Package 'fishgrowth'

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Contents	
fishgrowth-packagegcmgompertz	

2 fishgrowth-package

		_
Index		47
	vonberto	 42
	tags_skj	 36
	schnute3	 31
	richardso	 26
	richards	 21
	pred_band	 20
	otoliths_skj	 19
	gompertzo	 13

Description

Fit growth models to otoliths and/or tagging data, using the RTMB package and maximum likelihood.

The otoliths (or similar measurements of age) provide direct observed coordinates of age and length. The tagging data provide information about the observed length at release and length at recapture at a later time, where the age at release is unknown and estimated as a vector of parameters.

The growth models provided by this package can be fitted to otoliths only, tagging data only, or a combination of the two. Growth variability can be modelled as constant or increasing with length.

Details

Growth models:

Utilities:

Example data:

```
growth cessation
gcm
             Gompertz
gompertz
gompertzo
             Gompertz (old style)
             Richards
richards
richardso
             Richards (old style)
             Schnute Case 3
schnute3
             von Bertalanffy
vonbert
vonberto
             von Bertalanffy (old style)
                  prediction band
     pred_band
                   otoliths (haddock)
  otoliths_had
                   otoliths (skipjack)
  otoliths_skj
```

```
tags_skj tags (skipjack)
```

Author(s)

Arni Magnusson and Mark Maunder.

See Also

Useful links:

• https://github.com/arni-magnusson/fishgrowth

gcm

Growth Cessation Model

Description

Fit a growth cessation model (GCM) to otoliths and/or tags.

Usage

```
gcm(par, data, silent = TRUE, ...)
gcm_curve(t, L0, rmax, k, t50)
gcm_objfun(par, data)
```

Arguments

par	a parameter list.
data	a data list.
silent	passed to MakeADFun.
	passed to MakeADFun.
t	age (vector).
L0	predicted length at age 0.
rmax	shape parameter that determines the initial slope.
k	shape parameter that determines how quickly the growth curve reaches the asymptotic maximum.
t50	shape parameter that determines the logistic function midpoint.

Details

The main function gcm creates a model object, ready for parameter estimation. The auxiliary functions gcm_curve and gcm_objfun are called by the main function to calculate the regression curve and objective function value. The user can also call the auxiliary functions directly for plotting and model exploration.

The par list contains the following elements:

- L0, predicted length at age 0
- log_rmax, shape parameter that determines the initial slope
- log_k, shape parameter that determines how quickly the growth curve reaches the asymptotic
 maximum
- t50, shape parameter that determines the logistic function midpoint
- log_sigma_min, growth variability at the shortest observed length in the data
- log_sigma_max (*), growth variability at the longest observed length in the data
- log_age (*), age at release of tagged individuals (vector)
- *: The parameter log_sigma_max can be omitted to estimate growth variability that does not vary with length. The parameter vector log_age can be omitted to fit to otoliths only.

The data list contains the following elements:

- Aoto (*), age from otoliths (vector)
- Loto (*), length from otoliths (vector)
- Lrel (*), length at release of tagged individuals (vector)
- Lrec (*), length at recapture of tagged individuals (vector)
- liberty (*), time at liberty of tagged individuals in years (vector)

Value

The gcm function returns a TMB model object, produced by MakeADFun.

The gcm_curve function returns a numeric vector of predicted length at age.

The gcm_objfun function returns the negative log-likelihood as a single number, describing the goodness of fit of par to the data.

Note

The growth cessation model (Maunder et al. 2018) predicts length at age as:

$$\hat{L}_{t} = r_{\text{max}} \left[\frac{\log (1 + e^{-kt_{50}}) - \log (1 + e^{k(t - t_{50})})}{k} + t \right]$$

The variability of length at age increases linearly with length,

^{*:} The data vectors Aoto and Loto can be omitted to fit to tagging data only. The data vectors Lrel, Lrec, and liberty can be omitted to fit to otoliths only.

$$\sigma_L = \alpha + \beta \hat{L}$$

where the slope is $\beta = (\sigma_{\rm max} - \sigma_{\rm min})/(L_{\rm max} - L_{\rm min})$, the intercept is $\alpha = \sigma_{\rm min} - \beta L_{\rm min}$, and $L_{\rm min}$ and $L_{\rm max}$ are the shortest and longest observed lengths in the data. Alternatively, growth variability can be modelled as a constant $\sigma_L = \sigma_{\rm min}$ that does not vary with length, by omitting log_sigma_max from the parameter list (see above).

The negative log-likelihood is calculated by comparing the observed and predicted lengths:

```
nll_Loto <- -dnorm(Loto, Loto_hat, sigma_Loto, TRUE)
nll_Lrel <- -dnorm(Lrel, Lrel_hat, sigma_Lrel, TRUE)
nll_Lrec <- -dnorm(Lrec, Lrec_hat, sigma_Lrec, TRUE)
nll <- sum(nll_Loto) + sum(nll_Lrel) + sum(nll_Lrec)</pre>
```

References

Maunder, M.N., Deriso, R.B., Schaefer, K.M., Fuller, D.W., Aires-da-Silva, A.M., Minte-Vera, C.V., and Campana, S.E. (2018). The growth cessation model: a growth model for species showing a near cessation in growth with application to bigeye tuna (*Thunnus obesus*). *Marine Biology*, **165**, 76. doi:10.1007/s0022701833369.

See Also

gcm, gompertz, gompertzo, richards, richardso, schnute3, vonbert, and vonberto are alternative growth models.

pred_band calculates a prediction band for a fitted growth model.

otoliths_had, otoliths_skj, and tags_skj are example datasets.

fishgrowth-package gives an overview of the package.

Examples

```
# Model 1: Fit to haddock otoliths
# Explore initial parameter values
plot(len~age, otoliths_had, xlim=c(0,18), ylim=c(0,105), pch=16,
     col="#0080a010")
x \leftarrow seq(1, 18, 0.1)
lines(x, gcm_curve(x, L0=5, rmax=20, k=0.15, t50=0), lty=3)
# Prepare parameters and data
init <- list(L0=5, log_rmax=log(20), log_k=log(0.15), t50=-1,</pre>
             log_sigma_min=log(3), log_sigma_max=log(3))
dat <- list(Aoto=otoliths_had$age, Loto=otoliths_had$len)</pre>
gcm_objfun(init, dat)
# Fit model
model <- gcm(init, dat)</pre>
fit <- nlminb(model$par, model$fn, model$gr,</pre>
               control=list(eval.max=1e4, iter.max=1e4))
report <- model$report()</pre>
```

```
sdreport <- sdreport(model)</pre>
# Plot results
Lhat <- with(report, gcm_curve(x, L0, rmax, k, t50))
lines(x, Lhat, lwd=2, col=2)
legend("bottomright", c("initial curve", "model fit"), col=c(1,2), lty=c(3,1),
      lwd=c(1,2), bty="n", inset=0.02, y.intersp=1.25)
# Model summary
report[c("L0", "rmax", "k", "t50", "sigma_min", "sigma_max")]
fit[-1]
summary(sdreport)
# Plot 95% prediction band
band <- pred_band(x, model)</pre>
areaplot::confplot(cbind(lower,upper)~age, band, xlim=c(0,18), ylim=c(0,100),
        ylab="len", col="mistyrose")
points(len~age, otoliths_had, xlim=c(0,18), ylim=c(0,100),
      pch=16, col="#0080a010")
lines(x, Lhat, lwd=2, col=2)
lines(lower~age, band, lty=1, lwd=0.5, col=2)
lines(upper~age, band, lty=1, lwd=0.5, col=2)
# Model 2: Fit to skipjack otoliths and tags
# Explore initial parameter values
plot(len~age, otoliths_skj, xlim=c(0,4), ylim=c(0,100))
x < - seq(0, 4, 0.1)
points(lenRel~I(lenRel/60), tags_skj, col=4)
points(lenRec~I(lenRel/60+liberty), tags_skj, col=3)
lines(x, gcm_curve(x, L0=20, rmax=120, k=2, t50=0), lty=2)
legend("bottomright", c("otoliths","tag releases","tac recaptures",
       "initial curve"), col=c(1,4,3,1), pch=c(1,1,1,NA), lty=c(0,0,0,2),
      lwd=c(1.2,1.2,1.2,1), bty="n", inset=0.02, y.intersp=1.25)
# Prepare parameters and data
init <- list(L0=20, log_rmax=log(120), log_k=log(4), t50=0,</pre>
            log_sigma_min=log(3), log_sigma_max=log(3),
            log_age=log(tags_skj$lenRel/60))
dat <- list(Aoto=otoliths_skj$age, Loto=otoliths_skj$len,</pre>
           Lrel=tags_skj$lenRel, Lrec=tags_skj$lenRec,
           liberty=tags_skj$liberty)
gcm_objfun(init, dat)
# Fit model
model <- gcm(init, dat)</pre>
fit <- nlminb(model$par, model$fn, model$gr,</pre>
             control=list(eval.max=1e4, iter.max=1e4))
report <- model$report()</pre>
sdreport <- sdreport(model)</pre>
```

```
# Plot results
plot(len~age, otoliths_skj, xlim=c(0,4), ylim=c(0,100))
points(report$age, report$Lrel, col=4)
points(report$age+report$liberty, report$Lrec, col=3)
Lhat <- with(report, gcm_curve(x, L0, rmax, k, t50))
lines(x, Lhat, lwd=2)
legend("bottomright", c("otoliths","tag releases","tac recaptures",
       "model fit"), col=c(1,4,3,1), pch=c(1,1,1,NA), lty=c(0,0,0,1),
       lwd=c(1.2,1.2,1.2,2), bty="n", inset=0.02, y.intersp=1.25)
# Model summary
report[c("L0", "rmax", "k", "t50", "sigma_min", "sigma_max")]
fit[-1]
head(summary(sdreport), 6)
# Model 3: Stepwise estimation procedure, described by Maunder et al. (2018)
# - estimate L0 and rmax using linear regression on younger fish
# - estimate k and t0 using GCM and all data, keeping L0 and rmax fixed
# Estimate L0 and rmax
plot(otoliths\_skj, \ xlim=c(0,4), \ ylim=c(0,100))
fm \leftarrow lm(len~age, otoliths_skj)
abline(fm)
L0 \leftarrow coef(fm)[[1]]
rmax <- coef(fm)[[2]]</pre>
# Explore initial parameter values (k, t50, age)
t <- seq(0, 4, by=0.1)
points(t, gcm_curve(t, L0, rmax, k=3, t50=2), col="gray")
points(lenRel~I(lenRel/50), tags_skj, col=4)
points(lenRec~I(lenRel/50+liberty), tags_skj, col=3)
legend("bottomright", c("otoliths","tag releases","tac recaptures",
       "linear regression (otoliths)"), col=c(1,4,3,1), pch=c(1,1,1,NA),
       lty=c(0,0,0,1), lwd=c(1.2,1.2,1.2,2), bty="n", inset=0.02,
       y.intersp=1.25)
# Prepare parameters
init <- list(L0=L0, log_rmax = log(rmax), log_k = log(3), t50=2,
             log_sigma_min=log(3), log_sigma_max=log(3),
             log_age=log(tags_skj$lenRel/50))
# Fit model
map <- list(L0=factor(NA), log_rmax=factor(NA)) # fix L0 and rmax</pre>
model <- gcm(init, dat, map=map)</pre>
fit <- nlminb(model$par, model$fn, model$gr,</pre>
              control=list(eval.max=1e4,iter.max=1e4))
report <- model$report()</pre>
sdreport <- sdreport(model)</pre>
# Plot results
plot(len~age, otoliths_skj, xlim=c(0,4), ylim=c(0,100))
```

gompertz

Gompertz Growth Model

Description

Fit a Gompertz growth model to otoliths and/or tags, using the Schnute parametrization.

