Package ‘CytoMethIC’

June 6, 2024

Type Package

Title DNA methylation-based classification and regression

Description This package provides DNA methylation-based prediction of cancer type, molecular signature and clinical outcomes. It provides convenience functions for missing value imputation, probe ID conversion, model interpretation and visualization. The package links to our models on ExperimentHub. The package currently supports HM450, EPIC and EPICv2.

Version 1.0.0

License Artistic-2.0

Depends R (>= 4.4.0), ExperimentHub

Imports tibble, utils, stats, tools, sesame, sesameData, BiocParallel, BiocManager

VignetteBuilder knitr

Suggests BiocStyle, randomForest, testthat, knitr, rmarkdown, e1071, xgboost, keras, tensorflow

URL https://github.com/zhou-lab/CytoMethIC

BugReports https://github.com/zhou-lab/CytoMethIC/issues

biocViews ExperimentData, MicroarrayData, Genome, ExperimentHub, MethylationArrayData, CancerData, PackageTypeData

NeedsCompilation no

RoxygenNote 7.3.1

Encoding UTF-8

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<td>cmi_checkVersion</td>
<td>Check CytoMethIC versions</td>
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Description

print package version of cytometric and depended packages to help troubleshoot installation issues.

Usage

```r
cmi_checkVersion()
```

Value

print the versions of cytometric and dependencies

Examples

```r
cmi_checkVersion()
```

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<td>Master data frame for all model objects</td>
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Description

This is an internal object which will be updated on every new release

Value

master sheet of CytoMethIC model objects

Examples

```r
print(cmi_models[,c("EHID","Title")])
```
The `cmi_predict` function takes in a model and a sample, and uses the model to predict it. This function supports `randomForest`, `e1071::svm`, `xgboost`, and `keras/tensorflow` models. For `xgboost` and `keras` models, the features used in classification as well as a label mapping must be provided for output.

### Usage

```r
cmi_predict(
  betas,  # DNA methylation beta
  cmi_model,  # Cytomethic model downloaded from ExperimentHub
  source_platform = NULL,  # source platform If not given, will infer from probe ID.
  lift_over = FALSE,  # whether to allow mLiftOver to convert probe IDs
  verbose = FALSE,  # be verbose with warning
  BPPARAM = SerialParam()  # use MulticoreParam(n) for parallel processing
)
```

### Arguments

- **betas**: DNA methylation beta
- **cmi_model**: Cytomethic model downloaded from ExperimentHub
- **source_platform**: source platform If not given, will infer from probe ID.
- **lift_over**: whether to allow mLiftOver to convert probe IDs
- **verbose**: be verbose with warning
- **BPPARAM**: use MulticoreParam(n) for parallel processing

### Value

predicted cancer type label

### Examples

```r
library(sesame)
library(ExperimentHub)
library(CytomethIC)

## Cancer Type
```
model = ExperimentHub()['EH8395']

cmi_predict(openSesame(sesameDataGet("EPICv2.8.SigDF")[[1]]), model, lift_over=TRUE)
cmi_predict(openSesame(sesameDataGet("EPIC.1.SigDF"), model, lift_over=TRUE)
cmi_predict(sesameDataGet("HM450.1.TCGA.PAAD")$betas, model, lift_over=TRUE)
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