Package ‘drcSeedGerm’

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Description Utility functions to be used to analyse datasets obtained from seed germination/emergence assays. Fits several types of seed germination/emergence models, including those reported in Onofri et al. (2018) "Hydrothermal-time-to-event models for seed germination", European Journal of Agronomy, 101, 129-139 <doi:10.1016/j.eja.2018.08.011>. Contains several datasets for practicing.
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Description
The germination of alfalfa was assayed at 7 temperature levels, on three replicated Petri dishes per temperature and 100 seeds per Petri dish. Inspections were made in several times after the beginning of the assay.

Usage

data("alfalfaSG")

Format
A data frame with 21 observations on the following variables.

Dish a numeric vector with the coding for Petri dishes
Temp a numeric vector with the temperature level
nViable a numeric vector: number of viable seeds per dish
‘1’ a numeric vector: count at day 1
Details

Every line of data represents a Petri dish. There were 100 seeds per Petri dish. The columns represent the characteristics of each Petri dish. The columns from 4 to 32 represent the number of germinated seeds counted at each assessment time.

Author(s)

Andrea Onofri

Source

no reference yet
References


Examples

data(alfalfaSG)
head(alfalfaSG)

barley
A series of germination assays with barley

Description

The germination of barley was assayed at 9 temperature levels, on three replicated Petri dishes at each temperature and 50 seeds per Petri dish. Inspections were made in several times after the beginning of the assay.

Usage

data("barley")

Format

A data frame with 810 observations on the following 7 variables.

- **Dish**: a numeric vector with the coding for Petri dishes
- **Temp**: a numeric vector with the temperature level
- **timeBef**: a numeric vector, with the start time for each inspection interval
- **timeAf**: a numeric vector, with the end time for each inspection interval
- **nSeeds**: a numeric vector, with the number of germinated seeds at each inspection interval
- **nCum**: a numeric vector, with the cumulative number of germinated seeds at each assessment time
- **propCum**: a numeric vector, the cumulative proportion of germinated seeds at each assessment time

Details

The variable `timeAf` contains the value `Inf` (Infinity), that corresponds to the seeds which did not germinate during the assay, for which we the germination time might be comprised from the last assessment time to infinity.

Author(s)

Andrea Onofri

Source

no reference yet
References


Examples

data(barley)

---

excalibur Relationship between germination rate and water potential in oilseed rape (var. Excalibur)

Description

This file describes the relationship between germination rate and water potential in the substrate for seeds of oilseed rape (var. Excalibur). Three germination percentiles are considered for germination rate, i.e. GR10, GR30 and GR50.

Usage

data("excalibur")

Format

A data frame with 27 observations on the following 5 variables.

Perc  a numeric vector: the germination percentile
Psi   a numeric vector: water potential in the substrate (in MPa)
Tg    a numeric vector: germination time in days
SE    a numeric vector: standard errors for germination times (in days)
GR    a numeric vector: germination rates in d^-1

Source

Pace, R., Benincasa, P., Ghanem, M.E., Quinet, M., Lutts, S., 2012. GERMINATION OF UNTREATED AND PRIMED SEEDS IN RAPESEED (BRASSICA NAPUS VAR OLEIFERA DEL.) UNDER SALINITY AND LOW MATRIC POTENTIAL. Experimental Agriculture 48, 238–251.

References

Examples

```r
data(excalibur)
head(excalibur)
library(drc)

modGR1 <- drm(GR ~ Psi, fct=GRPsiLin(), data=excalibur, curveid=Perc)
summary(modGR1)
plot(modGR1, log="", legendPos=c(-0.9, 1))

modGR2 <- drm(GR ~ Psi, fct=GRPsiPol2(), data=excalibur, curveid=Perc)
summary(modGR2)
plot(modGR2, log="", legendPos=c(-0.9, 1))
```

---

**festuca**  
*Relationship between germination rate and water potential in Festuca arundinacea L.*

**Description**

This file describes the relationship between germination rate and Festuca arundinacea. Three germination percentiles are considered for germination rate, i.e. GR10, GR30 and GR50.

**Usage**

```r
data("festuca")
```

**Format**

A data frame with 36 observations on the following 3 variables.

- `g` a numeric vector: the germination percentile
- `Psi` a numeric vector: water potential in the substrate (in MPa)
- `GR` a numeric vector: germination rates in d^-1

**Source**


**References**

**GR_Psi_models**

**Examples**

```r
data(festuca)
modGR1 <- drm(GR ~ Psi, fct=GRPsiLin(),
              data=festuca, curveid=g)
summary(modGR1)
```

---

**GR_Psi_models**

*Hydro-time models based on the germination rate*

**Description**

These models describe the response of germination rate to water potential in the substrate.

**Usage**

- `GRPsiLin()`
- `GRPsiPol()`
- `GRPsiPol2()`
- `GRPsi.Lin()`
- `GRPsi.Pol()`
- `GRPsi.Pol2()`
- `GRPsiLin.fun(Psi, Psib, thetaH)`
- `GRPsiPol.fun(Psi, Psib, thetaH)`
- `GRPsiPol2.fun(Psi, Psib, thetaH)`

**Arguments**

The functions `GRPsiLin()`, `GRPsiPol()`, `GRPsiPol2()`, `GRPsi.Lin()`, `GRPsi.Pol()`, `GRPsi.Pol2()` have no arguments. The functions `GRPsiLin.fun()`, `GRPsiPol.fun()` and `GRPsiPol2.fun()` have the following arguments:

- `Psi` Water potential in the substrate
- `Psib` Base water potential within the population
- `thetaH` Hydro-time parameter

**Details**

The functions `GRPsiLin()`, `GRPsiPol()`, `GRPsiPol2()`, `GRPsi.Lin()`, `GRPsi.Pol()`, `GRPsi.Pol2()` are meant to be used with the `drm()` function in the `drc` package (`GRPsiLin()` and `GRPsi.Lin()`, `GRPsiPol()` and `GRPsi.Pol()`, `GRPsiPol2()` and `GRPsiPol2()` are totally equivalent, apart from the names). The functions `GRPsiLin.fun()`, `GRPsiPol.fun()` and `GRPsiPol2.fun()` are used for general purposes (plotting and other uses). Details about these functions and the meaning of parameters are described in Bradford (2002) and in the package documentation (see references below).
Value

The 'GRPsiLin.fun()', 'GRPsiPol.fun()' and 'GRPsiPol2.fun()' functions return the germination rate for any given values of water potential in the substrate. The 'GRPsiLin()', 'GRPsiPol()' and 'GRPsiPol2()' (and 'GRPsiLin()', 'GRPsiPol()' and 'GRPsiPol2()') functions return a list containing the nonlinear function, the self starter function, the parameter names and other items which are internally used by the 'drc()' function.