Usage

```
gompertz(par, data, silent = TRUE, ...)
gompertz_curve(t, L1, L2, k, t1, t2)
gompertz_objfun(par, data)
```

Arguments

par	a parameter list.
data	a data list.
silent	passed to MakeADFun.
	passed to MakeADFun.
t	age (vector).
L1	predicted length at age t1.
L2	predicted length at age t2.
k	growth coefficient.
t1	age where predicted length is L1.
t2	age where predicted length is L2

Details

The main function gompertz creates a model object, ready for parameter estimation. The auxiliary functions gompertz_curve and gompertz_objfun are called by the main function to calculate the regression curve and objective function value. The user can also call the auxiliary functions directly for plotting and model exploration.

The par list contains the following elements:

- log_L1, predicted length at age t1
- log_L2, predicted length at age t2
- log_k, growth coefficient
- log_sigma_min, growth variability at the shortest observed length in the data
- log_sigma_max (*), growth variability at the longest observed length in the data
- log_age (*), age at release of tagged individuals (vector)
- *: The parameter log_sigma_max can be omitted to estimate growth variability that does not vary with length. The parameter vector log_age can be omitted to fit to otoliths only.

The data list contains the following elements:

- Aoto (*), age from otoliths (vector)
- Loto (*), length from otoliths (vector)
- Lrel (*), length at release of tagged individuals (vector)
- Lrec (*), length at recapture of tagged individuals (vector)
- liberty (*), time at liberty of tagged individuals in years (vector)
- t1, age where predicted length is L1
- t2, age where predicted length is L2

Value

The gompertz function returns a TMB model object, produced by MakeADFun.

The gompertz_curve function returns a numeric vector of predicted length at age.

The gompertz_objfun function returns the negative log-likelihood as a single number, describing the goodness of fit of par to the data.

Note

The Schnute parametrization used in gompertz reduces parameter correlation and improves convergence reliability compared to the traditional parametrization used in gompertzo. Therefore, the gompertz parametrization can be recommended for general usage, as both parametrizations produce the same growth curve. However, there can be some use cases where the traditional parametrization (Linf, k, tau) is preferred over the Schnute parametrization (L1, L2, k).

^{*:} The data vectors Aoto and Loto can be omitted to fit to tagging data only. The data vectors Lrel, Lrec, and liberty can be omitted to fit to otoliths only.

Gompertz is a special case of the Richards (1959) model, where b = 0. If the best model fit of a richards model to a particular dataset involves a very small estimated value of b, then the gompertz model offers a preferable parametrization, as it produces the same curve using fewer parameters.

The Gompertz (1825) growth model, as parametrized by Schnute (1981, Eq. 16) predicts length at age as:

$$L = L_1 \exp \left[\log(L_2/L_1) \frac{1 - e^{-k(t-t_1)}}{1 - e^{-k(t_2-t_1)}} \right]$$

The variability of length at age increases linearly with length,

$$\sigma_L = \alpha + \beta \hat{L}$$

where the slope is $\beta = (\sigma_{\rm max} - \sigma_{\rm min})/(L_{\rm max} - L_{\rm min})$, the intercept is $\alpha = \sigma_{\rm min} - \beta L_{\rm min}$, and $L_{\rm min}$ and $L_{\rm max}$ are the shortest and longest observed lengths in the data. Alternatively, growth variability can be modelled as a constant $\sigma_L = \sigma_{\rm min}$ that does not vary with length, by omitting log_sigma_max from the parameter list (see above).

The negative log-likelihood is calculated by comparing the observed and predicted lengths:

```
nll_Loto <- -dnorm(Loto, Loto_hat, sigma_Loto, TRUE)
nll_Lrel <- -dnorm(Lrel, Lrel_hat, sigma_Lrel, TRUE)
nll_Lrec <- -dnorm(Lrec, Lrec_hat, sigma_Lrec, TRUE)
nll <- sum(nll_Loto) + sum(nll_Lrel) + sum(nll_Lrec)</pre>
```

References

Gompertz, B. (1825). On the nature of the function expressive of the law of human mortality, and on a new mode of determining the value of life contingencies. *Philosophical Transactions of the Royal Society*, **115**, 513-583.

Schnute, J. (1981). A versatile growth model with statistically stable parameters. *Canadian Journal of Fisheries and Aquatic Science*, **38**, 1128-1140. doi:10.1139/f81153.

See Also

gcm, gompertz, gompertzo, richards, richards, schnute3, vonbert, and vonberto are alternative growth models.

pred_band calculates a prediction band for a fitted growth model.

otoliths_had, otoliths_skj, and tags_skj are example datasets.

fishgrowth-package gives an overview of the package.

Examples

```
lines(x, gompertz_curve(x, L1=15, L2=70, k=0.4, t1=1, t2=10), lty=3)
# Prepare parameters and data
init <- list(log_L1=log(20), log_L2=log(70), log_k=log(0.1),\\
            log_sigma_min=log(3), log_sigma_max=log(3))
dat <- list(Aoto=otoliths_had$age, Loto=otoliths_had$len, t1=1, t2=10)</pre>
gompertz_objfun(init, dat)
# Fit model
model <- gompertz(init, dat)</pre>
fit <- nlminb(model$par, model$fn, model$gr,</pre>
             control=list(eval.max=1e4, iter.max=1e4))
report <- model$report()</pre>
sdreport <- sdreport(model)</pre>
# Plot results
Lhat <- with(report, gompertz_curve(x, L1, L2, k, t1, t2))
lines(x, Lhat, lwd=2, col=2)
legend("bottomright", c("initial curve", "model fit"), col=c(1,2), lty=c(3,1),
      lwd=c(1,2), bty="n", inset=0.02, y.intersp=1.25)
# Model summary
report[c("L1", "L2", "k", "sigma_min", "sigma_max")]
fit[-1]
summary(sdreport)
# Plot 95% prediction band
band <- pred_band(x, model)</pre>
areaplot::confplot(cbind(lower,upper)~age, band, xlim=c(0,18), ylim=c(0,100),
        ylab="len", col="mistyrose")
points(len~age, otoliths_had, xlim=c(0,18), ylim=c(0,100),
      pch=16, col="#0080a010")
lines(x, Lhat, lwd=2, col=2)
lines(lower~age, band, lty=1, lwd=0.5, col=2)
lines(upper~age, band, lty=1, lwd=0.5, col=2)
# Model 2: Fit to skipjack otoliths and tags
# Explore initial parameter values
plot(len~age, otoliths_skj, xlim=c(0,4), ylim=c(0,100))
x < - seq(0, 4, 0.1)
points(lenRel~I(lenRel/60), tags_skj, col=4)
points(lenRec~I(lenRel/60+liberty), tags_skj, col=3)
lines(x, gompertz_curve(x, L1=28, L2=74, k=1, t1=0, t2=4), lty=2)
legend("bottomright", c("otoliths","tag releases","tac recaptures",
       "initial curve"), col=c(1,4,3,1), pch=c(1,1,1,NA), lty=c(0,0,0,2),
      lwd=c(1.2,1.2,1.2,1), bty="n", inset=0.02, y.intersp=1.25)
# Prepare parameters and data
init <- list(log_L1=log(28), log_L2=log(74), log_k=log(1),</pre>
            log_sigma_min=log(3), log_sigma_max=log(3),
```

```
log_age=log(tags_skj$lenRel/60))
dat <- list(Aoto=otoliths_skj$age, Loto=otoliths_skj$len,</pre>
           Lrel=tags_skj$lenRel, Lrec=tags_skj$lenRec,
           liberty=tags_skj$liberty, t1=0, t2=4)
gompertz_objfun(init, dat)
# Fit model
model <- gompertz(init, dat)</pre>
fit <- nlminb(model$par, model$fn, model$gr,</pre>
             control=list(eval.max=1e4, iter.max=1e4))
report <- model$report()</pre>
sdreport <- sdreport(model)</pre>
# Plot results
plot(len~age, otoliths_skj, xlim=c(0,4), ylim=c(0,100))
points(report$age, report$Lrel, col=4)
points(report$age+report$liberty, report$Lrec, col=3)
Lhat <- with(report, gompertz_curve(x, L1, L2, k, t1, t2))
lines(x, Lhat, lwd=2)
legend("bottomright", c("otoliths","tag releases","tac recaptures",
       "model fit"), col=c(1,4,3,1), pch=c(1,1,1,NA), lty=c(0,0,0,1),
      lwd=c(1.2,1.2,1.2,2), bty="n", inset=0.02, y.intersp=1.25)
# Model summary
report[c("L1", "L2", "k", "sigma_min", "sigma_max")]
fit[-1]
head(summary(sdreport), 5)
# Model 3: Fit to skipjack otoliths only
init <- list(log_L1=log(28), log_L2=log(74), log_k=log(1),</pre>
            log_sigma_min=log(3), log_sigma_max=log(3))
dat <- list(Aoto=otoliths_skj$age, Loto=otoliths_skj$len, t1=0, t2=4)</pre>
model <- gompertz(init, dat)</pre>
fit <- nlminb(model$par, model$fn, model$gr,</pre>
             control=list(eval.max=1e4, iter.max=1e4))
model$report()[c("L1", "L2", "k", "sigma_min", "sigma_max")]
# Model 4: Fit to skipjack otoliths only,
# but now estimating constant sigma instead of sigma varying by length
# We do this by omitting log_sigma_max
init <- list(log_L1=log(28), log_L2=log(74), log_k=log(1),</pre>
            log_sigma_min=log(3))
dat <- list(Aoto=otoliths_skj$age, Loto=otoliths_skj$len, t1=0, t2=4)</pre>
model <- gompertz(init, dat)</pre>
fit <- nlminb(model$par, model$fn, model$gr,</pre>
                   control=list(eval.max=1e4, iter.max=1e4))
model$report()[c("L1", "L2", "k", "sigma_min")]
```

gompertzo

Gompertz Growth Model (Old Style)

Description

Fit a Gompertz growth model to otoliths and/or tags, using a traditional parametrization.

