict Cancer Subtypes Based on TCGA Data using Machine

Type Package

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2 centroids\_subtype

# **R** topics documented:

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Index																						1
	SubtypeClass-class		• •			•		•		•	•		٠	•	 •	•	•	 •	•	 •	٠	
	PlotHeat																					
	ml_subtype																					
	lusc_centroids																					
	luad_centroids																					
	load_dataset_from_gith	ub																				
	hnsc_centroids																					
	get_rf_pred																					
	get_median_centered .																			 		
	get_hvg																					
	example_fpkm																			 		
	centroids_subtype																					

#### **Description**

Predict the subtypes of selected cancer type based published papers

#### Usage

```
centroids_subtype(data, disease = "LUSC")
```

#### **Arguments**

data set to predict the subtypes which is a numeric matrix with row names of

features and column names of samples

disease character string of the disease to predict subtypes, currently support 'LUSC',

'LUAD'

#### Value

an object of class "SubtypeClass" with four slots: genes used for predictiong, predicted subtypes of samples, a matrix of predicting scores, and the method.

## **Examples**

```
## Not run:
library(OncoSubtype)
data <- get_median_centered(example_fpkm)
data <- assays(data)$centered
rownames(data) <- rowData(example_fpkm)$external_gene_name
centroids_subtype(data, disease = 'HNSC')
## End(Not run)</pre>
```

example\_fpkm 3

example\_fpkm

example FPKM data

## Description

example FPKM data

## Usage

```
example_fpkm
```

#### **Format**

SummarizedExperiment object

get\_hvg

select highly variable genes from a expression matrix

## Description

select highly variable genes from a expression matrix

## Usage

```
get_hvg(data, top = 1000)
```

## **Arguments**

data a (normalized) matrix with rownames of features and colnames of samples

top number of top highly variable genes to output

#### Value

subset with top ranked genes by the variances

## **Examples**

```
## Not run:
library(OncoSubtype)
data <- get_median_centered(example_fpkm)
data <- assays(data)$centered
get_hvg(data)
## End(Not run)</pre>
```

get\_rf\_pred

get\_median\_centered

convert expression matrix to median-centered

#### **Description**

convert expression matrix to median-centered

## Usage

```
get_median_centered(data, log2 = TRUE)
```

#### **Arguments**

data a numeric matrix or 'S4' object

logical, if 'TRUE' log2(x+1) will be applied before median centering. Defaut

'TRUE'.

#### Value

median-centered express matrix or an object with new slot "centered"

## **Examples**

```
## Not run:
get_median_centered(example_fpkm)
## End(Not run)
```

get\_rf\_pred

Predict the subtypes of selected cancer type

## **Description**

Predict the subtypes of selected cancer type

#### Usage

```
get_rf_pred(train_set, test_set, method = "rf", seed = NULL)
```

## **Arguments**

train\_set training set with rownames of samples, first column named 'mRNA\_subtype'

and the rest of features and expression values.

test\_set test set with rownames of features and colnames of samples.

method character string of the method to use currently support 'rf'.

seed integer seed to use.

hnsc\_centroids 5

#### Value

a matrix with column names of subtypes and predicted probabilities.

hnsc\_centroids

HNSC predictor centroids

#### **Description**

HNSC predictor centroids from https://www.nature.com/articles/nature14129

#### Usage

hnsc\_centroids

#### **Format**

A tibble with 728 features and four subtypes.

load\_dataset\_from\_github

Load Dataset from GitHub Repository

## Description

Downloads a specified dataset from a GitHub repository if it is not already present in the specified local directory, then loads the dataset into the global environment. This function is designed to help manage package size by storing data externally and loading it on-demand.

## Usage

```
load_dataset_from_github(disease, local_dir = path.expand(getwd()))
```

#### **Arguments**

disease A character string specifying the disease, which corresponds to the name of the

dataset to be loaded (e.g., "LUSC"). The function constructs the filename as

tolower(disease)\_tcga.rda and attempts to load this dataset.

local\_dir An optional character string specifying the path to the directory where datasets

should be stored locally. If not provided, defaults to a subdirectory named your\_package\_name\_data within the user's home directory. Users can specify their own directory path if they prefer to store data in a different location.

