# Package 'BioAge'

| June 13, 2020   |
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| Type Package  |
| Title Biological Age Calculations Using Several Biomarker Algorithms  |
| Version 0.1.0   |
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| <b>Description</b> This package measures biological aging using data from the National Health and Nutrition Examination Survey (NHANES). The package uses published biomarker algorithms to calculate three biological age measures: Klemera-Doubal Method biological age, phenotypic age, and homeostatic dysregulation. |
| License GPL-3   |
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| Imports dplyr, ggplot2  |
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| R topics documented:  |
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# Description

Calculate Klemera-Doubal Method (KDM) Biological Age

# Usage

```
bioage_calc(data, age, biomarkers, fit = NULL, s_ba2 = NULL)
```

# Arguments

| data       | The dataset for calculating KDM bioage  |
|------------|---|
| age        | A character vector (length=1) indicating the name of the variable for chronological age                             |
| biomarkers | A character vector indicating the names of the variables for the biomarkers to use in calculating KDM bioage        |
| fit        | An S3 object for model fit. If the value is NULL, then the parameters to use for training KDM bioage are calculated |
| s_ba2      | A particular fit parameter. Advanced users can modify this parameter to control the variance of bioage              |

## **Details**

Calculate KDM Biological Age

# Value

An object of class "bioage". This object is a list with two elements (data and fit)

# **Examples**

bioage\_nhanes 3

## **Description**

Calculate Klemera-Doubal Method (KDM) Biological Age using NHANES dataset

#### Usage

```
bioage_nhanes(biomarkers)
```

#### **Arguments**

biomarkers A character vector indicating the names of the variables for the biomarkers to

use in calculating bioage

#### **Details**

Calculate KDM Biological Age using NHANES dataset

#### Value

An object of class "bioage". This object is a list with two elements (data and fit)

# **Examples**

```
#Calculate KDM bioage
bioage = bioage_nhanes(biomarkers=c("fev","sbp","totchol","hba1c","albumin","creat","lncrp","alp","bun"))
#Extract bioage dataset
data = bioage$data
```

| hd_calc | hd_calc |  |  |
|---------|---------|--|--|
|---------|---------|--|--|

# Description

Calculate homeostatic dysregulation (HD)

# Usage

```
hd_calc(data, reference, biomarkers)
```

## **Arguments**

data The dataset for calculating HD

reference The reference dataset for calculating HD

biomarkers A character vector indicating the names of the variables for the biomarkers to

use in calculating HD

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#### **Details**

Calculate HD

#### Value

An object of class "hd". This object is a list with two elements (data and fit)

# **Examples**

hd\_nhanes

hd\_nhanes

## **Description**

Calculate homeostatic dysregulation (HD) using NHANES dataset

# Usage

```
hd_nhanes(biomarkers)
```

# **Arguments**

biomarkers

A character vector indicating the names of the variables for the biomarkers to use in calculating HD

## **Details**

Calculate HD using NHANES dataset

#### Value

An object of class "hd". This object is a list with two elements (data and fit)

# Examples

```
#Calculate HD
hd = hd_nhanes(biomarkers=c("albumin","lymph","mcv","glucose","rdw","creat","lncrp","alp","wbc"))
#Extract HD dataset
data = hd$data
```

phenoage\_calc 5

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## **Description**

Calculate Levine's Phenotypic Age

#### Usage

```
phenoage_calc(data, age, time, status, biomarkers, fit = NULL)
```

## **Arguments**

| data       | The dataset for calculating phenoage  |
|------------|---|
| age        | A character vector (length=1) indicating the name of the variable for chronological age                           |
| time       | A character vector (length=1) indicating the name of the variable for survival time                               |
| status     | A character vector (length=1) indicating the name of the variable for survival status                             |
| biomarkers | A character vector indicating the names of the variables for the biomarkers to use in calculating phenoage        |
| fit        | An S3 object for model fit. If the value is NULL, then the parameters to use for training phenoage are calculated |

# **Details**

Calculate Phenotypic Age

#### Value

An object of class "phenoage". This object is a list with two elements (data and fit)