Author(s)

Andrea Onofri

References

See package documentation at: https://www.statforbiology.com/_seedtutorial/

Examples

library(drcte)
# Observed data
Psi <- c(-2, -1.5, -1.2, -1, -0.8, -0.6, -0.4, -0.25, -0.12, -0.06, -0.03, 0)
GR <- c(0, 0, 0, 0, 0.0585, 0.094, 0.1231, 0.1351, 0.1418, 0.1453, 0.1458, 0.1459)
Psi2 <- c(-0.5, -0.6, -0.7, -0.8, -0.9, -1, -1.1, -1.2, -1.5)
GR2 <- c(1.4018, 1.0071, 0.5614, 0.3546, 0.2293, 0, 0, 0, 0, 0, 0, 0)

# Model fitting
modHT1 <- drm(GR ~ Psi, fct = GRPsiLin())
modHT2 <- drm(GR ~ Psi, fct = GRPsiPol())
modHT3 <- drm(GR ~ Psi2, fct = GRPsiPol2())
summary(modHT1)
summary(modHT2)
summary(modHT3)
**Usage**

```
GRTPsi.M()
GRTPsi.M.fun(Temp, Psi, k, Tb, ThetaHT, Psib)
GRTPsi.BS()
GRTPsi.BS.fun(Temp, Psi, k, Tb, To, ThetaHT, Psib)
```

**Arguments**

- `Temp` Temperature variable
- `Psi` Water potential variable
- `Psib` Base water potential variable
- `k` regression parameter
- `Tb` base temperature
- `To` optimal temperature
- `ThetaHT` Hydro-thermal-time parameter

**Details**

The 'GRT.M.fun()' is a generic R function, while the GRT.M() function is meant to be used with the 'drm()' function, within the 'drc' package.

**Value**

The 'GRT.M.fun()' functions returns a vector of responses, for given values of temperature, Tc, Tb and ThetaH. The GRT.M() function returns a list containing the nonlinear function, the self starter function, the parameter names and other items which are internally used by the 'drm()' function.

**Author(s)**

Andrea Onofri

**References**


**Examples**

```r
library(drcte)
Tval <- c(2, 5, 10, 15, 20, 25, 30, 35, 40)
GR <- c(0, 0, 0.209, 0.435, 0.759, 0.821, 0.417, 0.145, 0)
modM <- drm(GR ~ Tval, fct = GRT.M())
plot(modM, log="", xlim = c(0, 40), ylim=c(0,1.2),
     legendPos = c(5, 1.0), xlab = "Temperature (°C)"
```


GR_T_models

Thermal-time models based on the germination rate

Description

These models are used to describe the germination rate of a seed, depending on the environmental temperature.

Usage

- GRT.GH()
- GRT.GH2()
- GRT.YL()
- GRT.BS()
- GRT.BSb()
- GRT.Ex()
- GRT.Exb()
- GRT.M()
- GRT.Mb()
- GRT.RF()
- GRT.RFb()
- GRT.GH.fun(Temp, Tb, ThetaT)
- GRT.GH2.fun(Temp, Tb, beta)
- GRT.YL.fun
- GRT.BS.fun(Temp, k, Tb, To, ThetaT)
- GRT.BSb.fun(Temp, Tc, Tb, To, ThetaT)
- GRT.Ex.fun(Temp, k, Tb, Tc, ThetaT)
- GRT.Exb.fun(Temp, k, Tb, Tc, ThetaT)
- GRT.M.fun(Temp, k, Tb, ThetaT)
- GRT.Mb.fun(Temp, Tc, Tb, ThetaT)
- GRT.RF.fun(Temp, k, Tb, Td, ThetaT)
- GRT.RFb.fun(Temp, Tc, Tb, Td, ThetaT)

Arguments

The 'GR.funName()' functions have no arguments. The general purpose 'GR.funName.fun()' functions have some of the following arguments (depending on function):

- **Temp**: Temperature variable
- **k**: regression parameter
- **beta**: regression parameter
- **Tc**: ceiling temperature
- **Tb**: base temperature
- **To**: optimal temperature
- **Td**: close-to-optimal temperature (Rowse-Fintch-Savage equation)
- **ThetaT**: Hydro-time parameter
Details

All these functions are named according to this rule: 'GRT' (Germination Rate Temperature), followed by the 'function name' (e.g., BS, RF, M, Ex, YL). The R functions 'GR.funName().fun' are generic R function, that are meant to be used for general purposes, such as plotting or predicting. The corresponding 'GR.funName()' (without the '.fun' ending) are meant to be used for fitting with the 'drm()' function, within the 'drc' package.

Value

The 'GRT.funName.fun()' functions return a vector of responses, for given values of temperature and parameters. The 'GRT.funName()' functions return a list containing the nonlinear function, the self starter function, the parameter names and other items which are internally used by the 'drm()' function.

Author(s)

Andrea Onofri

References


Examples

```r
library(drcte)
Tval <- c(2, 5, 10, 15, 20, 25, 30, 35, 40)
GR <- c(0, 0, 0.209, 0.435, 0.759, 0.821, 0.417, 0.145, 0)
modM <- drm(GR ~ Tval, fct = GRT.M())
plot(modM, log="", xlim = c(0, 40), ylim=c(0,1.2),
       legendPos = c(5, 1.0), xlab = "Temperature (°C)")
```

hordeum

Germination of Hordeum spontaneum at different temperatures and water potentials

Description

This dataset was obtained from a germination assay with four replicated Petri dishes with 20 seeds, tested at six different water potential levels (0, -0.3, -0.6, -0.9, -1.2 and -1.5 MPa). Osmotic potentials were produced using variable amount of polyethylene glycol (PEG, molecular weight 8000) adjusted for the temperature level. Petri dishes were incubated at six constant temperature levels (8, 12, 16, 20, 24 and 28 °C), under a photoperiod of 12 h. Germinated seeds (radicle protrusion > 3 mm) were counted and removed daily for 20 days.

Usage

data("hordeum")
Format
A data frame with 3024 observations on the following 8 variables.

- **temp**: a numeric vector: temperature level
- **water**: a numeric vector: water potential level
- **Dish**: a numeric vector: code for Petri dishes
- **timeBef**: a numeric vector: beginning of scoring interval
- **timeAf**: a numeric vector: end of scoring interval
- **nViable**: a numeric vector: number of viable seeds at the beginning of assay, in each dish
- **nSeeds**: a numeric vector: number of germinated seeds, between timeBef and timeAf
- **nCum**: a numeric vector: cumulative number of germinated seeds at timeAf

Details
This dataset was analysed in the time-to-event framework in Onofri et al (2018). See Example 2.

Source

References

Examples
```r
# Fitting a hydrotime model
data(rape)
gmod <- drmte(nSeeds ~ timeBef + timeAf + Psi,
              fct=HTnorm(), data=rape)
summary(gmod)

jackGroupSE(gmod, rape, rape$Dish)
```
Hydrotime model with log-logistic distribution of germination time
(Onofri et al., 2018)

Description

This model relates the time-course of the proportion of germinated seeds to the water potential in the substrate. It is based on a truncated log-logistic distribution of germination time:

\[ P(t) = \frac{d}{1 + e^x \left[ b(\log(x) - \log(e)) \right]} \]

where two of the three usual parameters ('d' and 'e') are expressed as functions of water potential (\( \Psi \)). In this function, the two submodels are: (1) for the parameter 'd', we used a shifted exponential function:

\[ d = G \left[ 1 - \exp \left( \frac{\Psi - \Psi_b}{\sigma_{\Psi_b}} \right) \right] \]

while, (2) for the parameter 'e' we considered that its inverse corresponds to the median Germination Rate within the population (i.e. \( 1/e = GR_{50} \)) and modelled this latter parameter as:

\[ GR_{50} = \frac{\Psi - \Psi_b}{\theta_H} \]

The 'HTE1.fun()' is a generic function, which can be used for plotting or other applications, while the 'HTE1()' function is meant to be used for model fitting with the 'drmte()' function in the 'drcte()' package.