Usage

```
gompertzo(par, data, silent = TRUE, ...)
gompertzo_curve(t, Linf, k, tau)
gompertzo_objfun(par, data)
```

Arguments

par	a parameter list.
data	a data list.
silent	passed to MakeADFun.
	passed to MakeADFun.
t	age (vector).
Linf	asymptotic maximum length.
k	growth coefficient.
tau	location parameter.

Details

The main function gompertzo creates a model object, ready for parameter estimation. The auxiliary functions gompertzo_curve and gompertzo_objfun are called by the main function to calculate the regression curve and objective function value. The user can also call the auxiliary functions directly for plotting and model exploration.

The par list contains the following elements:

- log_Linf, asymptotic maximum length
- log_k, growth coefficient
- tau, location parameter
- log_sigma_min, growth variability at the shortest observed length in the data
- log_sigma_max (*), growth variability at the longest observed length in the data
- log_age (*), age at release of tagged individuals (vector)
- *: The parameter log_sigma_max can be omitted to estimate growth variability that does not vary with length. The parameter vector log_age can be omitted to fit to otoliths only.

The data list contains the following elements:

- Aoto (*), age from otoliths (vector)
- Loto (*), length from otoliths (vector)
- Lrel (*), length at release of tagged individuals (vector)
- Lrec (*), length at recapture of tagged individuals (vector)
- liberty (*), time at liberty of tagged individuals in years (vector)

Value

The gompertzo function returns a TMB model object, produced by MakeADFun.

The gompertzo_curve function returns a numeric vector of predicted length at age.

The gompertzo_objfun function returns the negative log-likelihood as a single number, describing the goodness of fit of par to the data.

Note

The Schnute parametrization used in gompertz reduces parameter correlation and improves convergence reliability compared to the traditional parametrization used in gompertzo. Therefore, the gompertz parametrization can be recommended for general usage, as both parametrizations produce the same growth curve. However, there can be some use cases where the traditional parametrization (Linf, k, tau) is preferred over the Schnute parametrization (L1, L2, k).

Gompertz is a special case of the Richards (1959) model, where b=0. If the best model fit of a richards model to a particular dataset involves a very small estimated value of b, then the gompertz model offers a preferable parametrization, as it produces the same curve using fewer parameters.

^{*:} The data vectors Aoto and Loto can be omitted to fit to tagging data only. The data vectors Lrel, Lrec, and liberty can be omitted to fit to otoliths only.

The Gompertz (1825) growth model, as parametrized by Ricker (1979, Eq. 23) predicts length at age as:

$$L = L_{\infty} \exp(-e^{-k(t-\tau)})$$

The variability of length at age increases linearly with length,

$$\sigma_L = \alpha + \beta \hat{L}$$

where the slope is $\beta = (\sigma_{\rm max} - \sigma_{\rm min})/(L_{\rm max} - L_{\rm min})$, the intercept is $\alpha = \sigma_{\rm min} - \beta L_{\rm min}$, and $L_{\rm min}$ and $L_{\rm max}$ are the shortest and longest observed lengths in the data. Alternatively, growth variability can be modelled as a constant $\sigma_L = \sigma_{\rm min}$ that does not vary with length, by omitting log_sigma_max from the parameter list (see above).

The negative log-likelihood is calculated by comparing the observed and predicted lengths:

```
nll_Loto <- -dnorm(Loto, Loto_hat, sigma_Loto, TRUE)
nll_Lrel <- -dnorm(Lrel, Lrel_hat, sigma_Lrel, TRUE)
nll_Lrec <- -dnorm(Lrec, Lrec_hat, sigma_Lrec, TRUE)
nll <- sum(nll_Loto) + sum(nll_Lrel) + sum(nll_Lrec)</pre>
```

References

Gompertz, B. (1825). On the nature of the function expressive of the law of human mortality, and on a new mode of determining the value of life contingencies. *Philosophical Transactions of the Royal Society*, **115**, 513-583.

Ricker, W.E. (1979). Growth rates and models. In: W.S. Hoar et al. (eds.) *Fish physiology 8: Bioenergetics and growth*. New York: Academic Press, pp. 677-743.

See Also

gcm, gompertz, gompertzo, richards, richards, schnute3, vonbert, and vonberto are alternative growth models.

pred_band calculates a prediction band for a fitted growth model.

otoliths_had, otoliths_skj, and tags_skj are example datasets.

fishgrowth-package gives an overview of the package.

Examples

```
log_sigma_min=log(3), log_sigma_max=log(3))
dat <- list(Aoto=otoliths_had$age, Loto=otoliths_had$len)</pre>
gompertzo_objfun(init, dat)
# Fit model
model <- gompertzo(init, dat)</pre>
fit <- nlminb(model$par, model$fn, model$gr,</pre>
             control=list(eval.max=1e4, iter.max=1e4))
report <- model$report()</pre>
sdreport <- sdreport(model)</pre>
# Plot results
Lhat <- with(report, gompertzo_curve(x, Linf, k, tau))</pre>
lines(x, Lhat, lwd=2, col=2)
legend("bottomright", c("initial curve","model fit"), col=c(1,2), lty=c(3,1),
      lwd=c(1,2), bty="n", inset=0.02, y.intersp=1.25)
# Model summary
report[c("Linf", "k", "tau", "sigma_min", "sigma_max")]
fit[-1]
summary(sdreport)
# Plot 95% prediction band
band <- pred_band(x, model)</pre>
areaplot::confplot(cbind(lower,upper)~age, band, xlim=c(0,18), ylim=c(0,100),
        ylab="len", col="mistyrose")
points(len~age, otoliths_had, xlim=c(0,18), ylim=c(0,100),
      pch=16, col="#0080a010")
lines(x, Lhat, lwd=2, col=2)
lines(lower~age, band, lty=1, lwd=0.5, col=2)
lines(upper~age, band, lty=1, lwd=0.5, col=2)
# Model 2: Fit to skipjack otoliths and tags
# Explore initial parameter values
plot(len~age, otoliths_skj, xlim=c(0,4), ylim=c(0,100))
x < - seq(0, 4, 0.1)
points(lenRel~I(lenRel/60), tags_skj, col=4)
points(lenRec~I(lenRel/60+liberty), tags_skj, col=3)
lines(x, gompertzo_curve(x, Linf=75, k=1, tau=0), lty=2)
legend("bottomright", c("otoliths","tag releases","tac recaptures",
       "initial curve"), col=c(1,4,3,1), pch=c(1,1,1,NA), lty=c(0,0,0,2),
      lwd=c(1.2,1.2,1.2,1), bty="n", inset=0.02, y.intersp=1.25)
# Prepare parameters and data
init <- list(log_Linf=log(75), log_k=log(1), tau=0,</pre>
            log_sigma_min=log(3), log_sigma_max=log(3),
            log_age=log(tags_skj$lenRel/60))
dat <- list(Aoto=otoliths_skj$age, Loto=otoliths_skj$len,</pre>
           Lrel=tags_skj$lenRel, Lrec=tags_skj$lenRec,
           liberty=tags_skj$liberty)
```

```
gompertzo_objfun(init, dat)
# Fit model
model <- gompertzo(init, dat)</pre>
fit <- nlminb(model$par, model$fn, model$gr,</pre>
            control=list(eval.max=1e4, iter.max=1e4))
report <- model$report()</pre>
sdreport <- sdreport(model)</pre>
# Plot results
plot(len~age, otoliths_skj, xlim=c(0,4), ylim=c(0,100))
points(report$age, report$Lrel, col=4)
points(report$age+report$liberty, report$Lrec, col=3)
Lhat <- with(report, gompertzo_curve(x, Linf, k, tau))</pre>
lines(x, Lhat, lwd=2)
legend("bottomright", c("otoliths","tag releases","tac recaptures",
      "model fit"), col=c(1,4,3,1), pch=c(1,1,1,NA), lty=c(0,0,0,1),
      lwd=c(1.2,1.2,1.2,2), bty="n", inset=0.02, y.intersp=1.25)
# Model summary
report[c("Linf", "k", "tau", "sigma_min", "sigma_max")]
fit[-1]
head(summary(sdreport), 5)
# Model 3: Fit to skipjack otoliths only
init <- list(log_Linf=log(75), log_k=log(1), tau=0,</pre>
           log_sigma_min=log(3), log_sigma_max=log(3))
dat <- list(Aoto=otoliths_skj$age, Loto=otoliths_skj$len)</pre>
model <- gompertzo(init, dat)</pre>
fit <- nlminb(model$par, model$fn, model$gr,</pre>
                control=list(eval.max=1e4, iter.max=1e4))
model$report()[c("Linf", "k", "tau", "sigma_min", "sigma_max")]
# Model 4: Fit to skipjack otoliths only,
# but now estimating constant sigma instead of sigma varying by length
# We do this by omitting log_sigma_max
init <- list(log_Linf=log(75), log_k=log(1), tau=0,</pre>
           log_sigma_min=log(3))
dat <- list(Aoto=otoliths_skj$age, Loto=otoliths_skj$len)</pre>
model <- gompertzo(init, dat)</pre>
fit <- nlminb(model$par, model$fn, model$gr,</pre>
            control=list(eval.max=1e4, iter.max=1e4))
model$report()[c("Linf", "k", "tau", "sigma_min")]
# Model 5: Fit to skipjack tags only
```

18 otoliths_had

otoliths_had

Otolith Data (Haddock)

Description

Otolith data for Icelandic haddock from the Icelandic spring bottom trawl survey 2011-2020.

