## Package 'AEenrich'

November 1, 2021

Version 1.1.0

**Title** Adverse Event Enrichment Tests

Type Package

Description We extend existing gene enrichment tests to perform adverse event enrichment analysis. Unlike the continuous gene expression data, adverse event data are counts. Therefore, adverse event data has many zeros and ties. We propose two enrichment tests. One is a modified Fisher's exact test based on pre-selected significant adverse events, while the other is based on a modified Kolmogorov-Smirnov statistic. We add Covariate adjustment to improve the analysis. "Adverse event enrichment tests using VAERS" Shuoran Li, Lili Zhao (2020) <arXiv:2007.02266>.

License GPL-2
Encoding UTF-8
LazyData true
Biarch true
Depends R (>= 3.5.0)
Imports dplyr, magrittr, qvalue, doParallel, tidyr, modelr, foreach, rlang, utils

biocViews

URL https://github.com/umich-biostatistics/AEenrich

BugReports https://github.com/umich-biostatistics/AEenrich/issues

RoxygenNote 7.1.1 Suggests testthat NeedsCompilation no

**F** 

Author Shuoran Li [aut], Hongfan Chen [aut], Lili Zhao [aut], Michael Kleinsasser [aut, cre]

Maintainer Michael Kleinsasser <mkleinsa@umich.edu>

**Repository** CRAN

**Date/Publication** 2021-11-01 15:20:07 UTC

2 AEenrich-package

## **R** topics documented:

AFen	rich-package	A	Een	ric	h:	Ad	lve	rse	• E	ve	nt	E	nri	ich	ım	en	t T	Te.	s <i>t</i> .s	,								
Index																												9
	group				•		•	•		•	•	•		•	•	•	•	•	•		•	•			•	•	•	8
	enrich																											
	covid2																											
	covid1																											4
	count_cases																											
	AEenrich-packag																											

## Description

The count\_cases function is used to convert data on the report level to aggregated data, grouping by specified covariates.

Use the function count\_cases to convert report level data into aggregated data.

See our Github home page or run ?count\_cases for examples.

Perform Adverse Event Enrichment Tests The enrich function is used to perform Adverse event (AE) enrichment analysis. Unlike the continuous gene expression data, AE data are counts. Therefore, AE data has many zeros and ties. We propose two enrichment tests. AEFisher is a modified Fisher's exact test based on pre-selected significant AEs, while AEKS is based on a modified Kolmogorov-Smirnov statistic.

Use the function enrich to fit models and inspect results.

See our Github home page or run ?enrich for examples.

#### Author(s)

Maintainer: Michael Kleinsasser <mkleinsa@umich.edu>

Authors:

- Shuoran Li <shuoranl@umich.edu>
- Hongfan Chen <chenhf@umich.edu>
- Lili Zhao <zhaolili@med.umich.edu>

Maintainer: Michael Kleinsasser <mkleinsa@umich.edu>

Authors:

- Shuoran Li <shuoranl@umich.edu>
- Hongfan Chen <chenhf@umich.edu>
- Lili Zhao <zhaolili@med.umich.edu>

count\_cases 3

## See Also

Useful links:

- https://github.com/umich-biostatistics/AEenrich
- Report bugs at https://github.com/umich-biostatistics/AEenrich/issues

#### Useful links:

- https://github.com/umich-biostatistics/AEenrich
- Report bugs at https://github.com/umich-biostatistics/AEenrich/issues

count\_cases

Convert data on the report level to aggregated data.

## **Description**

The count\_cases function is used to convert data on the report level to aggregated data, grouping by specified covariates.

## Usage

```
count_cases(
  data,
  drug.case = drug.case,
  drug.control = NULL,
  covar_disc = NULL,
  covar_cont = NULL,
  breaks = NULL,
  cores = detectCores(),
  min_AE = 10
```

## Arguments

data	a data.frame with at least 3 columns, consisting data on the report level, having ID, Drug type and AE name as the first 3 columns with covariates(optional) followed. The order of columns is not interchangeable.
drug.case	a character string for the target drug of interest.
drug.control	a character string for the reference drug. If $NULL(default)$ , all other drugs combined are the reference.
covar_disc	a character vector of categorical covariates.
covar_cont	a character vector of continuous covariates.