Usage

\[
\text{HTE1}(\text{fixed} = c(\text{NA}, \text{NA}, \text{NA}, \text{NA}, \text{NA}), \\
\text{names} = c(\text{"G"}, \text{"Psib"}, \text{"sigmaPsib"}, \text{"thetaH"}, \text{"b"})) \\
\text{HTE1.fun}(\text{time}, \text{Psi}, \text{G}, \text{Psib}, \text{sigmaPsib}, \text{thetaH}, \text{b})
\]

Arguments

These functions have the following arguments:

- **fixed**: numeric vector. Specifies which parameters are fixed and at what value they are fixed. NAs for parameter that are not fixed. At the moment, only the parameter G can be held fixed.
- **names**: a vector of character strings giving the names of the parameters. The default is reasonable.
- **time**: time
- **Psi**: water potential in the substrate
- **G**: maximum germination capability of the seed lot
Psib  base water potential
sigmaPsib  standard deviation for the base water potential within the seed lot
thetaH  hydro-time constant
b  shape parameter for the cumulative distribution function of germination time

Details

The detail of this time-to-event model and the meaning of parameters are described in Onofri et al. (2018). See Table 2, where 'HTE1()' is abbreviated as HTE.

Value

The 'HTE1.fun()' function returns the proportion of germinated seeds, for any given values of time and water potential in the substrate. The 'HTE1()' function returns a list containing the nonlinear function, the self-starter function, the parameter names and other items which are internally used by the 'drmte()' function.

Author(s)

Andrea Onofri

References


Examples

data(rape)
modHTE <- drmte( nSeeds ~ timeBef + timeAf + Psi, data=rape, fct=HTE1())
summary(modHTE)

HTE2
Hydrotime model with log-logistic distribution of germination time
(Onofri et al., 2018)

Description

This model relates the time-course of the proportion of germinated seeds to the water potential in the substrate. It is based on a truncated log-logistic distribution of germination time:

\[ P(t) = \frac{d}{1 + exp\left[b\log(x) - \log(e)\right]} \]
where two of the three usual parameters (‘d’ and ‘e’) are expressed as functions of water potential ($\Psi$). In this function, the two submodels are: (1) for the parameter ‘d’, we used a shifted exponential function:

$$d = G \left[ 1 - \exp \left( \frac{\Psi - \Psi_b}{\sigma_{\Psi_b}} \right) \right]$$

while, (2) for the parameter ‘e’ we considered that its inverse corresponds to the median Germination Rate within the population (i.e. $1/e = GR_{50}$) and modelled this latter parameter as:

$$GR_{50} = \frac{[\Psi - \Psi_b]^2}{\theta_H}$$

The difference with the 'HTE1()' function is that, in this case, the relationship between GR50 and water potential is not linear, but curvilinear (convex down). The 'HTE2.fun()' is a generic function, which can be used for plotting or other applications, while the 'HTE2()' function is meant to be used for model fitting with the 'drmte()' function in the 'drcte()' package.

Usage

```r
HTE2(fixed = c(NA, NA, NA, NA, NA),
      names = c("G", "Psib", "sigmaPsib", "thetaH", "b"))
HTE2.fun(time, Psi, G, Psib, sigmaPsib, thetaH, b)
```

Arguments

These functions have the following arguments:

- `fixed`: numeric vector. Specifies which parameters are fixed and at what value they are fixed. NAs for parameter that are not fixed. At the moment, only the parameter G can be held fixed.
- `names`: a vector of character strings giving the names of the parameters. The default is reasonable.
- `time`: time
- `Psi`: water potential in the substrate
- `G`: maximum germination capability of the seed lot
- `Psib`: base water potential
- `sigmaPsib`: standard deviation for the base water potential within the seed lot
- `thetaH`: hydro-time constant
- `b`: shape parameter for the cumulative distribution function of germination time

Details

No more detail, at the moment.
Value

The ’HTE2.fun()’ function returns the proportion of germinated seeds, for any given values of time and water potential in the substrate, depending on model parameters G, Psib, sigmaPsib, thetaH and b. The ’HTE2()’ function returns a list containing the nonlinear function, the self starter function, the parameter names and other items which are internally used by the ’drmte()’ function.

Author(s)

Andrea Onofri

References


Examples

# Fitting model
data(rape)
modHTE2 <- drmte( nSeeds ~ timeBef + timeAf + Psi,
data=rape, fct=HTE2())
summary(modHTE2)

Description

This model relates the time-course of the proportion of germinated seeds to the water potential in the substrate. It is based on a truncated log-logistic distribution of germination time:

\[ P(t) = \frac{d}{1 + \exp[b(\log(x) - \log(e))] \]  

where two of the three usual parameters (’d’ and ’e’) are expressed as functions of water potential (Ψ). In this function, the two submodels are: (1) for the parameter ’d’, we used a shifted exponential function:

\[ d = G \left[ 1 - \exp \left( \frac{\Psi - \Psi_b}{\sigma_{\Psi_b}} \right) \right] \] 

while, (2) for the parameter ’e’ we considered that its inverse corresponds to the median Germination Rate within the population (i.e. 1/e = GR50) and modelled this latter parameter as:

\[ GR_{50} = \frac{\Psi^2 - \Psi_b^2}{\theta_H} \]
The difference with the 'HTE1()' function is that, in this case, the relationship between GR50 and water potential is not linear, but curvilinear (convex up). The 'HTE3.fun()' is a generic function, which can be used for plotting or other applications, while the 'HTE3()' function is meant to be used for model fitting with the 'drmte()' function in the 'drcte()' package.

Usage

HTE3(fixed = c(NA, NA, NA, NA, NA),
    names = c("G", "Psib", "sigmaPsib", "thetaH", "b"))
HTE3.fun(time, Psi, G, Psib, sigmaPsib, thetaH, b)

Arguments

These functions have the following arguments:

- **fixed**: numeric vector. Specifies which parameters are fixed and at what value they are fixed. NAs for parameter that are not fixed. At the moment, only the parameter G can be held fixed.
- **names**: a vector of character strings giving the names of the parameters. The default is reasonable.
- **time**: time
- **Psi**: water potential in the substrate
- **G**: maximum germination capability of the seed lot
- **Psib**: base water potential
- **sigmaPsib**: standard deviation for the base water potential within the seed lot
- **thetaH**: hydro-time constant
- **b**: shape parameter for the cumulative distribution function of germination time

Details

No more detail, at the moment.

Value

The 'HTE3.fun()' function returns the proportion of germinated seeds, for any given values of time and water potential in the substrate, depending on model parameters G, Psib, sigmaPsib, thetaH and b. The 'HTE3()' function returns a list containing the nonlinear function, the self starter function, the parameter names and other items which are internally used by the 'drmte()' function.