Usage

otoliths had

Format

Data frame containing two columns:

```
age age (years)
len length (cm)
```

Note

The data were contributed by the Icelandic Marine and Freshwater Research Institute. The otoliths were collected following the sampling protocol described in the survey manual (Sólmundsson et al. 2020).

Source

Sólmundsson, J., Karlsson, H., Björnsson, H., Jónsdóttir, I.G., Jakobsdóttir, K.B., and Bogason, V. (2020). *A manual for the Icelandic groundfish survey in spring 2020*. Marine and Freshwater Research in Iceland HV 2020-08.

See Also

gcm, gompertz, gompertzo, richards, richardso, schnute3, vonbert, and vonberto are alternative growth models.

otoliths_had, otoliths_skj, and tags_skj are example datasets.

fishgrowth-package gives an overview of the package.

otoliths_skj

Examples

head(otoliths_had)

otoliths_skj

Otolith Data (Skipjack)

Description

Simulated otolith data, loosely based on a skipjack tuna dataset analyzed by Macdonald et al. (2022).

Usage

```
otoliths_skj
```

Format

Data frame containing two columns:

```
age age (years)
len length (cm)
```

Details

The simulation code that was used to produce this dataset is included in the package:

```
file.show(system.file(package="fishgrowth", "sim/simulate.R"))
```

Source

Macdonald, J., Day, J., Magnusson, A., Maunder, M., Aoki, Y., Matsubara, N., Tsuda, Y., McKechnie, S., Teears, T., Leroy, B., Castillo-Jordán, C., Hampton, J., and Hamer, P. (2022). *Review and new analyses of skipjack growth in the Western and Central Pacific Ocean.* Western and Central Pacific Fisheries Commission Report WCPFC-SC18-2022/SA-IP-06. https://meetings.wcpfc.int/node/16254.

See Also

gcm, gompertz, gompertzo, richards, richardso, schnute3, vonbert, and vonberto are alternative growth models.

otoliths_had, otoliths_skj, and tags_skj are example datasets.

fishgrowth-package gives an overview of the package.

Examples

```
otoliths_skj
```

20 pred_band

pred	band
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Prediction Band

Description

Calculate a prediction band for a fitted growth curve.

Usage

```
pred_band(age, model, level = 0.95)
```

Arguments

age a vector of ages to calculate the prediction band.

model a fitted growth model.

level significance level.

Value

A data frame containing five columns:

age age

Lhat predicted length sigma growth variability

lower limit of prediction band upper upper limit of prediction band

Note

The variability of length at age (sigma) increases linearly with length:

$$\sigma_L = \alpha + \beta \hat{L}$$

This calculation of sigma is demonstrated in the example below.

The lower and upper limits of the prediction band are calculated as $\hat{L}\pm 1.96\sigma_L$ at the 95% significance level.

See Also

gcm, gompertz, gompertzo, richards, richardso, schnute3, vonbert, and vonberto are alternative $growth\ models$.

fishgrowth-package gives an overview of the package.

Examples

```
# Fit a model
init <- list(\log_L 1 = \log(20), \log_L 2 = \log(70), \log_k = \log(0.1),
              log_sigma_min=log(3), log_sigma_max=log(3))
dat <- list(Aoto=otoliths_had$age, Loto=otoliths_had$len, t1=1, t2=10)</pre>
model <- vonbert(init, dat)</pre>
fit <- nlminb(model$par, model$fn, model$gr,</pre>
              control=list(eval.max=1e4, iter.max=1e4))
# Calculate 95% prediction band
x < - seq(1, 18, 0.5)
band <- pred_band(x, model)</pre>
# Plot 95% prediction band
areaplot::confplot(cbind(lower,upper)~age, band, xlim=c(0,18), ylim=c(0,100),
         ylab="len", col="mistyrose")
points(len~age, otoliths_had, xlim=c(0,18), ylim=c(0,100),
       pch=16, col="#0080a010")
lines(Lhat~age, band, lwd=2, col=2)
lines(lower~age, band, lty=1, lwd=0.5, col=2)
lines(upper~age, band, lty=1, lwd=0.5, col=2)
# Calculate sigma by hand
report <- model$report()</pre>
alpha <- report$sigma_intercept</pre>
beta <- report$sigma_slope</pre>
Lhat <- band$Lhat
alpha + beta * Lhat # same values as band$sigma calculated by pred_band()
```

richards

Richards Growth Model

Description

Fit a Richards growth model to otoliths and/or tags, using the Schnute parametrization.

Usage

```
richards(par, data, silent = TRUE, ...)
richards_curve(t, L1, L2, k, b, t1, t2)
richards_objfun(par, data)
```

Arguments

```
par a parameter list.
data a data list.
```

silent	passed to MakeADFun.
	passed to MakeADFun.
t	age (vector).
L1	predicted length at age t1.
L2	predicted length at age t2.
k	growth coefficient.
b	shape parameter.
t1	age where predicted length is L1.
t2	age where predicted length is L2.

Details

The main function richards creates a model object, ready for parameter estimation. The auxiliary functions richards_curve and richards_objfun are called by the main function to calculate the regression curve and objective function value. The user can also call the auxiliary functions directly for plotting and model exploration.

The par list contains the following elements:

- log_L1, predicted length at age t1
- log_L2, predicted length at age t2
- log_k, growth coefficient
- b, shape parameter
- log_sigma_min, growth variability at the shortest observed length in the data
- log_sigma_max (*), growth variability at the longest observed length in the data
- log_age (*), age at release of tagged individuals (vector)
- *: The parameter log_sigma_max can be omitted to estimate growth variability that does not vary with length. The parameter vector log_age can be omitted to fit to otoliths only.

The data list contains the following elements:

- Aoto (*), age from otoliths (vector)
- Loto (*), length from otoliths (vector)
- Lrel (*), length at release of tagged individuals (vector)
- Lrec (*), length at recapture of tagged individuals (vector)
- liberty (*), time at liberty of tagged individuals in years (vector)
- t1, age where predicted length is L1
- t2, age where predicted length is L2

^{*:} The data vectors Aoto and Loto can be omitted to fit to tagging data only. The data vectors Lrel, Lrec, and liberty can be omitted to fit to otoliths only.

Value

The richards function returns a TMB model object, produced by MakeADFun.

The richards_curve function returns a numeric vector of predicted length at age.

The richards_objfun function returns the negative log-likelihood as a single number, describing the goodness of fit of par to the data.

Note

The Schnute parametrization used in richards reduces parameter correlation and improves convergence reliability compared to the traditional parametrization used in richardso. Therefore, the richards parametrization can be recommended for general usage, as both parametrizations produce the same growth curve. However, there can be some use cases where the traditional parametrization (Linf, k, tau, b) is preferred over the Schnute parametrization (L1, L2, k, b).

The Richards (1959) growth model, as parametrized by Schnute (1981, Eq. 15), predicts length at age as:

$$\hat{L}_t = \left[L_1^b + (L_2^b - L_1^b) \frac{1 - e^{-k(t-t_1)}}{1 - e^{-k(t_2-t_1)}} \right]^{1/b}$$

The variability of length at age increases linearly with length,

$$\sigma_L = \alpha + \beta \hat{L}$$

where the slope is $\beta = (\sigma_{\rm max} - \sigma_{\rm min})/(L_{\rm max} - L_{\rm min})$, the intercept is $\alpha = \sigma_{\rm min} - \beta L_{\rm min}$, and $L_{\rm min}$ and $L_{\rm max}$ are the shortest and longest observed lengths in the data. Alternatively, growth variability can be modelled as a constant $\sigma_L = \sigma_{\rm min}$ that does not vary with length, by omitting log_sigma_max from the parameter list (see above).

The negative log-likelihood is calculated by comparing the observed and predicted lengths:

```
nll_Loto <- -dnorm(Loto, Loto_hat, sigma_Loto, TRUE)
nll_Lrel <- -dnorm(Lrel, Lrel_hat, sigma_Lrel, TRUE)
nll_Lrec <- -dnorm(Lrec, Lrec_hat, sigma_Lrec, TRUE)
nll <- sum(nll_Loto) + sum(nll_Lrel) + sum(nll_Lrec)</pre>
```

References

Richards, F.J. (1959). A flexible growth function for empirical use. *Journal of Experimental Botany*, **10**, 290-300. https://www.jstor.org/stable/23686557.

Schnute, J. (1981). A versatile growth model with statistically stable parameters. *Canadian Journal of Fisheries and Aquatic Science*, **38**, 1128-1140. doi:10.1139/f81153.

See Also

gcm, gompertz, gompertzo, richards, richardso, schnute3, vonbert, and vonberto are alternative growth models.

pred_band calculates a prediction band for a fitted growth model.

otoliths_had, otoliths_skj, and tags_skj are example datasets.

fishgrowth-package gives an overview of the package.

