

Package: TestingSimilarity (via r-universe)

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Type Package

Title Bootstrap Test for the Similarity of Dose Response Curves
Concerning the Maximum Absolute Deviation

Version 1.1

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Description Provides a bootstrap test which decides whether two dose response curves can be assumed as equal concerning their maximum absolute deviation. A plenty of choices for the model types are available, which can be found in the 'DoseFinding' package, which is used for the fitting of the models. See [<doi:10.1080/01621459.2017.1281813>](https://doi.org/10.1080/01621459.2017.1281813) for details.

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Beta model:

$$m(d, \beta) = E_0 + E_{max} B(\delta_1, \delta_2) (d/scal)^{\delta_1} (1 - d/scal)^{\delta_2}$$

with

$$B(\delta_1, \delta_2) = (\delta_1 + \delta_2)^{\delta_1 + \delta_2} / (\delta_1^{\delta_1} \delta_2^{\delta_2})$$

and *scal* is a fixed dose scaling parameter.**Usage**

```
betaMod(d, e, scal)
```

Arguments

- | | |
|------|--|
| d | real-valued argument to the function (dose variable) |
| e | model parameter |
| scal | fixed dose scaling parameter |

Value

Response value.

bootstrap_test	<i>Bootstrap test for the equivalence of dose response curves via the maximum absolute deviation</i>
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Description

Function for testing whether two dose response curves can be assumed as equal concerning the hypotheses

$$H_0 : \max_{d \in \mathcal{D}} |m_1(d, \beta_1) - m_2(d, \beta_2)| \geq \epsilon \text{ vs. } H_1 : \max_{d \in \mathcal{D}} |m_1(d, \beta_1) - m_2(d, \beta_2)| < \epsilon,$$

where

$$\mathcal{D}$$

denotes the dose range. See <https://doi.org/10.1080/01621459.2017.1281813> for details.

Usage

```
bootstrap_test(data1, data2, m1, m2, epsilon, B = 2000, bnds1 = NULL,
               bnds2 = NULL, plot = FALSE, scal = NULL, off = NULL)
```

Arguments

data1, data2	data frame for each of the two groups containing the variables referenced in dose and resp
m1, m2	model types. Built-in models are "linlog", "linear", "quadratic", "emax", "exponential", "sigEmax", "betaMod" and "logistic"
epsilon	positive argument specifying the hypotheses of the test
B	number of bootstrap replications. If missing, default value of B is 5000
bnds1, bnds2	bounds for the non-linear model parameters. If not specified, they will be generated automatically
plot	if TRUE, a plot of the absolute difference curve of the two estimated models will be given
scal, off	fixed dose scaling/offset parameter for the Beta/ Linear in log model. If not specified, they are 1.2*max(dose) and 1 respectively

Value

A list containing the p.value, the maximum absolute difference of the models, the estimated model parameters and the number of bootstrap replications. Furthermore plots of the two models are given.

References

<https://doi.org/10.1080/01621459.2017.1281813>

Examples

```
data(IBScovars)
male<-IBScovars[1:118,]
female<-IBScovars[119:369,]
bootstrap_test(male,female,"linear","emax",epsilon=0.35,B=300)
```

dff

Implementation of absolute difference function

Description

Function calculating the absolute difference of two dose response models:

$$dff(d, \beta_1, \beta_2) = |m_1(d, \beta_1) - m_2(d, \beta_2)|$$

Usage

```
dff(d, beta1, beta2, m1, m2)
```

Arguments

- | | |
|--------------|--|
| d | real-valued argument to the function (dose variable) |
| beta1, beta2 | model parameters (real vectors) |
| m1, m2 | model types. Built-in models are "linlog", "linear", "quadratic", "emax", "exponential", "sigEmax", "betaMod" and "logistic" |

Value

Response value for the absolute difference of two models.

emax

Implementation of EMAX models

Description

Emax model:

$$m(d, \beta) = E_0 + E_{max} \frac{d}{ED_{50} + d}$$

Usage

```
emax(d, e)
```

Arguments

- | | |
|---|--|
| d | real-valued argument to the function (dose variable) |
| e | model parameter |

Value

Response value.

exponential*Implementation of exponential models*

Description

Exponential model:

$$m(d, \beta) = E_0 + E_1(\exp(d/\delta) - 1)$$

Usage

`exponential(d, e)`

Arguments

- | | |
|---|--|
| d | real-valued argument to the function (dose variable) |
| e | model parameter |

Value

Response value.

linear*Implementation of linear models*

Description

Linear model:

$$m(d, \beta) = E_0 + \delta d$$

Usage

`linear(d, e)`

Arguments

- | | |
|---|--|
| d | real-valued argument to the function (dose variable) |
| e | model parameter |

Value

Response value.

linlog*Implementation of linear in log models***Description**

Linear in log Model model:

$$m(d, \beta) = E_0 + \delta \log(d + off)$$

and *off* is a fixed offset parameter.**Usage**`linlog(d, e, off)`**Arguments**

<i>d</i>	real-valued argument to the function (dose variable)
<i>e</i>	model parameter
<i>off</i>	fixed offset parameter

Value

Response value.

logistic*Implementation of logistic models***Description**

Logistic model:

$$m(d, \beta) = E_0 + \frac{E_{max}}{1 + exp[(ED_{50} - d)/\delta]}$$

Usage`logistic(d, e)`**Arguments**

<i>d</i>	real-valued argument to the function (dose variable)
<i>e</i>	model parameter

Value

Response value.

quadratic

*Implementation of quadratic models***Description**

Quadratic model:

$$m(d, \beta) = E_0 + \beta_1 d + \beta_2 d^2$$

Usage`quadratic(d, e)`**Arguments**

- | | |
|---|--|
| d | real-valued argument to the function (dose variable) |
| e | model parameter |

Value

Response value.

sigEmax

*Implementation of Sigmoid Emax models***Description**

Sigmoid Emax Model model:

$$m(d, \beta) = E_0 + E_{max} \frac{d^h}{ED_{50}^h + d^h}$$

Usage`sigEmax(d, e)`**Arguments**

- | | |
|---|--|
| d | real-valued argument to the function (dose variable) |
| e | model parameter |

Value

Response value

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