4 covid1

breaks	a list consists of vectors used for creating specific bins to transform continuous covariates into categorical. Breaks Should have the same length as covar_cont. Given a vector of non-decreasing breakpoints in breaks[i], find the interval containing each element of covar_cont[i]; i.e., for each index j in breaks[i], value j is assigned to covar_cont[i] if and only if breaks[i][j] <= covar_cont[i] < breaks[i][j+1].
cores	the number of cores to use for parallel execution.
min_AE	the minimum number of cases required to start counting for a specific AE. De-

#### Value

A data.frame consists of aggregated data.

fault 10.

The returned data.frame contains the following columns:

- DRUG\_TYPE: type of the drug, DrugYes for target drug and DrugNo for referenced drug
- AE\_NAME: the name of the adverse event
- AEyes: number of observations that have this AE
- AEno: number of observations that do not have this AE
- covariates: covariates specifed by user

## **Examples**

```
# count_cases(data = covid1, drug.case = "COVID19", drug.control = "OTHER",
# covar_cont = c("AGE"), covar_disc = c("SEX"),
# breaks = list(c(16,30,50,65,120)))
```

covid1

Covid Vaccine Adverse Event Data

#### **Description**

Adverse event data in the long format. Each row is a single adverse event, along with covariates.

## Usage

covid1

#### **Format**

An object of class tbl\_df (inherits from tbl, data.frame) with 12500 rows and 5 columns.

covid2 5

## **Details**

- VAERS\_ID Event ID
- VAX\_LABEL Vaccine type
- AE\_NAME Adverse event name
- AGE covariate
- · SEX covariate

covid2

Covid Vaccine Adverse Event Data

## Description

Adverse event data in the short format. Each row is a count of adverse events with the given name.

## Usage

covid2

#### **Format**

An object of class tbl\_df (inherits from tbl, data.frame) with 2656 rows and 6 columns.

## **Details**

- DRUG\_TYPE Vaccine type
- AE\_NAME Adverse event name
- AEYes Number of observations that have this AE
- AENo Number of observations that do not have this AE
- AGE covariate
- SEX covariate

6 enrich

enrich

Perform Adverse Event Enrichment Tests

#### **Description**

The enrich function is used to perform Adverse event (AE) enrichment analysis. Unlike the continuous gene expression data, AE data are counts. Therefore, AE data has many zeros and ties. We propose two enrichment tests. AEFisher is a modified Fisher's exact test based on pre-selected significant AEs, while AEKS is based on a modified Kolmogorov-Smirnov statistic.

## Usage

```
enrich(
  data.
  dd.group,
  drug.case,
  drug.control = NULL,
  method = "aeks",
  n_perms = 1000,
  covar = NULL,
  p = 0,
  q.cut = 0.1,
  or.cut = 1.5,
  zero = FALSE,
 min_size = 5,
 min_AE = 10,
  cores = detectCores()
)
```

#### **Arguments**

data

a data.frame. Two data types are allowed. Type I data consisting data on the report level, having ID, Drug type and AE name as the first 3 columns with covariates(optional) followed. Type II data have drug type and AE name as the first two columns, with the 3rd and 4th Columns giving the numbers of successes(have AE) and failures(Do not have AE) respectively, then followed by covariates. See example data for details.

dd.group

a data.frame with AE name and Group name. This data.frame have the group information for each individual AE.

drug.case

a character string for the target drug of interest.

drug.control

a character string for the reference drug. If NULL(default), all other drugs com-

bined are the reference.

method

a character string specifying the method for the enrichment test. It must take "aeks" (default) or "aefisher"; "aeks" is the rank-based enrichment test, and "aefisher" is the Fisher enrichment test. See details described in the paper (see

reference section of this document).