Examples

```
# Model 1: Fit to haddock otoliths
# Explore initial parameter values
plot(len~age, otoliths_had, xlim=c(0,18), ylim=c(0,105), pch=16,
     col="#0080a010")
x < - seq(1, 18, 0.1)
lines(x, richards_curve(x, L1=18, L2=67, k=0.1, b=1, t1=1, t2=10), lty=3)
# Prepare parameters and data
init <- list(log_L1=log(18), log_L2=log(67), log_k=log(0.1), b=1,</pre>
            log_sigma_min=log(3), log_sigma_max=log(3))
dat <- list(Aoto=otoliths_had$age, Loto=otoliths_had$len, t1=1, t2=10)</pre>
richards_objfun(init, dat)
# Fit model
model <- richards(init, dat)</pre>
fit <- nlminb(model$par, model$fn, model$gr,</pre>
             control=list(eval.max=1e4, iter.max=1e4))
report <- model$report()</pre>
sdreport <- sdreport(model)</pre>
# Plot results
Lhat <- with(report, richards_curve(x, L1, L2, k, b, t1, t2))
lines(x, Lhat, lwd=2, col=2)
legend("bottomright", c("initial curve", "model fit"), col=c(1,2), lty=c(3,1),
      lwd=c(1,2), bty="n", inset=0.02, y.intersp=1.25)
# Model summary
report[c("L1", "L2", "k", "b", "sigma_min", "sigma_max")]
fit[-1]
summary(sdreport)
# Plot 95% prediction band
band <- pred_band(x, model)</pre>
areaplot::confplot(cbind(lower,upper)~age, band, xlim=c(0,18), ylim=c(0,100),
        ylab="len", col="mistyrose")
points(len~age, otoliths_had, xlim=c(0,18), ylim=c(0,100),
      pch=16, col="#0080a010")
lines(x, Lhat, lwd=2, col=2)
lines(lower~age, band, lty=1, lwd=0.5, col=2)
lines(upper~age, band, lty=1, lwd=0.5, col=2)
# Model 2: Fit to skipjack otoliths and tags
# Explore initial parameter values
plot(len~age, otoliths_skj, xlim=c(0,4), ylim=c(0,100))
x < - seq(0, 4, 0.1)
points(lenRel~I(lenRel/60), tags_skj, col=4)
points(lenRec~I(lenRel/60+liberty), tags_skj, col=3)
```

```
lines(x, richards_curve(x, L1=25, L2=75, k=0.8, b=1, t1=0, t2=4), lty=2)
legend("bottomright", c("otoliths","tag releases","tac recaptures",
      "initial curve"), col=c(1,4,3,1), pch=c(1,1,1,NA), lty=c(0,0,0,2),
      lwd=c(1.2,1.2,1.2,1), bty="n", inset=0.02, y.intersp=1.25)
# Prepare parameters and data
init <- list(log_L1=log(25), log_L2=log(75), log_k=log(0.8), b=1,</pre>
            log_sigma_min=log(3), log_sigma_max=log(3),
            log_age=log(tags_skj$lenRel/60))
dat <- list(Aoto=otoliths_skj$age, Loto=otoliths_skj$len,</pre>
           Lrel=tags_skj$lenRel, Lrec=tags_skj$lenRec,
           liberty=tags_skj$liberty, t1=0, t2=4)
richards_objfun(init, dat)
# Fit model
model <- richards(init, dat)</pre>
fit <- nlminb(model$par, model$fn, model$gr,</pre>
             control=list(eval.max=1e4, iter.max=1e4))
report <- model$report()</pre>
sdreport <- sdreport(model)</pre>
# Plot results
plot(len~age, otoliths_skj, xlim=c(0,4), ylim=c(0,100))
points(report$age, report$Lrel, col=4)
points(report$age+report$liberty, report$Lrec, col=3)
Lhat <- with(report, richards_curve(x, L1, L2, k, b, t1, t2))
lines(x, Lhat, lwd=2)
legend("bottomright", c("otoliths","tag releases","tac recaptures",
      "model fit"), col=c(1,4,3,1), pch=c(1,1,1,NA), lty=c(0,0,0,1),
      lwd=c(1.2,1.2,1.2,2), bty="n", inset=0.02, y.intersp=1.25)
# Model summary
report[c("L1", "L2", "k", "b", "sigma_min", "sigma_max")]
fit[-1]
head(summary(sdreport), 6)
# Model 3: Fit to skipjack otoliths only
init <- list(log_L1=log(25), log_L2=log(75), log_k=log(0.8), b=1,</pre>
            log_sigma_min=log(3), log_sigma_max=log(3))
dat <- list(Aoto=otoliths_skj$age, Loto=otoliths_skj$len, t1=0, t2=4)</pre>
model <- richards(init, dat)</pre>
fit <- nlminb(model$par, model$fn, model$gr,</pre>
             control=list(eval.max=1e4, iter.max=1e4))
model$report()[c("L1", "L2", "k", "b", "sigma_min", "sigma_max")]
# Model 4: Fit to skipjack otoliths only,
# but now estimating constant sigma instead of sigma varying by length
```

```
# We do this by omitting log_sigma_max
init <- list(log_L1=log(25), log_L2=log(75), log_k=log(0.8), b=1,</pre>
            log_sigma_min=log(3))
dat <- list(Aoto=otoliths_skj$age, Loto=otoliths_skj$len, t1=0, t2=4)</pre>
model <- richards(init, dat)</pre>
fit <- nlminb(model$par, model$fn, model$gr,</pre>
             control=list(eval.max=1e4, iter.max=1e4))
model$report()[c("L1", "L2", "k", "b", "sigma_min")]
# Model 5: Fit to skipjack tags only
init <- list(log_L1=log(25), log_L2=log(75), log_k=log(0.8), b=1,</pre>
            log_sigma_min=log(3), log_sigma_max=log(3),
            log_age=log(tags_skj$lenRel/60))
dat <- list(Lrel=tags_skj$lenRel, Lrec=tags_skj$lenRec,</pre>
           liberty=tags_skj$liberty, t1=0, t2=4)
model <- richards(init, dat)</pre>
fit <- nlminb(model$par, model$fn, model$gr,</pre>
             control=list(eval.max=1e4, iter.max=1e4))
model$report()[c("L1", "L2", "k", "b", "sigma_min", "sigma_max")]
```

richardso

Richards Growth Model (Old Style)

Description

Fit a Richards growth model to otoliths and/or tags, using a traditional parametrization.

Usage

```
richardso(par, data, silent = TRUE, ...)
richardso_curve(t, Linf, k, tau, b)
richardso_objfun(par, data)
```

Arguments

par	a parameter list.
data	a data list.
silent	passed to MakeADFun.
• • •	passed to MakeADFun.
t	age (vector).
Linf	asymptotic maximum length.

k	growth coefficient.
tau	location parameter.
b	shape parameter.

Details

The main function richardso creates a model object, ready for parameter estimation. The auxiliary functions richardso_curve and richardso_objfun are called by the main function to calculate the regression curve and objective function value. The user can also call the auxiliary functions directly for plotting and model exploration.

The par list contains the following elements:

- log_Linf, asymptotic maximum length
- log_k, growth coefficient
- tau, location parameter
- · b, shape parameter
- log_sigma_min, growth variability at the shortest observed length in the data
- log_sigma_max (*), growth variability at the longest observed length in the data
- log_age (*), age at release of tagged individuals (vector)
- *: The parameter log_sigma_max can be omitted to estimate growth variability that does not vary with length. The parameter vector log_age can be omitted to fit to otoliths only.

The data list contains the following elements:

- Aoto (*), age from otoliths (vector)
- Loto (*), length from otoliths (vector)
- Lrel (*), length at release of tagged individuals (vector)
- Lrec (*), length at recapture of tagged individuals (vector)
- liberty (*), time at liberty of tagged individuals in years (vector)

Value

The richardso function returns a TMB model object, produced by MakeADFun.

The richardso_curve function returns a numeric vector of predicted length at age.

The richardso_objfun function returns the negative log-likelihood as a single number, describing the goodness of fit of par to the data.

^{*:} The data vectors Aoto and Loto can be omitted to fit to tagging data only. The data vectors Lrel, Lrec, and liberty can be omitted to fit to otoliths only.

Note

The Schnute parametrization used in richards reduces parameter correlation and improves convergence reliability compared to the traditional parametrization used in richardso. Therefore, the richards parametrization can be recommended for general usage, as both parametrizations produce the same growth curve. However, there can be some use cases where the traditional parametrization (Linf, k, tau, b) is preferred over the Schnute parametrization (L1, L2, k, b).

The Richards (1959) growth model, as parametrized by Tjørve and Tjørve (2010, Eq. 4), predicts length at age as:

$$\hat{L}_t = L_{\infty} \left(1 - \frac{1}{b} e^{-k(t-\tau)} \right)^b$$

The variability of length at age increases linearly with length,

$$\sigma_L = \alpha + \beta \hat{L}$$

where the slope is $\beta = (\sigma_{\max} - \sigma_{\min})/(L_{\max} - L_{\min})$, the intercept is $\alpha = \sigma_{\min} - \beta L_{\min}$, and L_{\min} and L_{\max} are the shortest and longest observed lengths in the data. Alternatively, growth variability can be modelled as a constant $\sigma_L = \sigma_{\min}$ that does not vary with length, by omitting log_sigma_max from the parameter list (see above).

The negative log-likelihood is calculated by comparing the observed and predicted lengths:

```
nll_Loto <- -dnorm(Loto, Loto_hat, sigma_Loto, TRUE)
nll_Lrel <- -dnorm(Lrel, Lrel_hat, sigma_Lrel, TRUE)
nll_Lrec <- -dnorm(Lrec, Lrec_hat, sigma_Lrec, TRUE)
nll <- sum(nll_Loto) + sum(nll_Lrel) + sum(nll_Lrec)</pre>
```

References

Richards, F.J. (1959). A flexible growth function for empirical use. *Journal of Experimental Botany*, **10**, 290-300. https://www.jstor.org/stable/23686557.

Tjørve, E. and Tjørve, K.M.C. (2010). A unified approach to the Richards-model family for use in growth analyses: Why we need only two model forms. *Journal of Theoretical Biology*, **267**, 417-425.

See Also

gcm, gompertz, gompertzo, richards, richardso, schnute3, vonbert, and vonberto are alternative growth models.

pred_band calculates a prediction band for a fitted growth model.

otoliths_had, otoliths_skj, and tags_skj are example datasets.

fishgrowth-package gives an overview of the package.