Author(s)

Andrea Onofri

References

Examples

# Fitting model
data(rape)
modHTE3 <- drmte( nSeeds ~ timeBef + timeAf + Psi,
data = rape, fct=HTE3())
summary(modHTE3)

HTex

Hydrotime model with Type II Extreme Value distribution of base water potential (Mesgaran et al., 2013)

Description

This model relates the time-course of the proportion of germinated seeds to the water potential in the substrate. It is based on Type II Extreme Value distribution of base water potential within the seed lot. The equation is:

\[
P(t) = 1 - \exp \left\{ -\exp \left[ \frac{\Psi - \left(\frac{\theta_H}{t}\right) - \mu}{\sigma} \right] \right\}
\]

In contrast to other hydrotime models (e.g., 'HTE1()', 'HTE2()', 'HTE3()', describing the distribution of germination time), this models describes the distribution of base water potential within the population, while the distribution of germination times is indirectly modelled, but it is not, in itself, gaussian (you see that 't' is at the denominator). The 'HTex.fun()' is a generic function, which can be used for plotting or other applications, while the 'HTex()' function is meant to be used for model fitting with the 'drmte()' function in the 'drcte()' package.

Usage

HTex()
HTex.fun(time, Psi, thetaH, mu, sigma)

Arguments

The 'HTex()' function has no arguments. The 'HTex.fun()' function has the following arguments:

time
time
Psi water potential in the substrate
thetaH hydro-time constant
mu location parameter for the Type II Extreme Value distribution
sigma scale parameter for the Type II Extreme Value distribution

Details

The detail of this time-to-event model and the meaning of parameters are described in Mesgaran et al. (2013).
Value

The 'HTex.fun()' function returns the proportion of germinated seeds, for any given values of time and water potential in the substrate. The 'HTex()' function returns a list containing the nonlinear function, the self-starter function, the parameter names and other items which are internally used by the 'drmte()' function.

Note

This function is for use with the function 'drmte()'

Author(s)

Andrea Onofri

References


Examples

data(rape)
modg <- drm(propCum ~ timeAf + Psi, fct=HTex(), data=rape)
summary(modg)

---

HTG  
Hydrotime model with Gumbel distribution of base water potential  
(Mesgaran et al., 2013)

Description

This model relates the time-course of the proportion of germinated seeds to the water potential in the substrate. It is based on a Gumbel distribution of base water potential within the seed lot. The equation is:

\[ P(t) = \exp\left\{- \exp\left[- \left( \frac{\Psi - (\theta_H/t) - \mu}{\sigma} \right) \right]\right\} \]

In contrast to other hydrotime models (e.g., 'HTE1()', 'HTE2()', 'HTE3()', describing the distribution of germination time), this model describes the distribution of base water potential within the population, while the distribution of germination times is indirectly modelled, but it is not, in itself, Gaussian (you see that 't' is at the denominator). The 'HTG.fun()' is a generic function, which can be used for plotting or other applications, while the 'HTG()' function is meant to be used for model fitting with the 'drmte()' function in the 'drcte()' package.
Usage

HTG()
HTG.fun(time, Psi, ThetaH, mu, sigma)

Arguments

The 'HTG()' function has no arguments. The 'HTG.fun()' function has the following arguments:

- **time**
- **Psi** water potential in the substrate
- **ThetaH** hydro-time constant
- **mu** location parameter for the Gumbel distribution
- **sigma** scale parameter for the Gumbel distribution

Details

The detail of this time-to-event model and the meaning of parameters are described in Mesgaran et al. (2013).

Value

The 'HTG.fun()' function returns the proportion of germinated seeds, for any given values of time and water potential in the substrate. The 'HTG()' function returns a list containing the nonlinear function, the self-starter function, the parameter names and other items which are internally used by the 'drmte()' function.

Note

This function is for use with the function 'drmte()'

Author(s)

Andrea Onofri

References


Examples

data(rape)
modg <- drm(propCum ~ timeAf + Psi, fct=HTG(), data=rape)
summary(modg)
Hydrotime model with logistic distribution of base water potential
(Mesgaran et al., 2013)

Description
This model relates the time-course of the proportion of germinated seeds to the water potential in the substrate. It is based on a logistic distribution of base water potential within the seed lot. The equation is:

\[
P(t) = \frac{1}{1 + e^{\frac{-\Psi - (\theta H/t) - \Psi_b}{\sigma}}}
\]

In contrast to other hydrotime models (e.g., 'HTE1()', 'HTE2()', 'HTE3()', describing the distribution of germination time), this model describes the distribution of base water potential within the population, while the distribution of germination times is indirectly modelled, but it is not, in itself, gaussian (you see that ‘t’ is at the denominator). The 'HTL.fun()' is a generic function, which can be used for plotting or other applications, while the 'HTL()' function is meant to be used for model fitting with the 'drmte()' function in the 'drcte()' package.

Usage

HTL()
HTL.fun(time, Psi, ThetaH, Psib50, sigma)

Arguments
The 'HTL()' function has no arguments. The 'HTL.fun()' function has the following arguments:

time  time
Psi   water potential in the substrate
ThetaH hydro-time constant
Psib50 median base water potential
sigma scale parameter for the logistic distribution

Details
The detail of this time-to-event model and the meaning of parameters are described in Mesgaran et al. (2013).

Value
The 'HTL.fun()' function returns the proportion of germinated seeds, for any given values of time and water potential in the substrate. The 'HTL()' function returns a list containing the nonlinear function, the self-starter function, the parameter names and other items which are internally used by the 'drmte()' function.
Note
This function is for use with the function 'drmte()'

Author(s)
Andrea Onofri

References

Examples
```
data(rape)
modg <- drm(propCum ~ timeAf + Psi, fct=HTL(), data=rape)
summary(modg)
```

HTLL

Hydrotime model with log-logistic distribution of base water potential
(Mesgaran et al., 2013)

Description
This model relates the time-course of the proportion of germinated seeds to the water potential in the substrate. It is based on a log-logistic distribution of base water potential within the seed lot. The equation is:

\[
P(t) = \frac{1}{1 + \exp \left\{ \frac{\log \left( \frac{\Psi - \left( \frac{\Psi_b}{t} \right) + \delta}{\sigma} \right) - \log(\Psi_b + \delta)}{\sigma} \right\}}
\]

In contrast to other hydrotimes (e.g., 'HTE1()', 'HTE2()', 'HTE3()', describing the distribution of germination time), this models describes the distribution of base water potential within the population, while the distribution of germination times is indirectly modelled, but it is not, in itself, gaussian (you see that 't' is at the denominator). The 'HTLL.fun()' is a generic function, which can be used for plotting or other applications, while the 'HTLL()' function is meant to be used for model fitting with the 'drmte()' function in the 'drcte()' package.

Usage
```
HTLL()
HTLL.fun(time, Psi, thetaH, delta, Psib50, sigma)
```
Arguments

The 'HTLL()' function has no arguments. The 'HTLL.fun()' function has the following arguments:

- `time`: time
- `Psi`: water potential in the substrate
- `thetaH`: hydro-time constant
- `delta`: shifting parameter to allow for negative water potential levels
- `Psib50`: median base water potential
- `sigma`: scale parameter for the log-logistic distribution

Details

The detail of this time-to-event model and the meaning of parameters are described in Mesgaran et al. (2013).

Value

The 'HTLL.fun()' function returns the proportion of germinated seeds, for any given values of time and water potential in the substrate. The 'HTLL()' function returns a list containing the nonlinear function, the self-starter function, the parameter names and other items which are internally used by the 'drmte()' function.