# **Examples**

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phenoage\_nhanes

phenoage\_nhanes

#### **Description**

Calculate Levine's Phenotypic Age using NHANES dataset

#### Usage

```
phenoage_nhanes(biomarkers)
```

#### **Arguments**

biomarkers

A character vector indicating the names of the variables for the biomarkers to use in calculating phenoage

#### **Details**

Calculate Levine's Phenotypic Age using NHANES dataset

#### Value

An object of class "phenoage". This object is a list with two elements (data and fit)

# **Examples**

plot\_ba

plot\_ba

## **Description**

Plot correlations between biological aging measures and chronological age

## Usage

```
plot_ba(data, agevar, label)
```

# **Arguments**

data The dataset for plotting correlations

agevar A character vector indicating the names of the biological aging measures

label A character vector indicating the labels of the biological aging measures

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#### **Details**

Plot correlations between biological aging measures and chronological age

#### Note

Chronological age and gender variables need to be named "age" and "gender"

## **Examples**

plot\_baa plot\_baa

# **Description**

Plot correlations between biological age advancement (BAA) and chronological age

## Usage

```
plot_baa(data, agevar, label, axis_type)
```

#### **Arguments**

data
The dataset for plotting corplot

agevar
A character vector indicating the names of the interested biological aging measures

label
A character vector indicating the labels of the biological aging measures Values should be formatted for displaying along diagonal of the plot Names should be used to match variables and order is preserved

axis\_type
A character vector indicating the axis type (int or float) Use variable name to define the axis type

## **Details**

Plot correlations between BAA and chronological age

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#### **Examples**

```
#Create corplot of BAA with chronologicl age
agevar = c("bioage_advance0",
           "phenoage_advance0",
           "bioage_advance",
           "phenoage_advance",
           "hd",
           "hd_log")
label = c("bioage_advance0"="KDM\nBiological\nAge",
           "phenoage_advance0"="Levine\nPhenotypic\nAge",
           "bioage_advance"="Modified-KDM\nBiological\nAge",
           "phenoage_advance"="Modified-Levine\nPhenotypic\nAge",
           "hd" = "Mahalanobis\nDistance",
           "hd_log" = "Log\nMahalanobis\nDistance")
axis_type = c("bioage_advance0"="float",
              "phenoage_advance0"="float",
              "bioage_advance"="float",
              "phenoage_advance"="flot",
              "hd"="flot",
              "hd_log"="float")
f2 = plot_baa(data, agevar, labels, axis_type)
f2
```

table\_health

table\_health

## **Description**

Association with current health status outcomes

## Usage

```
table_health(data, agevar, outcome, label)
```

## **Arguments**

data The dataset for linear regression table

agevar A character vector indicating the names of the interested biological aging mea-

sures

outcome A character vector indicating the name of the interested current health status

outcomes

label A character vector indicating the labels of the biological aging measures

## **Details**

Association with current health status outcomes, adjusting for chronological age and gender and stratified by gender, race, and age

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#### Note

Chronological age and gender variables need to be named "age" and "gender"

## **Examples**

table\_ses

table ses

## **Description**

Association with socioeconomic variables

# Usage

```
table_ses(data, agevar, exposure, label)
```

## **Arguments**

data The dataset for linear regression table

agevar A character vector indicating the names of the interested biological aging mea-

sures

exposure A character vector indicating the name of the interested socioeconomic variables

#### **Details**

Association with socioeconomic variables, adjusting for chronological age and gender and stratified by gender, race, and age

## Note

Chronological age and gender variables need to be named "age" and "gender"

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## **Examples**

table\_surv

table\_surv

## **Description**

Mortality validation for biological aging measures

# Usage

```
table_surv(data, agevar, time, status, label)
```

## **Arguments**

| data   | The dataset for mortality table   |
|--------|---|
| agevar | A character vector indicating the names of the interested biological aging measures   |
| time   | A character vector (length=1) indicating the name of the variable for survival time   |
| status | A character vector (length=1) indicating the name of the variable for survival status |
| label  | A character vector indicating the labels of the biological aging measures             |

## **Details**

Mortality validation for biological age measures, adjusting for chronological age and gender and stratified by gender, race, and age

## Note

Chronological age, gender, and race/ethnicity variables need to be named "age", "gender", and "race"

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# **Examples**

table1

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