enrich 7

n_perms	an integer value specifying the number of permutations in permutation test.
covar	a character vector specifying the columns of covariates, default NULL.
p	a numerical value to control the weight of the step, can take any value between 0 and 1. If $0$ (default), reduces to the standard Kolmogorov-Smirnov statistics.
q.cut	a numerical value specifying the significance cut for q value of AEs in aefisher.
or.cut	a numerical value specifying the significance cut for odds ratio of AEs in ae-fisher.
zero	logical, default FALSE.If TRUE, add zero indicator to enrichment score.
min_size	the minimum size of group required for enrichment analysis.
min_AE	the minimum number of cases required to start counting for a specific AE.
cores	the number of cores to use for parallel execution.

#### Value

A list containing 2 data.frames named **Final\_result** and **AE\_info**.

The **Final\_result** data.frame contains the following columns:

• GROUP\_NAME: AE group names

• ES: enrichment score

• p\_value: p value of the enrichment test

• GROUP\_SIZE: number of AEs per group

The **AE\_info** contains the following columns:

• AE NAME: AE names

• OR: odds ratio for each individual AE

• p\_value: p value for AE-drug association

• 95Lower: lower bound of 95 percent confidence interval of odds ratio

• 95Lower: upper bound of 95 percent confidence interval of odds ratio

• se(logOR): standard error of log odds ratio

#### References

Li, S. and Zhao, L. (2020). Adverse event enrichment tests using VAERS. arXiv:2007.02266.

Subramanian, A.e.a. (2005). Gene set enrichment analysis: a knowledge-based approach for interpreting genome-wide expression profiles. Proc Natl Acad Sci U S A. Proceedings of the National Academy of Sciences. 102. 15545-15550.

Tian, Lu & Greenberg, Steven & Kong, Sek Won & Altschuler, Josiah & Kohane, Isaac & Park, Peter. (2005). Discovering statistically significant pathways in expression profiling studies. Proceedings of the National Academy of Sciences of the United States of America. 102. 13544-9. 10.1073/pnas.0506577102.

8 group

#### **Examples**

```
# AEKS
### Type I data: data on report level
# enrich(data = covid1, covar = c("SEX", "AGE"), p = 0, method = "aeks",
        n_perms = 1000, drug.case = "COVID19", dd.group = group, cores = 2,
        drug.control = "OTHER", min_size = 5, min_AE = 10, zero = FALSE)
#
## Type II data: aggregated data
# enrich(data = covid2, covar = c("SEX", "AGE"), p = 0, method = "aeks",
        n_perms = 1000, drug.case = "DrugYes", dd.group = group, cores = 2,
        drug.control = "DrugNo", min_size = 5, min_AE = 10)
# AEFISHER
## Type I data: data on report level
# enrich(data = covid1, covar = c("SEX", "AGE"), p = 0, method = "aefisher",
        n_perms = 1000, drug.case = "COVID19", dd.group = group,
        drug.control = "OTHER", min_size = 5, min_AE = 10, q.cut = 0.05,
#
        or.cut = 1.5, cores = 2)
#
## Type II data: aggregated data
# enrich(data = covid2, covar = c("SEX", "AGE"), p = 0, method = "aefisher",
        n_perms = 1000, drug.case = "DrugYes", dd.group = group,
#
        drug.control = "DrugNo", min_size = 5, min_AE = 10, cores = 2)
```

group

Group Structure Data

#### **Description**

Identifies which group each set of adverse events belongs.

## Usage

group

#### **Format**

An object of class NULL of length 0.

#### **Details**

- AE\_NAME Adverse event name
- GROUP\_NAME Group name

# **Index**

```
* datasets
      covid1, 4
      covid2, 5
      group, 8

AEenrich (AEenrich-package), 2
AEenrich-package, 2

count_cases, 3
  covid1, 4
  covid2, 5

enrich, 6

group, 8
```