Note

This function is for use with the function 'drmte()'

Author(s)

Andrea Onofri

References


Examples

```r
data(rape)
modg <- drm( propCum ~ timeAf + Psi, fct=HTLL(), data=rape)
summary(modg)
```
Hydrotime model with normal distribution of base water potential (Bradford, 2002)

Description

This model relates the time-course of the proportion of germinated seeds to the water potential in the substrate. It is based on a normal distribution of base water potential within the seed lot. The equation is:

\[
P(t) = \Phi \left\{ \frac{\Psi - (\theta_H/t) - \Psi_b}{\sigma_{\Psi_b}} \right\}
\]

where \( \Phi \) is a gaussian cumulative distribution function for base water potential. In contrast to other hydrotime models (e.g., 'HTE1()', 'HTE2()', 'HTE3()', describing the distribution of germination time), this models describes the distribution of base water potential within the population, while the distribution of germination times is indirectly modelled, but it is not, in itself, gaussian (you see that 't' is at the denominator). The 'HTNorm.fun()' is a generic function, which can be used for plotting or other applications, while the 'HTNorm()' function is meant to be used for model fitting with the 'drmte()' function in the 'drcte()' package.

Usage

HTnorm()

HTnorm.fun(time, Psi, ThetaH, Psib50, sigmaPsib)

Arguments

The 'HTNorm()' function has no arguments. The 'HTNorm.fun()' function has the following arguments:

time time
Psi water potential in the substrate
ThetaH hydro-time constant
Psib50 median base water potential
sigmaPsib standard deviation for the base water potential within the seed lot

Details

The detail of this time-to-event model and the meaning of parameters are described in Bradford (2002).

Value

The 'HTNorm.fun()' function returns the proportion of germinated seeds, for any given values of time and water potential in the substrate. The 'HTNorm()' function returns a list containing the nonlinear function, the self-starter function, the parameter names and other items which are internally used by the 'drmte()' function.
Note
This function is for use with the R function 'drmte()'

Author(s)
Andrea Onofri

References

Examples
```r
data(rape)
modg <- drm( propCum ~ timeAf + Psi, fct=HTnorm(), data=rape)
summary(modg)
```

Description
This model relates the time-course of the proportion of germinated seeds to the water potential and temperature in the substrate. It is based on a truncated log-logistic distribution of germination time:

\[
P(t) = \frac{d}{1 + \exp\left[b(\log(x) - \log(e))\right]}
\]

where two of the three usual parameters (‘d’ and ‘e’) are expressed as functions of water potential (Ψ) and temperature (T). In the function ‘HTTEM()’, we implemented the following submodels: (1) for the parameter ‘d’, we implemented a shifted exponential function:

\[
d = G \left[1 - \exp\left(\frac{\Psi - \Psi_b - k(T - T_b)}{\sigma_{\Psi_b}}\right)\right]
\]

while, (2) for the parameter ‘e’ we considered that its inverse corresponds to the median Germination Rate within the population (i.e. \(1/e = GR_{50}\)) and modelled this latter parameter as:

\[
GR_{50} = \frac{T - T_b}{\theta_{HT}} [\Psi - \Psi_b - k(T - T_b)]
\]

The ‘HTTEM.fun()’ is a generic function, which can be used for plotting or other applications, while the ‘HTTEM()’ function is meant to be used for model fitting with the ‘drmte()’ function in the ‘drcte()’ package.
Usage

`HTTEM()`

`HTTEM.fun(time, Psi, Temp, G, Psib, kt, Tb, sigmaPsib, ThetaHT, b)`

Arguments

The 'HTTEM' function has no arguments. The 'HTTEM.fun()' has the following arguments:

- `time`: time
- `Psi`: water potential in the substrate
- `Temp`: temperature
- `G`: maximum germination capability of the seed lot
- `Psib`: base water potential
- `kt`: parameter measuring the effect of temperature on base water potential
- `Tb`: base temperature
- `sigmaPsib`: standard deviation for the base water potential within the seed lot
- `ThetaHT`: hydro-thermal-time parameter
- `b`: shape parameter for the cumulative distribution function of germination time

Details

The detail of this time-to-event model and the meaning of parameters are described in Onofri et al. (2018). See Table 2, where 'HTTEM()' is abbreviated as HTTE.

Value

The 'HTTEM.fun()' function returns the proportion of germinated seeds, for any given values of time and water potential in the substrate. The 'HTTEM()' function returns a list containing the nonlinear function, the parameter names and other items which are internally used by the 'drmte()' function. At the moment, there is no self-starting function and starting parameters for fitting must be provided within the 'drcte' function.

Author(s)

Andrea Onofri

References

Examples

```r
data(hordeum)

modHTTEM <- drmte(nSeeds ~ timeBef + timeAf + water + temp,
                  data=hordeum,
                  fct = HTTEM(),
                  start=c(0.8,-2, 0.05, 3, 0.2, 2000, 0.5))
summary(modHTTEM)
```

---

**HTTLL**

*Hydro-thermal-time model with log-logistic distribution of base water potential (Mesgaran et al., 2017)*

**Description**

This model relates the time-course of the proportion of germinated seeds to the water potential and temperature in the substrate and it is based on a log-logistic distribution of base water potential within the seed lot. Two similar functions are available within the ‘drcSeedGerm’ package: the first one is ‘HTTLL.M()’ that assumes that the base water potential decreases with temperature for any \( T > T_b \). The equation is:

\[
P(t, T, \Psi) = \frac{1}{1 + \exp\left\{-\frac{\log\left[\Psi - \left(\frac{\theta_{HT} T}{T - T_b}\right) + \delta\right] - \log[\Psi_b + K_t(T - T_b) + \delta]}{\sigma_{\Psi_b}}\right\}}
\]

The second function is ‘HTTLL.BS()’ and it assumes that the base water potential decreases with temperature only for any \( T > T_o \), where \( T_o \) is the optimal temperature level. For this case, the element \( K_t(T - T_b) \) is modified as \( K_t[\max(T, T_o) - T_o] \) and the optimal temperature \( T_o \) is included as an explicit parameter.

The ‘HTTLL.M.fun()’ and ‘HTTLL.BS.fun()’ are two generic functions, which can be used for plotting or other applications, while the ‘HTTLL.M()’ and ‘HTTLL.BS()’ functions are meant to be used for model fitting with the ‘drmte()’ function in the ‘drcte()’ package.

**Usage**

```r
HTTLL.M()
HTTLL.BS()
HTTLL.M.fun(time, Psi, Temp, thetaHT, Tb, Psib50, Kt, delta, sigmaPsib)
HTTLL.BS.fun(time, Psi, Temp, thetaHT, Tb, To, Psib50, Kt, delta, sigmaPsib)
```

**Arguments**

The ‘HTTLL.M()’ and ‘HTTLL.BS()’ functions have no arguments. The ‘HTTLL.M.fun()’ and the ‘HTTLL.BS.fun()’ functions have the following arguments:

- `time`
<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Psi</td>
<td>water potential in the substrate</td>
</tr>
<tr>
<td>Temp</td>
<td>temperature</td>
</tr>
<tr>
<td>thetaHT</td>
<td>hydro-time parameter</td>
</tr>
<tr>
<td>Tb</td>
<td>base temperature</td>
</tr>
<tr>
<td>To</td>
<td>optimal temperature</td>
</tr>
<tr>
<td>Psib50</td>
<td>median base water potential</td>
</tr>
<tr>
<td>Kt</td>
<td>parameter measuring the effect of temperature on base water potential</td>
</tr>
<tr>
<td>delta</td>
<td>shifting parameter to allow for a log-logistic distribution of negative values for base water potential</td>
</tr>
<tr>
<td>sigmaPsib</td>
<td>scale parameter for log-logistic distribution</td>
</tr>
</tbody>
</table>

**Details**

The detail of this time-to-event model and the meaning of parameters are described in Mesgaran et al. (2017).

