# Package 'exHExpPred'

# March 22, 2023

	ExpoHExpPredict module is designed to predict blood concentrations of chemicals and prioritize chemicals of health concern
Versi	concern.  on 1.0.0

**Description** Developing a prediction model on the annotation of chemicals in human blood can provide new insight into

the distribution and extent of exposures to a wide range of chemicals in humans. Our objective of this module is

to develop a machine learning model (eg., random forest) to predict blood concentrations of chemicals and prioritize chemicals of health concern.

**License** GPL (>= 3)

**Encoding** UTF-8

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Imports httr,writex1,ggplot2,readx1,vroom

NeedsCompilation no

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# **R** topics documented:

	ConvToExpoI	D.												 				2
	FuncExit																	2
	InitHEP																	3
	LoadHEP																	4
	PredBlood													 				4
	VizPredBlood													 				5
Index																		7

2 FuncExit

ConvToExpoID

Convert different IDs to the unified ExposomeX IDs

#### **Description**

Convert the IDs of exposure, chemicals, metabolites, or proteins to the unified ExposomeX ID, i.e., unified identifier in ExposomeX platform

# Usage

```
ConvToExpoID(PID,OutPath="default")
```

# **Arguments**

PID chr. Program ID. It must be the same with the PID generated by InitHEP.

OutPath chr. Output file directory, e.g. "D:/test". It should be noted that the slash symbol

is "/", not "\". If "default", the current working directory will be set.

#### Value

A data frame containing the converted ID information.

#### Author(s)

Mingliang Fang, Weinan Lin, Bin Wang (corresponding author)

# **Examples**

```
res = InitHEP()
  res1 = LoadHEP(PID = res$PID,UseExample = "example#1")
  res2 = ConvToExpoID(PID = res$PID)
```

FuncExit

End the module analysis

# **Description**

End the module analysis

#### Usage

FuncExit(PID)

# **Arguments**

PID chr. Program ID. It must be the same with the PID generated by any initial

functions.

# Value

Exit status

InitHEP 3

### Author(s)

Bin Wang (corresponding author)

# **Examples**

```
res = InitHEP()
  res = LoadHEP(PID = res$PID,UseExample = "example#1")
  FuncExit(PID = res$PID)
```

InitHEP

Initialize HExpPridict module

# Description

Initialize HExpPridict module analysis. It can generate an R6 class object integrating all the analysis information.

# Usage

InitHEP()

#### **Details**

ExpoHExpPredict module is designed to predict blood concentrations of chemicals and prioritize chemicals of health concern.

# Value

An R6 class object.

# Author(s)

Mingliang Fang, Bin Wang,(corresponding author)

# **Examples**

```
res <- InitHEP()</pre>
```

4 PredBlood

LoadHEP	Load data file for HExpPridict module
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# **Description**

Load data file for HExpPridict module.

# Usage

```
LoadHEP(PID, UseExample = "default", DataPath=NULL)
```

## **Arguments**

PID chr. Program ID. It must be the same with the PID generated by InitHEP.

UseExample chr. Method of uploading data. If "default", user should upload their own data

files, or use "example#1" provided by this module.

DataPath chr. Input file directory, e.g. "D:/test/eg\_hep.xlsx". It should be noted that the

slash symbol is "/", not "\".

#### Value

An R6 class object containing the input data.

#### Author(s)

Mingliang Fang, Bin Wang (corresponding author)

# **Examples**

```
res <- InitHEP()
  res = LoadHEP(PID = res$PID, UseExample = "example#1")</pre>
```

PredBlood

Predict blood concentrations of chemicals

#### **Description**

Predict blood concentrations of chemicals

#### Usage

```
PredBlood(PID, OutPath="default",MC = "F",N=100)
```

# Arguments

PID chr. Program ID. It must be the same with the PID generated by InitHEP.

OutPath chr. Output file directory. e.g. "D:/test". It should be noted that the slash symbol

is "/", not "\". If "default", the current working directory will be set.

MC lgl. T (or TRUE) and F (or FALSE). Whether to perform Mentocaro simulation.

N num. Times of Mentocaro simulation.

VizPredBlood 5

#### Value

A list object containing the prediction results.

#### Author(s)

Mingliang Fang, Bin Wang (corresponding author)

# **Examples**

```
res = InitHEP()
  res1 = LoadHEP(PID = res$PID,UseExample = "example#1")
  res2 = ConvToExpoID(PID = res$PID)
  res3 = PredBlood(PID = res$PID, OutPath="default", MC ='F',N=1000)
```

VizPredBlood

Visualize the PredBlood

## **Description**

Visualize the PredBlood.

# Usage

# **Arguments**

PID chr. Program ID. It must be the same with the PID generated by InitBioLink.

OutPath chr. Output file directory. e.g. "D:/test". It should be noted that the slash symbol

is "/", not "\". If "default", the current working directory will be set.

Layout chr. Visualization layout. Available options include "forest" and "boxplot".

Brightness chr. Visualization brightness. Available options include "light" and "dark".

Palette chr. Visualization palette. Available options include "default1", "default2" and

several journal preference styles (i.e., cell, nature, science, lancet, nejm, and

jama).

#### Value

A list object containing the plot of the biological link. This plot can be further processed using ggplot2 package.

# Author(s)

Mingliang Fang, Ning Gao, Bin Wang (corresponding author)

6 VizPredBlood

# Examples

```
res = InitHEP()
    res1 = LoadHEP(PID = res$PID,UseExample = "example#1")
    res2 = ConvToExpoID(PID = res$PID)
    res3 = PredBlood(PID = res$PID, OutPath="default", MC ='F', N=1000)
    res4 = VizPredBlood(PID = res$PID, OutPath ="default",Layout="forest",
    Brightness="light",Palette= "science" )
```

# Index

```
ConvToExpoID, 2
FuncExit, 2
InitHEP, 3
LoadHEP, 4
PredBlood, 4
VizPredBlood, 5
```