Package: vaxpmx (via r-universe)

August 27, 2024

Type Package

Title Vaccines Pharmacometrics

Version 0.0.3

Depends R (>= 4.0)

Date 2024-02-20

Maintainer Julie Dudasova (MSD) < julie.dudasova@merck.com>

Description Estimate vaccine efficacy (VE) using immunogenicity data. The inclusion of immunogenicity data in regression models can increase precision in VE. The methods are described in the publication ``Elucidating vaccine efficacy using a correlate of protection, demographics, and logistic regression" by Julie Dudasova, Zdenek Valenta, and Jeffrey R. Sachs (2024).

License GPL-3

Copyright Copyright © 2024 Merck & Co., Inc., Rahway, NJ, USA and its affiliates. All rights reserved.

Encoding UTF-8

LazyData true

RoxygenNote 7.1.2

Imports methods (>= 3.5.2), stats, MASS (>= 7.3-51.6), dplyr (>= 1.0.0),

Suggests knitr, rmarkdown, testthat

NeedsCompilation no

Author Julie Dudasova (MSD) [aut, cre]

Date/Publication 2024-02-28 11:50:02 UTC

Repository https://vavrycuk-zz.r-universe.dev

RemoteUrl https://github.com/cran/vaxpmx

RemoteRef HEAD

RemoteSha e3d06b2828e9b97e68081df9be32504ef917a954

2 EfficacyCI

Contents

data_	_temp	Exa	атр	ole	of	a i	hy	poi	the	etio	cai	l v	ac	cir	ıe	cli	ini	ca	l t	ria	ıl a	lat	ʻa	sei	t				
Index																													
	ve			•		٠	•			٠				•	•	•			•	•		•	•		•		•	٠	-
	vaxpmx																												
	glmParametricSamp	oling																											3
	data_temp EfficacyCI																												

Description

A dataset containing immunogenicity data, and clinical outcome data in the vaccinated and control groups. The dataset is provided in the form of a data frame.

Usage

data_temp

Format

Data frame:

ID identification of subjects

nAb1 value of neutralizing titer for serotype 1

nAb2 value of neutralizing titer for serotype 2

group binary indicator of a baseline demographic characteristics of interest

vaccine binary indicator of treatment arm, with value 1 in vaccinated and 0 in control subjects

type_disease serotype of disease

disease_any binary indicator of disease caused by any serotype

EfficacyCI

Efficacy summary (mean, median, confidence intervals)

Description

Function summarizes efficacy statistics (mean, median, confidence intervals) based on the set of estimated efficacy values and chosen condfidence interval.

Usage

```
EfficacyCI(efficacySet, ci = 0.95)
```

glmParametricSampling

3

Arguments

```
efficacySet numeric vector - vector of estimated efficacy values

ci numeric - required confidence level
```

Details

Confidence intervals are calculated using quantiles of estimated efficacy values.

Value

```
named list - mean, median, CILow, CIHigh
```

Examples

```
# Load required packages
library(dplyr)

# Load an example dataset
data(data_temp)
Data.vaccinated <- filter(data_temp, vaccine == 1)
Data.control <- filter(data_temp, vaccine == 0)

# Fit logistic model relating neutralizing titer to disease status, specific to serotype 2
logisticFit <- glm(disease_any ~ nAb1, data = data_temp, family = binomial())

# Estimate 95\% confidence interval of vaccine efficacy based on the fitted model
efficacySet <- glmParametricSampling(logisticFit, nboot = 500, Data.vaccinated, Data.control)
EfficacyCI(efficacySet)</pre>
```

 ${\tt glmParametricSampling} \begin{tabular}{ll} Accounting for the uncertainty on the fitted "glm" model and observed \\ data \\ \end{tabular}$

Description

glmParametricSampling is used for vaccine efficacy confidence interval construction. It provides a vector of vaccine efficacy values, with length of nboot. 95% confidence interval, defined by 2.5th and 97.5th percentile of this vector, accounts for the uncertainty on the model fit (via parametric resampling of the posterior distribution of the model parameters) and observed data (via bootstrapping).

Usage

```
glmParametricSampling(Fit, nboot = 2000, Data.vaccinated, Data.control)
```

4 vaxpmx

Arguments

Fit an object of class inheriting from "glm" representing the fitted model

nboot a numeric value for number of bootstrap samples for confidence interval con-

struction

Data.vaccinated

a data frame for the vaccinated group, containing the variables in the fitted model; data must include a column called "vaccine" with binary indicator of

vaccination status

Data.control a data frame for the control group, containing the variables in the fitted model;

data must include a column called "vaccine" with binary indicator of vaccination

status

Value

a vector of vaccine efficacy values VE_set, with length of nboot

Examples

```
# Load required packages
library(dplyr)

# Load an example dataset
data(data_temp)
Data.vaccinated <- filter(data_temp, vaccine == 1)
Data.control <- filter(data_temp, vaccine == 0)

# Fit logistic model relating neutralizing titer to disease status, specific to serotype 2
logisticFit <- glm(disease_any ~ nAb1, data = data_temp, family = binomial())

# Estimate 95\% confidence interval of vaccine efficacy based on the fitted model
efficacySet <- glmParametricSampling(logisticFit, nboot = 500, Data.vaccinated, Data.control)
CI <- lapply(EfficacyCI(efficacySet),"*", 100)</pre>
```

vaxpmx

vaxpmx

Description

pharmacometric modeling in vaccines

Author(s)

Julie Dudasova

ve 5

Description

Calculates vaccine efficacy and confidence interval as described in Dudasova et al., 2024, BMC Med Res Methodol

Usage

```
ve(Fit, Data, nboot = 2000)
```

Arguments

Fit	an object of class inheriting from "glm" representing the fitted model
Data	a data frame containing the variables in the fitted model; data must include a column called "vaccine" with binary indicator of vaccination status
nboot	a numeric value for number of bootstrap samples for confidence interval construction

a value of vaccine efficacy VE and lower and upper bound of confidence interval CI

Examples

Value

```
# Load an example dataset
data(data_temp)

# Fit logistic model relating neutralizing titer to disease status
logisticFit <- glm(disease_any ~ nAb1, data = data_temp, family = binomial())

# Estimate vaccine efficacy and 95\% confidence interval based on the fitted model
ve(logisticFit, data_temp, nboot = 500)</pre>
```

Index