**Value**

The 'HTTLL.M.fun()' and 'HTTLL.BS.fun()' functions return the proportion of germinated seeds, for any given values of time, water potential and temperature in the substrate. The 'HTTLL.M()' and 'HTTLL.BS()' functions return a list containing the nonlinear function, the self-starter function, the parameter names and other items which are internally used by the 'drmte()' function.

**Note**

This function is for use with the function 'drmte()'

**Author(s)**

Andrea Onofri

**References**


**Examples**

data(hordeum)

modHTTLL.M <- drmte(nSeeds ~ timeBef + timeAf + water + temp, 
data=hordeum, 
   fct = HTTLL.M(), 
   start=c(832,-2.5, -3, 0.07, 3, 0.5))

summary(modHTTLL.M)

modHTTLL.BS <- drmte(nSeeds ~ timeBef + timeAf + water + temp, 
data=hordeum, 
   fct = HTTLL.BS(), 
   start=c(832,-2.5, -3, 0.07, 3, 0.5))

summary(modHTTLL.BS)
fct = HTTLL.BS()
start=c(932,-2.5, 15, -3, 0.07, 3, 0.5))
summary(modHTTLL.BS)

\[ P(t, T, \Psi) = \Phi\left( \frac{\Psi - \left[ \frac{\Psi_{b}}{t(T-T_{b})} \right] - [\Psi_{b} + K_{t}(T-T_{b})]}{\sigma_{\Psi_{b}}} \right) \]

where \( \Phi \) is a gaussian cumulative distribution function for base water potential.

The second function is 'HTTNorm.BS()' and it assumes that the base water potential decreases with temperature only for any \( T > T_{o} \), where \( T_{o} \) is the optimal temperature level. For this case, the element \( K_{t}(T-T_{b}) \) is modified as \( K_{t} \{ \text{max}(T, T_{o}) - T_{o} \} \) and the optimal temperature \( T_{o} \) is included as an explicit parameter.

The 'HTTNorm.M.fun()' and 'HTTNorm.BS.fun()' are two generic functions, which can be used for plotting or other applications, while the 'HTTNorm.M()' and 'HTTNorm.BS()' functions are meant to be used for model fitting with the 'drmte()' function in the 'drcte()' package.

Usage

HTTNorm.M()
HTTNorm.BS()
HTTNorm.M.fun(time, psi, temp, thetaHT, Tb, Psib50, Kt, sigmaPsib)
HTTNorm.BS.fun(time, psi, temp, thetaHT, Tb, To, Psib50, Kt, sigmaPsib)

Arguments

The 'HTTNorm.M()' and 'HTTNorm.BS()' functions have no arguments. The 'HTTNorm.M.fun()' and the 'HTTNorm.BS.fun()' functions have the following arguments:

time
time
Psi
water potential in the substrate
Temp
temperature
thetaHT
hydro-time parameter
Tb       base temperature
To       optimal temperature
Psib50   median base water potential
Kt       parameter measuring the effect of temperature on base water potential
sigmaPsib standard deviation for the base water potential within the seed lot

Details
The detail of this time-to-event model and the meaning of parameters are described in Bradford (2002).

Value
The 'HTTnorm.M.fun()' and 'HTTnorm.BS.fun()' functions return the proportion of germinated seeds, for any given values of time, water potential and temperature in the substrate. The 'HTTnorm.M()' and 'HTTnorm.BS()' functions return a list containing the nonlinear function, the self-starter function, the parameter names and other items which are internally used by the 'drmte()' function.

Note
This function is for use with the function 'drmte()'

Author(s)
Andrea Onofri

References

Examples
```
data(hordeum)
modHTTnorm.M <- drmte(nSeeds ~ timeBef + timeAf + water + temp, 
                        data=hordeum, 
                        fct = HTTnorm.M(), 
                        start=c(932, -2.5, -3, 0.07, 0.5))
summary(modHTTnorm.M)
```
Hydrotime model with Weibull Type I distribution of base water potential (Mesgaran et al., 2013)

Description

This model relates the time-course of the proportion of germinated seeds to the water potential in the substrate. It is based on a Weibull Type I distribution of base water potential within the seed lot. The equation is:

\[ P(t) = \exp \left\{ -\exp \left[ -\log \left( \Psi - \left( \frac{\theta_H \mu}{\tau} \right) + \delta \right) \right] - \log(\Psi_b + \delta) \sigma \right\} \]

In contrast to other hydrotime models (e.g., 'HTE1()', 'HTE2()', 'HTE3()', describing the distribution of germination time), this model describes the distribution of base water potential within the population, while the distribution of germination times is indirectly modelled, but it is not, in itself, gaussian (you see that 't' is at the denominator). The 'HTW1.fun()' is a generic function, which can be used for plotting or other applications, while the 'HTW1()' function is meant to be used for model fitting with the 'drmte()' function in the 'drcte()' package.

Usage

HTW1()

HTW1.fun(time, Psi, thetaH, delta, mu, sigma)

Arguments

The 'HTW1()' function has no arguments. The 'HTW1.fun()' function has the following arguments:

- **time**: time
- **Psi**: water potential in the substrate
- **thetaH**: hydro-time constant
- **delta**: shifting parameter to allow for negative water potential levels
- **mu**: location parameter for the Type I Weibull distribution
- **sigma**: scale parameter for the Type I Weibull Value distribution

Details

The detail of this time-to-event model and the meaning of parameters are described in Mesgaran et al. (2013).

Value

The 'HTW1.fun()' function returns the proportion of germinated seeds, for any given values of time and water potential in the substrate. The 'HTW1()' function returns a list containing the nonlinear function, the self-starter function, the parameter names and other items which are internally used by the 'drmte()' function.
Note

This function is for use with the function 'drmte()'

Author(s)

Andrea Onofri

References


Examples

data(rape)
modg <- drm( propCum ~ timeAf + Psi, fct=HTW1(), data=rape)
summary(modg)

Description

This model relates the time-course of the proportion of germinated seeds to the water potential in the substrate. It is based on a Weibull Type II distribution of base water potential within the seed lot. The equation is:

\[
P(t) = 1 - \exp\left\{ - \exp\left[ \log\left( \frac{\Psi - (\frac{\theta H}{\mu}) + \delta}{\sigma} \right) - \log(\Psi_b + \delta) \right] \right\}
\]

In contrast to other hydrotime models (e.g., 'HTE1()', 'HTE2()', 'HTE3()', describing the distribution of germination time), this model describes the distribution of base water potential within the population, while the distribution of germination times is indirectly modelled, but it is not, in itself, gaussian (you see that 't' is at the denominator). The 'HTW2.fun()' is a generic function, which can be used for plotting or other applications, while the 'HTW2()' function is meant to be used for model fitting with the 'drmte()' function in the 'drcte()' package.