Examples

```
# Model 1: Fit to haddock otoliths
# Explore initial parameter values
plot(len~age, otoliths_had, xlim=c(0,18), ylim=c(0,105), pch=16,
     col="#0080a010")
x < - seq(1, 18, 0.1)
lines(x, richardso_curve(x, Linf=100, k=0.1, tau=-1, b=1), lty=3)
# Prepare parameters and data
init <- list(log_Linf=log(100), log_k=log(0.1), tau=-1, b=1,</pre>
             log_sigma_min=log(3), log_sigma_max=log(3))
dat <- list(Aoto=otoliths_had$age, Loto=otoliths_had$len)</pre>
richardso_objfun(init, dat)
# Fit model
model <- richardso(init, dat)</pre>
fit <- nlminb(model$par, model$fn, model$gr,</pre>
              control=list(eval.max=1e4, iter.max=1e4))
report <- model$report()</pre>
sdreport <- sdreport(model)</pre>
# Plot results
Lhat <- with(report, richardso_curve(x, Linf, k, tau, b))</pre>
lines(x, Lhat, lwd=2, col=2)
legend("bottomright", c("initial curve", "model fit"), col=c(1,2), lty=c(3,1),
       lwd=c(1,2), bty="n", inset=0.02, y.intersp=1.25)
# Model summary
report[c("Linf", "k", "tau", "b", "sigma_min", "sigma_max")]
fit[-1]
summary(sdreport)
# Plot 95% prediction band
band <- pred_band(x, model)</pre>
areaplot::confplot(cbind(lower,upper)~age, band, xlim=c(0,18), ylim=c(0,100),
         ylab="len", col="mistyrose")
points(len~age, otoliths_had, xlim=c(0,18), ylim=c(0,100),
       pch=16, col="#0080a010")
lines(x, Lhat, lwd=2, col=2)
lines(lower~age, band, lty=1, lwd=0.5, col=2)
lines(upper~age, band, lty=1, lwd=0.5, col=2)
# Model 2: Fit to skipjack otoliths and tags
# Explore initial parameter values
plot(len~age, otoliths_skj, xlim=c(0,4), ylim=c(0,100))
x < - seq(0, 4, 0.1)
points(lenRel~I(lenRel/60), tags_skj, col=4)
points(lenRec~I(lenRel/60+liberty), tags_skj, col=3)
```

```
lines(x, richardso_curve(x, Linf=80, k=0.8, tau=-0.5, b=1), lty=2)
legend("bottomright", c("otoliths","tag releases","tac recaptures",
      "initial curve"), col=c(1,4,3,1), pch=c(1,1,1,NA), lty=c(0,0,0,2),
      lwd=c(1.2,1.2,1.2,1), bty="n", inset=0.02, y.intersp=1.25)
# Prepare parameters and data
init <- list(log_Linf=log(80), log_k=log(0.8), tau=-0.5, b=1,</pre>
            log_sigma_min=log(3), log_sigma_max=log(3),
            log_age=log(tags_skj$lenRel/60))
dat <- list(Aoto=otoliths_skj$age, Loto=otoliths_skj$len,</pre>
           Lrel=tags_skj$lenRel, Lrec=tags_skj$lenRec,
           liberty=tags_skj$liberty)
richardso_objfun(init, dat)
# Fit model
model <- richardso(init, dat)</pre>
fit <- nlminb(model$par, model$fn, model$gr,</pre>
             control=list(eval.max=1e4, iter.max=1e4))
report <- model$report()</pre>
sdreport <- sdreport(model)</pre>
# Plot results
plot(len~age, otoliths_skj, xlim=c(0,4), ylim=c(0,100))
points(report$age, report$Lrel, col=4)
points(report$age+report$liberty, report$Lrec, col=3)
Lhat <- with(report, richardso_curve(x, Linf, k, tau, b))</pre>
lines(x, Lhat, lwd=2)
legend("bottomright", c("otoliths","tag releases","tac recaptures",
      "model fit"), col=c(1,4,3,1), pch=c(1,1,1,NA), lty=c(0,0,0,1),
      lwd=c(1.2,1.2,1.2,2), bty="n", inset=0.02, y.intersp=1.25)
# Model summary
report[c("Linf", "k", "tau", "b", "sigma_min", "sigma_max")]
fit[-1]
head(summary(sdreport), 6)
# Model 3: Fit to skipjack otoliths only
init <- list(log_Linf=log(80), log_k=log(0.8), tau=-0.5, b=1,
            log_sigma_min=log(3), log_sigma_max=log(3))
dat <- list(Aoto=otoliths_skj$age, Loto=otoliths_skj$len)</pre>
model <- richardso(init, dat)</pre>
fit <- nlminb(model$par, model$fn, model$gr,</pre>
             control=list(eval.max=1e4, iter.max=1e4))
model$report()[c("Linf", "k", "tau", "b", "sigma_min", "sigma_max")]
# Model 4: Fit to skipjack otoliths only,
# but now estimating constant sigma instead of sigma varying by length
```

schnute3 31

```
# We do this by omitting log_sigma_max
init <- list(log_Linf=log(80), log_k=log(0.8), tau=-0.5, b=1,</pre>
            log_sigma_min=log(3))
dat <- list(Aoto=otoliths_skj$age, Loto=otoliths_skj$len)</pre>
model <- richardso(init, dat)</pre>
fit <- nlminb(model$par, model$fn, model$gr,</pre>
             control=list(eval.max=1e4, iter.max=1e4))
model$report()[c("Linf", "k", "tau", "b", "sigma_min")]
# Model 5: Fit to skipjack tags only
init <- list(log_Linf=log(80), log_k=log(0.8), tau=-0.5, b=1,</pre>
            log_sigma_min=log(3), log_sigma_max=log(3),
            log_age=log(tags_skj$lenRel/60))
dat <- list(Lrel=tags_skj$lenRel, Lrec=tags_skj$lenRec,</pre>
           liberty=tags_skj$liberty)
model <- richardso(init, dat)</pre>
                                       # using 1e3 to keep CRAN checks fast,
fit <- nlminb(model$par, model$fn, model$gr,</pre>
                                                       # but try 1e4 to get
             control=list(eval.max=1e3, iter.max=1e3)) # better convergence
model$report()[c("Linf", "k", "tau", "b", "sigma_min", "sigma_max")]
```

schnute3

Schnute Case 3 Model

Description

Fit a Schnute Case 3 model to otoliths and/or tags.

Usage

```
schnute3(par, data, silent = TRUE, ...)
schnute3_curve(t, L1, L2, b, t1, t2)
schnute3_objfun(par, data)
```

Arguments

par	a parameter list.
data	a data list.
silent	passed to MakeADFun.
	passed to MakeADFun.
t	age (vector).
L1	predicted length at age t1.

32 schnute3

- L2 predicted length at age t2.
 b shape parameter.
 t1 age where predicted length is L1.
- t2 age where predicted length is L2.

Details

The main function schnute3 creates a model object, ready for parameter estimation. The auxiliary functions schnute3_curve and schnute3_objfun are called by the main function to calculate the regression curve and objective function value. The user can also call the auxiliary functions directly for plotting and model exploration.

The par list contains the following elements:

- log_L1, predicted length at age t1
- log_L2, predicted length at age t2
- b, shape parameter
- log_sigma_min, growth variability at the shortest observed length in the data
- log_sigma_max (*), growth variability at the longest observed length in the data
- log_age (*), age at release of tagged individuals (vector)
- *: The parameter log_sigma_max can be omitted to estimate growth variability that does not vary with length. The parameter vector log_age can be omitted to fit to otoliths only.

The data list contains the following elements:

- Aoto (*), age from otoliths (vector)
- Loto (*), length from otoliths (vector)
- Lrel (*), length at release of tagged individuals (vector)
- Lrec (*), length at recapture of tagged individuals (vector)
- liberty (*), time at liberty of tagged individuals in years (vector)
- t1, age where predicted length is L1
- t2, age where predicted length is L2

Value

The schnute3 function returns a TMB model object, produced by MakeADFun.

The schnute3_curve function returns a numeric vector of predicted length at age.

The schnute3_objfun function returns the negative log-likelihood as a single number, describing the goodness of fit of par to the data.

^{*:} The data vectors Aoto and Loto can be omitted to fit to tagging data only. The data vectors Lrel, Lrec, and liberty can be omitted to fit to otoliths only.

schnute3 33

Note

The Schnute Case 3 model is a special case of the Richards (1959) model, where k=0. If the best model fit of a richards model to a particular dataset involves a very small estimated value of k, then the schnute3 model offers a preferable parametrization, as it produces the same curve using fewer parameters.

The Schnute Case 3 model (Schnute 1981, Eq. 17) predicts length at age as:

$$L = \left[L_1^b + (L_2^b - L_1^b) \frac{t - t_1}{t_2 - t_1} \right]^{1/b}$$

The variability of length at age increases linearly with length,

$$\sigma_L = \alpha + \beta \hat{L}$$

where the slope is $\beta = (\sigma_{\rm max} - \sigma_{\rm min})/(L_{\rm max} - L_{\rm min})$, the intercept is $\alpha = \sigma_{\rm min} - \beta L_{\rm min}$, and $L_{\rm min}$ and $L_{\rm max}$ are the shortest and longest observed lengths in the data. Alternatively, growth variability can be modelled as a constant $\sigma_L = \sigma_{\rm min}$ that does not vary with length, by omitting log_sigma_max from the parameter list (see above).

The negative log-likelihood is calculated by comparing the observed and predicted lengths:

```
nll_Loto <- -dnorm(Loto, Loto_hat, sigma_Loto, TRUE)
nll_Lrel <- -dnorm(Lrel, Lrel_hat, sigma_Lrel, TRUE)
nll_Lrec <- -dnorm(Lrec, Lrec_hat, sigma_Lrec, TRUE)
nll <- sum(nll_Loto) + sum(nll_Lrel) + sum(nll_Lrec)</pre>
```

References

Richards, F.J. (1959). A flexible growth function for empirical use. *Journal of Experimental Botany*, **10**, 290-300. https://www.jstor.org/stable/23686557.

Schnute, J. (1981). A versatile growth model with statistically stable parameters. *Canadian Journal of Fisheries and Aquatic Science*, **38**, 1128-1140. doi:10.1139/f81153.

See Also

gcm, gompertz, gompertzo, richards, richardso, schnute3, vonbert, and vonberto are alternative growth models.

pred_band calculates a prediction band for a fitted growth model.

otoliths_had, otoliths_skj, and tags_skj are example datasets.

fishgrowth-package gives an overview of the package.

