

Package ‘planet’

April 10, 2023

Title Placental DNA methylation analysis tools

Version 1.6.0

URL <https://victor.rbind.io/planet>, <http://github.com/wvictor14/planet>

BugReports <http://github.com/wvictor14/planet/issues>

Description This package contains R functions to infer additional biological variables to supplemental DNA methylation analysis of placental data. This includes inferring ethnicity/ancestry, gestational age, and cell composition from placental DNA methylation array (450k/850k) data. The package comes with an example processed placental dataset.

Depends R (>= 4.0)

Imports methods, tibble, magrittr, dplyr

Suggests ggplot2, testthat, tidyr, scales, minfi, EpiDISH, knitr, rmarkdown

License GPL-2

Encoding UTF-8

LazyData false

RoxygenNote 7.1.1

VignetteBuilder knitr

biocViews Software, DifferentialMethylation, Epigenetics, Microarray, MethylationArray, DNAMethylation, CpGIsland

Roxygen list(markdown = TRUE)

git_url <https://git.bioconductor.org/packages/planet>

git_branch RELEASE_3_16

git_last_commit 20b35d5

git_last_commit_date 2022-11-01

Date/Publication 2023-04-10

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ageCpGs	<i>Placental gestational age CpGs</i>
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Description

Coefficients from the three placental gestational age clocks from [Lee Y et al. 2019](#).

Reference: Lee Y, Choufani S, Weksberg R, et al. Placental epigenetic clocks: estimating gestational age using placental DNA methylation levels. *Aging (Albany NY)*. 2019;11(12):4238–4253. doi:10.18632/aging.102049. PMID: 31235674

Usage

```
data(ageCpGs)
```

Format

A [tibble](#) with coefficients for the RPC, CPC, and refined RPC.

ethnicityCpGs	<i>CpGs to predict ethnicity</i>
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Description

1860 CpGs used to predict ethnicity.

See [Yuan et al. 2019](#) for details.

Usage

```
data(ethnicityCpGs)
```

Format

A character vector of length 1860

Source

<https://pubmed.ncbi.nlm.nih.gov/31399127/>

planet-deprecated *Deprecated functions in **planet***

Description

These functions still work but will be removed (defunct) in the next version.

Details

- `pl_infer_ethnicity`: This function has been renamed `predictEthnicity`
- `pl_infer_age`: This function has been renamed `predictAge`

plBetas *Example placental DNA methylation data*

Description

6 DNA methylation profiles from preeclampsia and healthy control placentas. This data was downloaded from:

- [GSE75196](#)

"Genome wide DNA methylation profiling of normal and preeclampsia placental samples. Illumina Infinium HumanMethylation450 BeadChip (450K array) was used to obtain DNA methylation profiles in placental samples. Samples included 16 samples from healthy uncomplicated pregnancies and 8 samples from pregnancies affected by preeclampsia." - from [Yeung et al.](#)

The DNA methylation data for 24 placental samples were downloaded from [GSE75196](#). After normalizing using `minfi::preprocessNoob` and `watermelon::BMIQ`, the data were filtered to 6/24 samples and 10,000 random CpGs + those CpGs used in the gestational age clock and ethnicity classifier.

Reference: Yeung KR, Chiu CL, Pidsley R, Makris A et al. DNA methylation profiles in preeclampsia and healthy control placentas. *Am J Physiol Heart Circ Physiol* 2016 May 15;310(10):H1295-303. [PMID:26968548](#)

Usage

```
data(plBetas)
```

Format

A matrix

Source

<https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE75196>

plCellCpGsFirst *First trimester placental cell type coefficients*

Description

First trimester coefficients for placental cellular deconvolution from [YuanVetal.2020](#).

Reference: to be edited PMID: to be edited

Usage

```
data(plCellCpGsFirst)
```

Format

A [matrix](#) with coefficients for Trophoblasts, Stromal, Endothelial, Hofbauer cells, nRBCs, and Syncytiotrophoblasts.

plCellCpGsThird *Third trimester placental cell type coefficients*

Description

Third trimester coefficients for placental cellular deconvolution from [YuanVetal.2020](#).

Reference: to be edited PMID: to be edited

Usage

```
data(plCellCpGsThird)
```

Format

A [matrix](#) with coefficients for Trophoblasts, Stromal, Endothelial, Hofbauer cells, nRBCs, and Syncytiotrophoblasts.

plColors	<i>A color palette for placental cell types</i>
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Description

A nice color palette for placental cell types.

Used in [YuanVetal.2020](#).

Contains colors for:

- Syncytiotrophoblast
- Trophoblast
- Stromal
- Hofbauer
- Endothelial
- nRBCs

Usage

```
data(plColors)
```

Format

An object of class character of length 6.

plPhenoData	<i>Sample information accompanying pl_betas</i>
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Description

Sex, disease, and gestational age information associated with pl_betas.

Downloaded from the GEO accession:

- [GSE75196](#)

Reference: Yeung KR, Chiu CL, Pidsley R, Makris A et al. DNA methylation profiles in preeclampsia and healthy control placentas. Am J Physiol Heart Circ Physiol 2016 May 15;310(10):H1295-303. [PMID: 26968548](#)

Usage

```
data(plPhenoData)
```

Format

A [tibble](#)

Source

<https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE75196>

predictAge	<i>Predicts gestational age using placental DNA methylation microarray data</i>
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Description

predictAge Multiplies the coefficients from one of three epigenetic gestational age clocks, by the corresponding CpGs in a supplied betas data.frame.

Usage

```
predictAge(betas, type = "RPC")
```

Arguments

betas	An n by m dataframe of methylation values on the beta scale (0, 1), where the CpGs are arranged in rows, and samples in columns. Should contain all CpGs used in each clock
type	One of the following: "RPC" (Robust), "CPC", (Control) or "RRPC" (Refined Robust).

Details

Predicts gestational age using one of 3 placental gestational age clocks: RPC, CPC, or refined RPC. Requires placental DNA methylation measured on the Infinium 27K/450k/EPIC methylation array. Ensure as many predictive CpGs are present in your data, otherwise accuracy may be impacted.

It's recommended that you have all predictive CpGs, otherwise accuracy may vary.

Value

A vector of length m, containing inferred gestational age.

Examples

```
# Load placenta DNAm data
library(dplyr)
data(plBetas)
data(plPhenoData)

plPhenoData %>%
  mutate(inferred_ga = predictAge(plBetas, type = "RPC"))
```

predictEthnicity *Predicts ethnicity using placental DNA methylation microarray data*

Description

Uses 1860 CpGs to predict self-reported ethnicity on placental microarray data.

Usage

```
predictEthnicity(betas, threshold = 0.75)
```

Arguments

betas	n x m dataframe of methylation values on the beta scale (0, 1), where the variables are arranged in rows, and samples in columns. Should contain all 1860 predictors and be normalized with NOOB and BMIQ.
threshold	A probability threshold ranging from (0, 1) to call samples 'ambiguous'. Defaults to 0.75.

Details

Predicts self-reported ethnicity from 3 classes: Africans, Asians, and Caucasians, using placental DNA methylation data measured on the Infinium 450k/EPIC methylation array. Will return membership probabilities that often reflect genetic ancestry composition.

The input data should contain all 1860 predictors (cpgs) of the final GLMNET model.

It's recommended to use the same normalization methods used on the training data: NOOB and BMIQ.

Value

a [tibble](#)

Examples

```
## To predict ethnicity on 450k/850k samples  
  
# Load placenta DNAm data  
data(plBetas)  
predictEthnicity(plBetas)
```

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