Usage

HTW2()
HTW2.fun(time, Psi, thetaH, delta, mu, sigma)
Arguments

The 'HTW2()' function has no arguments. The 'HTW2.fun()' function has the following arguments:

- time: time
- Psi: water potential in the substrate
- thetaH: hydro-time constant
- delta: shifting parameter to allow for negative water potential levels
- mu: location parameter for the Type II Weibull distribution
- sigma: scale parameter for the Type II Weibull Value distribution

Details

The detail of this time-to-event model and the meaning of parameters are described in Mesgaran et al. (2013).

Value

The 'HTW2.fun()' function returns the proportion of germinated seeds, for any given values of time and water potential in the substrate. The 'HTW2()' function returns a list containing the nonlinear function, the self-starter function, the parameter names and other items which are internally used by the 'drmte()' function.

Note

This function is for use with the function 'drmte()'

Author(s)

Andrea Onofri

References


Examples

data(rape)
modg <- drm(propCum ~ timeAf + Psi, fct=HTW2(), data=rape)
summary(modg)
Delete-a-Petri-dish jackknife

Description

This function estimates standard errors for regression model parameters by using the fully-iterated delete-a-group jackknife (Yu and Peng, 2008). This is asymptotically equivalent to the cluster-robust 'sandwich' variance estimator (Lipsitz, 1994).

Usage

`jackGroupSE(obj, data, units)`

Arguments

- `obj`: a fitted model object
- `data`: dataset containing the variables used in `obj`
- `units`: a vector containing the code for Petri dishes

Details

Care should be taken to ensure that the assumptions for using a fully-iterated jackknife are valid.

Value

This function returns a data.frame containing the estimated parameters, the SEs and the robust SEs

Author(s)

Andrea Onofri

References


Examples

data(rape)

```r
modHTE <- drmte(nSeeds ~ timeBef + timeAf + Psi,
                data=rape, fct=HTE1())
robust <- jackGroupSE(modHTE, rape, rape$Dish)
robust
```
Reshape a seed germination datasets for time-to-event analyses (deprecated)

Description

MakeDrm: this function reshapes a common field book for germination assays into the form that is required for time-to-event analyses with drm() in the drc package and drmte() in the drcte package. The common field book has one row per each Petri dish and the counts of germinated seeds at each assessment time are listed in different columns.

Usage

makeDrm(counts, treat, nViable, moniTimes)

Arguments

counts Dataframe listing the counts of germinated seeds in each Petri dish (rows) at each assessment time (columns).
treat Dataframe listing for each dish the levels of each treatment. It has as many columns as there are treatments in the experiment.
nViable A vector with the number of viable seeds per dish, at the beginning of the assay
moniTimes Vector of monitoring times. It needs to be of the same length as the number of columns in 'counts'.

Value

This function returns a dataframe

Author(s)

Andrea Onofri

Reshape a seed germination datasets for time-to-event model fitting.

Description

This function reshapes a datasets organised as necessary for nonlinear regression into the kind of dataset required by the drmte() function in the 'drcte' package. It works with either the counts of germinated seeds at each monitoring time or the cumulative counts at each monitoring time.

Usage

makeDrm2(counts, treat, nViable, moniTimes, Dish, cumulative = TRUE)
Arguments

- **counts**: Vector listing the counts of germinated seeds in each Petri dish at each assessment time.
- **treat**: Dataframe listing, for each row of data, the corresponding level of experimental factors (one factor per column)
- **nViable**: A vector listing the number of viable seeds, at the beginning of the assay. This number is the same for all observations belonging to the same dish.
- **moniTimes**: Vector of monitoring times.
- **Dish**: Vector of codes for dishes.
- **cumulative**: Logical: True if counts are cumulative, False if they are not.

Value

Returns a dataframe

Author(s)

Andrea Onofri

Examples

```r
# makeDrm2 (deprecated)
data(lotusCum)
moniTime <- lotusCum$Time
count <- lotusCum$nCum
nViable <- rep(25, length(lotusCum[,1]))
Dish <- as.factor(lotusCum$Dish)
treatGroups <- lotusCum[,1]
dataset_sd <- makeDrm2(count, treatGroups, nViable, moniTime, Dish)
head(dataset_sd)
count <- lotusCum$nSeeds
dataset_sd <- makeDrm2(count, treatGroups, nViable, moniTime, Dish, cumulative = FALSE)
head(dataset_sd)
```

---

Data frame containing the counts of germinated seeds in each Petri dish at each assessment time.

**Description**

The germination of Phalaris minor was assayed at 6 mater potential levels and 6 temperature levels, on four replicated Petri dishes at each temperature and water potential. Twenty-five seeds per Petri dish were used. Inspections were made daily for twenty days.

**Usage**

data("phalaris")
Format

A data frame with 3024 observations on the following 9 variables.

- `temp` a numeric vector with the temperature level
- `water` a numeric vector with the water potential level
- `Dish` a numeric vector with the coding for Petri dishes
- `timeBef` a numeric vector, with the start time for each inspection interval
- `timeAf` a numeric vector, with the end time for each inspection interval
- `nViable` a numeric vector, with the number of viable seeds per dish, at the beginning of the assay
- `nSeeds` a numeric vector, with the number of germinated seeds during each inspection interval
- `nCum` a numeric vector, with the cumulative number of germinated seeds at each assessment time
- `propCum` a numeric vector, the cumulative proportion of germinated seeds at each assessment time

Details

The variable 'timeAf' contains the value 'Inf' (Infinity), that corresponds to the seeds which did not germinate during the assay, for which the germination times might be comprised from the last assessment time to infinity.

Author(s)

Mohsen Mesgaran

Source


References


Examples

data(phalaris)
Effect of environmental water potential or temperature on the germination capability of a seed lot

Description
These models are used to describe the germination capability of a seed lot, depending on the environmental water potential or temperature.

Usage
PmaxPsi1(fixed = c(NA, NA, NA), names = c("G", "Psib", "sigma"))
PmaxT1(fixed = c(NA, NA, NA), names = c("G", "Tc", "sigmaTc"))
PmaxPsi1.fun(Psi, G, Psib, sigma)
PmaxT1.fun(Temp, G, Tc, sigmaTc)

Arguments
- **fixed**: numeric vector. Specifies which parameters are fixed and at what value they are fixed. Defaults to NAs for parameter that are not fixed.
- **names**: a vector of character strings giving the names of the parameters (should not contain ":"). The default is reasonable.
- **Psi**: Water potential level
- **Temp**: Temperature level
- **G**: Maximum proportion of germinated seeds
- **Psib**: Base water potential level
- **sigma**: Variability of base water potential within the seed lot
- **Tc**: Ceiling temperature level
- **sigmaTc**: Variability of ceiling temperature level within the population

Details
The R functions 'PmaxPsi1()' and 'PmaxT1()' are meant to be used for model fitting with the 'drm()' function, within the 'drc' package. On the other hand, 'PmaxPsi1.fun()' and 'PmaxT1.fun()' are general purpose functions, to be used for plotting or other applications.

Value
'PmaxPsi1()' and 'PmaxT1()' return a list containing the nonlinear function, the self starter function and the parameter names, that are internally used for model fitting. 'PmaxPsi1.fun()' and 'PmaxT1.fun()' return the maximum proportion of germinated seeds for any given level of temperature or water potential in the substrate.