Examples

34 schnute3

```
x \leftarrow seq(1, 18, 0.1)
lines(x, schnute3_curve(x, L1=15, L2=70, b=2, t1=1, t2=10), lty=3)
# Prepare parameters and data
init <- list(log_L1=log(15), log_L2=log(70), b=2,</pre>
             log_sigma_min=log(3), log_sigma_max=log(3))
dat <- list(Aoto=otoliths_had$age, Loto=otoliths_had$len, t1=1, t2=10)</pre>
schnute3_objfun(init, dat)
# Fit model
model <- schnute3(init, dat)</pre>
fit <- nlminb(model$par, model$fn, model$gr,</pre>
              control=list(eval.max=1e4, iter.max=1e4))
report <- model$report()</pre>
sdreport <- sdreport(model)</pre>
# Plot results
Lhat <- with(report, schnute3_curve(x, L1, L2, b, t1, t2))
lines(x, Lhat, lwd=2, col=2)
legend("bottomright", c("initial curve", "model fit"), col=c(1,2), lty=c(3,1),
       lwd=c(1,2), bty="n", inset=0.02, y.intersp=1.25)
# Model summary
report[c("L1", "L2", "b", "sigma_min", "sigma_max")]
fit[-1]
summary(sdreport)
# Plot 95% prediction band
band <- pred_band(x, model)</pre>
areaplot::confplot(cbind(lower,upper)^age, band, xlim=c(0,18), ylim=c(0,100),
        ylab="len", col="mistyrose")
points(len~age, otoliths_had, xlim=c(0,18), ylim=c(0,100),
       pch=16, col="#0080a010")
lines(x, Lhat, lwd=2, col=2)
lines(lower~age, band, lty=1, lwd=0.5, col=2)
lines(upper~age, band, lty=1, lwd=0.5, col=2)
# Model 2: Fit to skipjack otoliths and tags
# Explore initial parameter values
plot(len~age, otoliths\_skj, xlim=c(0,4), ylim=c(0,100))
x < - seq(0, 4, 0.1)
points(lenRel~I(lenRel/60), tags_skj, col=4)
points(lenRec~I(lenRel/60+liberty), tags_skj, col=3)
lines(x, schnute3_curve(x, L1=25, L2=75, b=3, t1=0, t2=4), lty=2)
legend("bottomright", c("otoliths","tag releases","tac recaptures",
       "initial curve"), col=c(1,4,3,1), pch=c(1,1,1,NA), lty=c(0,0,0,2),
       lwd=c(1.2,1.2,1.2,1), bty="n", inset=0.02, y.intersp=1.25)
# Prepare parameters and data
init <- list(log_L1=log(25), log_L2=log(75), b=3,</pre>
```

schnute3 35

```
log_sigma_min=log(3), log_sigma_max=log(3),
            log_age=log(tags_skj$lenRel/60))
dat <- list(Aoto=otoliths_skj$age, Loto=otoliths_skj$len,</pre>
           Lrel=tags_skj$lenRel, Lrec=tags_skj$lenRec,
           liberty=tags_skj$liberty, t1=0, t2=4)
schnute3_objfun(init, dat)
# Fit model
model <- schnute3(init, dat)</pre>
fit <- nlminb(model$par, model$fn, model$gr,</pre>
             control=list(eval.max=1e4, iter.max=1e4))
report <- model$report()</pre>
sdreport <- sdreport(model)</pre>
# Plot results
plot(len~age, otoliths_skj, xlim=c(0,4), ylim=c(0,100))
points(report$age, report$Lrel, col=4)
points(report$age+report$liberty, report$Lrec, col=3)
Lhat <- with(report, schnute3_curve(x, L1, L2, b, t1, t2))
lines(x, Lhat, lwd=2)
legend("bottomright", c("otoliths","tag releases","tac recaptures",
       "model fit"), col=c(1,4,3,1), pch=c(1,1,1,NA), lty=c(0,0,0,1),
      lwd=c(1.2,1.2,1.2,2), bty="n", inset=0.02, y.intersp=1.25)
# Model summary
report[c("L1", "L2", "b", "sigma_min", "sigma_max")]
fit[-1]
head(summary(sdreport), 5)
# Model 3: Fit to skipjack otoliths only
init <- list(log_L1=log(25), log_L2=log(75), b=3,</pre>
            log_sigma_min=log(3), log_sigma_max=log(3))
dat <- list(Aoto=otoliths_skj$age, Loto=otoliths_skj$len, t1=0, t2=4)</pre>
model <- schnute3(init, dat)</pre>
fit <- nlminb(model$par, model$fn, model$gr,</pre>
             control=list(eval.max=1e4, iter.max=1e4))
model$report()[c("L1", "L2", "b", "sigma_min", "sigma_max")]
# Model 4: Fit to skipjack otoliths only,
# but now estimating constant sigma instead of sigma varying by length
# We do this by omitting log_sigma_max
init <- list(log_L1=log(25), log_L2=log(75), b=3,</pre>
            log_sigma_min=log(3))
dat <- list(Aoto=otoliths_skj$age, Loto=otoliths_skj$len, t1=0, t2=4)</pre>
model <- schnute3(init, dat)</pre>
fit <- nlminb(model$par, model$fn, model$gr,</pre>
             control=list(eval.max=1e4, iter.max=1e4))
```

36 tags_skj

tags_skj

Tagging Data (Skipjack)

Description

Simulated tagging data, loosely based on a skipjack tuna dataset analyzed by Macdonald et al. (2022).

Usage

tags_skj

Format

Data frame containing three columns:

lenRel length at release (cm)
lenRec length at recapture (cm)
liberty time at liberty (years)

Details

The simulation code that was used to produce this dataset is included in the package:

```
file.show(system.file(package="fishgrowth", "sim/simulate.R"))
```

Source

Macdonald, J., Day, J., Magnusson, A., Maunder, M., Aoki, Y., Matsubara, N., Tsuda, Y., McKechnie, S., Teears, T., Leroy, B., Castillo-Jordán, C., Hampton, J., and Hamer, P. (2022). *Review and new analyses of skipjack growth in the Western and Central Pacific Ocean*. Western and Central Pacific Fisheries Commission Report WCPFC-SC18-2022/SA-IP-06. https://meetings.wcpfc.int/node/16254.

See Also

```
gcm, gompertz, gompertzo, richards, richardso, schnute3, vonbert, and vonberto are alternative growth models.
```

```
otoliths_had, otoliths_skj, and tags_skj are example datasets.
```

fishgrowth-package gives an overview of the package.

Examples

```
head(tags_skj)
```

vonbert

Von Bertalanffy Growth Model

Description

Fit a von Bertalanffy growth model to otoliths and/or tags, using the Schnute-Fournier parametrization.

Usage

```
vonbert(par, data, silent = TRUE, ...)
vonbert_curve(t, L1, L2, k, t1, t2)
vonbert_objfun(par, data)
```

Arguments

par	a parameter list.
data	a data list.
silent	passed to MakeADFun.
• • •	passed to MakeADFun.
t	age (vector).
L1	predicted length at age t1.
L2	predicted length at age t2.
k	growth coefficient.
t1	age where predicted length is L1.
t2	age where predicted length is L2.

Details

The main function vonbert creates a model object, ready for parameter estimation. The auxiliary functions vonbert_curve and vonbert_objfun are called by the main function to calculate the regression curve and objective function value. The user can also call the auxiliary functions directly for plotting and model exploration.

The par list contains the following elements:

- log_L1, predicted length at age t1
- log_L2, predicted length at age t2
- log_k, growth coefficient
- log_sigma_min, growth variability at the shortest observed length in the data
- log_sigma_max (*), growth variability at the longest observed length in the data
- log_age (*), age at release of tagged individuals (vector)
- *: The parameter log_sigma_max can be omitted to estimate growth variability that does not vary with length. The parameter vector log_age can be omitted to fit to otoliths only.

The data list contains the following elements:

- Aoto (*), age from otoliths (vector)
- Loto (*), length from otoliths (vector)
- Lrel (*), length at release of tagged individuals (vector)
- Lrec (*), length at recapture of tagged individuals (vector)
- liberty (*), time at liberty of tagged individuals in years (vector)
- t1, age where predicted length is L1
- t2, age where predicted length is L2
- *: The data vectors Aoto and Loto can be omitted to fit to tagging data only. The data vectors Lrel, Lrec, and liberty can be omitted to fit to otoliths only.

Value

The vonbert function returns a TMB model object, produced by MakeADFun.

The vonbert_curve function returns a numeric vector of predicted length at age.

The vonbert_objfun function returns the negative log-likelihood as a single number, describing the goodness of fit of par to the data.

Note

The Schnute-Fournier parametrization used in vonbert reduces parameter correlation and improves convergence reliability compared to the traditional parametrization used in vonberto. Therefore, the vonbert parametrization can be recommended for general usage, as both parametrizations produce the same growth curve. However, there can be some use cases where the traditional parametrization (Linf, k, t0) is preferred over the Schnute-Fournier parametrization (L1, L2, k).

The von Bertalanffy (1938) growth model, as parametrized by Schnute and Fournier (1980), predicts length at age as:

$$\hat{L}_t = L_1 + (L_2 - L_1) \frac{1 - e^{-k(t-t_1)}}{1 - e^{-k(t_2-t_1)}}$$

The variability of length at age increases linearly with length,

$$\sigma_L = \alpha + \beta \hat{L}$$

where the slope is $\beta = (\sigma_{\max} - \sigma_{\min})/(L_{\max} - L_{\min})$, the intercept is $\alpha = \sigma_{\min} - \beta L_{\min}$, and L_{\min} and L_{\max} are the shortest and longest observed lengths in the data. Alternatively, growth variability can be modelled as a constant $\sigma_L = \sigma_{\min}$ that does not vary with length, by omitting log_sigma_max from the parameter list (see above).

The negative log-likelihood is calculated by comparing the observed and predicted lengths:

```
nll_Loto <- -dnorm(Loto, Loto_hat, sigma_Loto, TRUE)
nll_Lrel <- -dnorm(Lrel, Lrel_hat, sigma_Lrel, TRUE)
nll_Lrec <- -dnorm(Lrec, Lrec_hat, sigma_Lrec, TRUE)
nll <- sum(nll_Loto) + sum(nll_Lrel) + sum(nll_Lrec)</pre>
```

References

von Bertalanffy, L. (1938). A quantitative theory of organic growth. *Human Biology*, **10**, 181-213. https://www.jstor.org/stable/41447359.

Schnute, J. and Fournier, D. (1980). A new approach to length-frequency analysis: Growth structure. *Canadian Journal of Fisheries and Aquatic Science*, **37**, 1337-1351. doi:10.1139/f80172.

See Also

gcm, gompertz, gompertzo, richards, richardso, schnute3, vonbert, and vonberto are alternative growth models.

pred_band calculates a prediction band for a fitted growth model.

otoliths_had, otoliths_skj, and tags_skj are example datasets.

fishgrowth-package gives an overview of the package.