#### Value

Invisible NULL. The function is primarily used for its side effect of loading a dataset into the global environment. However, the function itself does not return the dataset directly.

6 lusc\_centroids

## **Examples**

```
## Not run:
   load_dataset_from_github("LUSC")
## End(Not run)
```

luad\_centroids

LUAD predictor centroids

## Description

LUAD predictor centroids from Wilkerson (2012)

## Usage

luad\_centroids

#### **Format**

A tibble with 506 features and three subtypes bronchioid, magnoid, and squamoid.

 ${\tt lusc\_centroids}$ 

LUSC predictor centroids

## Description

LUSC predictor centroids from Wilkerson (2010)

## Usage

lusc\_centroids

## **Format**

A tibble with 208 features and four subtypes: primitive, classical, secretory, and basal.

ml\_subtype 7

ml_subtype Predict the subtypes of selected cancer type using machine learning	
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## **Description**

Predict the subtypes of selected cancer type using machine learning

## Usage

```
ml_subtype(
   data,
   disease = "LUSC",
   method = "rf",
   removeBatch = TRUE,
   seed = NULL
)
```

## Arguments

data	data set to predict the subtypes which is a numeric matrix with row names of features and column names of samples
disease	character string of the disease to predict subtypes, currently support 'LUSC', 'LUAD', and 'BLCA'.
method	character string of the method to use currently support 'rf'.
removeBatch	whether do batch effect correction using limma::removeBatchEffect, default TRUE.
seed	integer seed to use.

#### Value

An object of class "SubtypeClass" with four slots: genes used for predictiong, predicted subtypes of samples, a matrix of predicting scores, and the method.

#### References

- Wilkerson MJ, Yin X, Hayes D, et al. (2010). "Lung Squamous Cell Carcinoma mRNA Expression Subtypes Are Reproducible, Clinically Important, and Correspond to Normal Cell Types." Clin Cancer Res, 16(19), 4864-4875.
- 2. Wilkerson MJ, Yin X, Hayes D, et al. (2012). "Differential pathogenesis of lung adenocarcinoma subtypes involving sequence mutations, copy number, chromosomal instability, and methylation." *Plos One*, **7**(5), e36530.
- 3. Network TCGA (2015). "Comprehensive genomic characterization of head and neck squamous cell carcinomas." *Nature*, **517**, e36530.

8 PlotHeat

#### **Examples**

```
## Not run:
library(OncoSubtype)
data <- get_median_centered(example_fpkm)
data <- assays(data)$centered
rownames(data) <- rowData(example_fpkm)$external_gene_name
ml_subtype(data, disease = 'LUAD', method = 'rf', seed = 123)
## End(Not run)</pre>
```

PlotHeat

Plot heatmap of the train set or test set

## **Description**

Plot heatmap of the train set or test set

#### Usage

```
PlotHeat(object, set = "test", ...)
```

#### **Arguments**

```
object a SubtypeClass object
set options could be 'test', 'train' or 'both'. Default 'test'.
... Parameters passed to pheatmap.
```

#### Value

a pheatmap object

## **Examples**

```
## Not run:
library(OncoSubtype)
data <- get_median_centered(example_fpkm)
data <- assays(data)$centered
rownames(data) <- rowData(example_fpkm)$external_gene_name
object <- MLSubtype(data, disease = 'LUSC')
PlotHeat(object, set = 'both', fontsize = 10, show_rownames = FALSE, show_colnames = FALSE)
## End(Not run)</pre>
```

SubtypeClass-class 9

SubtypeClass-class Set the SubtypeClass
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## Description

Set the SubtypeClass

## Value

an object of SubtypeClass with three empty solts

# **Index**

```
* datasets
     example_fpkm, 3
     hnsc_centroids, 5
     luad_centroids, 6
     lusc_centroids, 6
centroids\_subtype, \textcolor{red}{2}
\verb|example_fpkm|, 3
get_hvg, 3
{\tt get\_median\_centered, 4}
get_rf_pred, 4
\verb|hnsc_centroids|, 5
load\_dataset\_from\_github, 5
luad_centroids, 6
\verb|lusc_centroids|, 6
ml_subtype, 7
{\tt PlotHeat}, {\color{red} 8}
SubtypeClass(SubtypeClass-class), 9
SubtypeClass-class, 9
```