Author(s)
Andrea Onofri
References


Examples

library(drcte)
# Pmax vs Psi (shifted exponential)
Psi <- seq(-2.2, 0, by = 0.2)
Pmax <- c(0, 0, 0.076, 0.413, 0.514, 0.643, 0.712,
         0.832, 0.865, 0.849, 0.89, 0.90)
mod <- drm(Pmax ~ Psi, fct = PmaxPsi1())
summary(mod)

# Pmax vs Psi (shifted exponential, with asymptote)
Psi <- seq(-2.2, 0, by = 0.2)
Pmax <- c(0, 0, 0.076, 0.413, 0.514, 0.643, 0.712,
         0.832, 0.865, 0.849, 0.89, 0.90)
mod <- drm(Pmax ~ Psi, fct = PmaxPsi1(fixed = c(1, NA, NA)))
summary(mod)

# Pmax vs temperature
Tval <- c(0, 2.5, 5, 7.5, 10, 12.5, 15, 17.5,
         20, 22.5, 25, 27.5, 30, 32.5, 35)
Pmax2 <- c(0.79, 0.81, 0.807, 0.776, 0.83,
         0.73, 0.744, 0.73, 0.828, 0.818,
         0.805, 0.706, 0.41, 0.0002, 0)
mod2 <- drm(Pmax2 ~ Tval, fct = PmaxT1())
summary(mod2)

rape

Germination data from an assay of rapeseed at decreasing water potential levels

Description

This file describes the germination of rapeseed (cv. Excalibur) at different water potential levels in the substrate.

Usage

data("rape")

Format

A data frame with 294 observations on the following 7 variables.

Psi a numeric vector: waterpotential level
Dish a numeric vector: code for Petri dishes
timeBef a numeric vector: beginning of scoring interval
timeAf  a numeric vector: end of scoring interval  
nSeeds  a numeric vector: number of germinated seeds, between timeBef and timeAf  
nCum  a numeric vector: cumulative number of germinated seeds at timeAf  
propCum  a numeric vector: cumulative proportion of germinated seeds at timeAf

**Source**


**References**


**Examples**

```r
#Fitting a hydrotime model
library(drcte)
data(rape)
gmod <- drmte(nSeeds ~ timeBef + timeAf + Psi,  
               data = rape, fct = HTE1())
summary(gmod)
```

**Description**

This file describes the germination of rapeseed (cv. Excalibur and Toccata) at different water potential levels in the substrate.

**Usage**

```r
data("rape")
```

**Format**

A data frame with 294 observations on the following 8 variables.

- **CV**: a factor: rape genotype
- **Psi**: a numeric vector: water potential level
- **Dish**: a numeric vector: code for Petri dishes
- **timeBef**: a numeric vector: beginning of scoring interval
- **timeAf**: a numeric vector: end of scoring interval
- **nSeeds**: a numeric vector: number of germinated seeds, between timeBef and timeAf
- **nCum**: a numeric vector: cumulative number of germinated seeds at timeAf
- **propCum**: a numeric vector: cumulative proportion of germinated seeds at timeAf
Thermal-Time\_models

- \texttt{timeAf} a numeric vector: end of scoring interval
- \texttt{nSeeds} a numeric vector: number of germinated seeds, between \texttt{timeBef} and \texttt{timeAf}
- \texttt{nCum} a numeric vector: cumulative number of germinated seeds at \texttt{timeAf}
- \texttt{Prop} a numeric vector: cumulative proportion of germinated seeds at \texttt{timeAf}

\textbf{Source}


\textbf{References}


\textbf{Examples}

\begin{verbatim}
#Fitting a hydrotime-to-event model
data(rape2G)
modHTE <- drmte( nSeeds ~ timeBef + timeAf + Psi,
data=rape2G, fct=HTE1(),
curveid = CV)
summary(modHTE)
\end{verbatim}

\underline{Thermal-Time\_models}  \textit{Thermal-time models with log-logistic distribution of germination time (Onofri et al., 2018)}

\textbf{Description}

These models relates the time-course of germinations to the environmental temperature and they are based on a truncated log-logistic distribution of germination time:

\[
P(t) = \frac{d}{1 + \exp \left[ b(\log(x) - \log(e)) \right]}
\]

where the usual parameters ('b', 'd' and 'e') are expressed as functions of \textit{temperature} \((T)\). In the function 'TTEM()', we implemented the following submodels: (1) for the parameter 'd', we implemented the following submodels:

\[
d = G \left[ 1 - \exp \left( - \frac{T_c - T}{\sigma T_c} \right) \right]
\]

\[
1/[e(T)] = GR_{50}(T) = \frac{T - T_b}{\theta T_c - T_b} \left[ 1 - \frac{T_c - T_b}{T_c - T_b} \right]
\]
while 'b' was regarded as constant and independent from temperature.

In the 'TTERF()' function, the last submodel was modified, according to Rowse and Finch-Savage (2003):

\[
1/[e(T)] = GR_{50}(T) = \begin{cases} 
\frac{T - T_b}{\sigma_T} & \text{if } T_b < T < T_d \\
\frac{T - T_b}{\sigma_T} \left[1 - \frac{T - T_b}{T_c - T_d}\right] & \text{if } T_d < T < T_c \\
0 & \text{if } T \leq T_b \text{ or } T \geq T_c
\end{cases}
\]

In the In the 'TTERFc()' function a further submodel was introduced, to model the effect of temperature in the shape parameter 'b':

\[
\sigma(T) = \frac{1}{b} = \frac{1}{b_0} + s(T - T_b)
\]

The 'TTEM.fun()', 'TTERF.fun()' and 'TTERFc.fun()' are generic functions, which can be used for plotting or other applications, while the 'TTEM()', 'TTERF()' and 'TTERFc()' functions are meant to be used for model fitting with the 'drmte()' function in the 'drcte()' package.

### Usage

TTEM()
TTEM.fun(time, Temp, G, Tc, sigmaTc, Tb, ThetaT, b)
TTERF()
TTERF.fun(time, Temp, G, Tc, sigmaTc, Td, Tb, ThetaT, b)
TTERFc()
TTERFc.fun(time, Temp, G, Tc, sigmaTc, Td, Tb, ThetaT, b0, s)

### Arguments

The TTEM(), TTERF() and TTERFc() functions have no arguments. The TTEM.fun(), TTERF.fun() and TTERFc.fun() functions have the following arguments:

- **time**: time
- **Temp**: temperature
- **G**: maximum germination capability of the seed lot
- **Tc**: ceiling temperature
- **sigmaTc**: standard deviation for the ceiling temperature within the seed lot
- **Tb**: base temperature
- **Td**: close to optimal temperature
- **ThetaT**: Thermal-time parameter
- **b**: shape parameter for the cumulative distribution function of germination time
- **b0**: shape parameter for the cumulative distribution function of germination time at \( T = T_b \)
- **s**: parameter for the effect of temperature on the scale parameter for the log-logistic distribution of germination times
Details

The detail of these functions are described in Onofri et al. (2018).

Value

The 'TTEm.fun()', 'TTERF.fun()' and 'TTERFc.fun()' functions return the proportion of germinated seeds, for given values of time and temperature. The 'TTEm()', 'TTERF()' and 'TTERFc()' functions return a list containing the nonlinear function, the self starter function, the parameter names and other items which are internally used by the 'drmte()' function.

Author(s)

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References


Examples

data(barley)
modTTE <- drmte(nSeeds ~ timeBef + timeAf + Temp, data = barley, 
  fct = TTERF())
summary(modTTE)
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