Examples

```
# Fit model
model <- vonbert(init, dat)</pre>
fit <- nlminb(model$par, model$fn, model$gr,</pre>
              control=list(eval.max=1e4, iter.max=1e4))
report <- model$report()</pre>
sdreport <- sdreport(model)</pre>
# Plot results
Lhat <- with(report, vonbert_curve(x, L1, L2, k, t1, t2))
lines(x, Lhat, lwd=2, col=2)
legend("bottomright", c("initial curve", "model fit"), col=c(1,2), lty=c(3,1),
       lwd=c(1,2), bty="n", inset=0.02, y.intersp=1.25)
# Model summary
report[c("L1", "L2", "k", "sigma_min", "sigma_max")]
fit[-1]
summary(sdreport)
# Plot 95% prediction band
band <- pred_band(x, model)</pre>
areaplot::confplot(cbind(lower,upper)~age, band, xlim=c(0,18), ylim=c(0,100),
         ylab="len", col="mistyrose")
points(len~age, otoliths_had, xlim=c(0,18), ylim=c(0,100),
       pch=16, col="#0080a010")
lines(x, Lhat, lwd=2, col=2)
lines(lower~age, band, lty=1, lwd=0.5, col=2)
lines(upper~age, band, lty=1, lwd=0.5, col=2)
# Model 2: Fit to skipjack otoliths and tags
# Explore initial parameter values
plot(len~age, otoliths_skj, xlim=c(0,4), ylim=c(0,100))
x < - seq(0, 4, 0.1)
points(lenRel~I(lenRel/60), tags_skj, col=4)
points(lenRec~I(lenRel/60+liberty), tags_skj, col=3)
lines(x, vonbert_curve(x, L1=25, L2=75, k=0.8, t1=0, t2=4), lty=2)
legend("bottomright", c("otoliths","tag releases","tac recaptures",
       "initial curve"), col=c(1,4,3,1), pch=c(1,1,1,NA), lty=c(0,0,0,2),
       lwd=c(1.2,1.2,1.2,1), bty="n", inset=0.02, y.intersp=1.25)
# Prepare parameters and data
init <- list(log_L1=log(25), log_L2=log(75), log_k=log(0.8),</pre>
             log_sigma_min=log(3), log_sigma_max=log(3),
             log_age=log(tags_skj$lenRel/60))
dat <- list(Aoto=otoliths_skj$age, Loto=otoliths_skj$len,</pre>
            Lrel=tags_skj$lenRel, Lrec=tags_skj$lenRec,
            liberty=tags_skj$liberty, t1=0, t2=4)
vonbert_objfun(init, dat)
# Fit model
```

```
model <- vonbert(init, dat)</pre>
fit <- nlminb(model$par, model$fn, model$gr,</pre>
            control=list(eval.max=1e4, iter.max=1e4))
report <- model$report()</pre>
sdreport <- sdreport(model)</pre>
# Plot results
plot(len~age, otoliths_skj, xlim=c(0,4), ylim=c(0,100))
points(report$age, report$Lrel, col=4)
points(report$age+report$liberty, report$Lrec, col=3)
Lhat <- with(report, vonbert_curve(x, L1, L2, k, t1, t2))
lines(x, Lhat, lwd=2)
legend("bottomright", c("otoliths","tag releases","tac recaptures",
      "model fit"), col=c(1,4,3,1), pch=c(1,1,1,NA), lty=c(0,0,0,1),
      lwd=c(1.2,1.2,1.2,2), bty="n", inset=0.02, y.intersp=1.25)
# Model summary
report[c("L1", "L2", "k", "sigma_min", "sigma_max")]
fit[-1]
head(summary(sdreport), 5)
# Model 3: Fit to skipjack otoliths only
init \leftarrow list(log_L1=log(25), log_L2=log(75), log_k=log(0.8),
            log_sigma_min=log(3), log_sigma_max=log(3))
dat <- list(Aoto=otoliths_skj$age, Loto=otoliths_skj$len, t1=0, t2=4)</pre>
model <- vonbert(init, dat)</pre>
fit <- nlminb(model$par, model$fn, model$gr,</pre>
            control=list(eval.max=1e4, iter.max=1e4))
model$report()[c("L1", "L2", "k", "sigma_min", "sigma_max")]
# Model 4: Fit to skipjack otoliths only,
# but now estimating constant sigma instead of sigma varying by length
# We do this by omitting log_sigma_max
init <- list(log_L1 = log(25), log_L2 = log(75), log_k = log(0.8),
            log_sigma_min=log(3))
dat <- list(Aoto=otoliths_skj$age, Loto=otoliths_skj$len, t1=0, t2=4)</pre>
model <- vonbert(init, dat)</pre>
fit <- nlminb(model$par, model$fn, model$gr,</pre>
             control=list(eval.max=1e4, iter.max=1e4))
model$report()[c("L1", "L2", "k", "sigma_min")]
# Model 5: Fit to skipjack tags only
init \leftarrow list(log_L1=log(25), log_L2=log(75), log_k=log(0.8),
            log_sigma_min=log(3), log_sigma_max=log(3),
```

vonberto

Von Bertalanffy Growth Model (Old Style)

Description

Fit a von Bertalanffy growth model to otoliths and/or tags, using a traditional parametrization.

Usage

```
vonberto(par, data, silent = TRUE, ...)
vonberto_curve(t, Linf, k, t0)
vonberto_objfun(par, data)
```

Arguments

par	a parameter list.
data	a data list.
silent	passed to MakeADFun.
	passed to MakeADFun.
t	age (vector).
Linf	asymptotic maximum length.
k	growth coefficient.
t0	age where the predicted length is zero, the x-intercept.

Details

The main function vonberto creates a model object, ready for parameter estimation. The auxiliary functions vonberto_curve and vonberto_objfun are called by the main function to calculate the regression curve and objective function value. The user can also call the auxiliary functions directly for plotting and model exploration.

The par list contains the following elements:

- log_Linf, asymptotic maximum length
- log_k, growth coefficient

- to, age where the predicted length is zero, the x-intercept
- log_sigma_min, growth variability at the shortest observed length in the data
- log_sigma_max (*), growth variability at the longest observed length in the data
- log_age (*), age at release of tagged individuals (vector)
- *: The parameter log_sigma_max can be omitted to estimate growth variability that does not vary with length. The parameter vector log_age can be omitted to fit to otoliths only.

The data list contains the following elements:

- Aoto (*), age from otoliths (vector)
- Loto (*), length from otoliths (vector)
- Lrel (*), length at release of tagged individuals (vector)
- Lrec (*), length at recapture of tagged individuals (vector)
- liberty (*), time at liberty of tagged individuals in years (vector)
- *: The data vectors Aoto and Loto can be omitted to fit to tagging data only. The data vectors Lrel, Lrec, and liberty can be omitted to fit to otoliths only.

Value

The vonberto function returns a TMB model object, produced by MakeADFun.

The vonberto_curve function returns a numeric vector of predicted length at age.

The vonberto_objfun function returns the negative log-likelihood as a single number, describing the goodness of fit of par to the data.

Note

The Schnute-Fournier parametrization used in vonbert reduces parameter correlation and improves convergence reliability compared to the traditional parametrization used in vonberto. Therefore, the vonbert parametrization can be recommended for general usage, as both parametrizations produce the same growth curve. However, there can be some use cases where the traditional parametrization (Linf, k, t0) is preferred over the Schnute-Fournier parametrization (L1, L2, k).

The von Bertalanffy (1938) growth model, as parametrized by Beverton and Holt (1957), predicts length at age as:

$$\hat{L}_t = L_{\infty} \left(1 - e^{-k(t-t_0)} \right)$$

The variability of length at age increases linearly with length,

$$\sigma_L = \alpha + \beta \hat{L}$$

where the slope is $\beta = (\sigma_{\rm max} - \sigma_{\rm min})/(L_{\rm max} - L_{\rm min})$, the intercept is $\alpha = \sigma_{\rm min} - \beta L_{\rm min}$, and $L_{\rm min}$ and $L_{\rm max}$ are the shortest and longest observed lengths in the data. Alternatively, growth variability can be modelled as a constant $\sigma_L = \sigma_{\rm min}$ that does not vary with length, by omitting log_sigma_max from the parameter list (see above).

The negative log-likelihood is calculated by comparing the observed and predicted lengths:

```
nll_Loto <- -dnorm(Loto, Loto_hat, sigma_Loto, TRUE)
nll_Lrel <- -dnorm(Lrel, Lrel_hat, sigma_Lrel, TRUE)
nll_Lrec <- -dnorm(Lrec, Lrec_hat, sigma_Lrec, TRUE)
nll <- sum(nll_Loto) + sum(nll_Lrel) + sum(nll_Lrec)</pre>
```

References

von Bertalanffy, L. (1938). A quantitative theory of organic growth. *Human Biology*, **10**, 181-213. https://www.jstor.org/stable/41447359.

Beverton, R.J.H. and Holt, S.J. (1957). *On the dynamics of exploited fish populations*. London: Her Majesty's Stationery Office.

See Also

gcm, gompertz, gompertzo, richards, richards, schnute3, vonbert, and vonberto are alternative growth models.

pred_band calculates a prediction band for a fitted growth model.

otoliths_had, otoliths_skj, and tags_skj are example datasets.

fishgrowth-package gives an overview of the package.

Examples

```
# Model 1: Fit to haddock otoliths
# Explore initial parameter values
plot(len~age, otoliths_had, xlim=c(0,18), ylim=c(0,105), pch=16,
     col="#0080a010")
x \leftarrow seq(1, 18, 0.1)
lines(x, vonberto_curve(x, Linf=100, k=0.1, t0=-1), lty=3)
# Prepare parameters and data
init <- list(log_Linf=log(100), log_k=log(0.1), t0=-1,</pre>
              log_sigma_min=log(3), log_sigma_max=log(3))
dat <- list(Aoto=otoliths_had$age, Loto=otoliths_had$len)</pre>
vonberto_objfun(init, dat)
# Fit model
model <- vonberto(init, dat)</pre>
fit <- nlminb(model$par, model$fn, model$gr,</pre>
              control=list(eval.max=1e4, iter.max=1e4))
report <- model$report()</pre>
sdreport <- sdreport(model)</pre>
# Plot results
Lhat <- with(report, vonberto_curve(x, Linf, k, t0))</pre>
lines(x, Lhat, lwd=2, col=2)
legend("bottomright", c("initial curve", "model fit"), col=c(1,2), lty=c(3,1),
       lwd=c(1,2), bty="n", inset=0.02, y.intersp=1.25)
# Model summary
```

```
report[c("Linf", "k", "t0", "sigma_min", "sigma_max")]
fit[-1]
summary(sdreport)
# Plot 95% prediction band
band <- pred_band(x, model)</pre>
areaplot::confplot(cbind(lower,upper)~age, band, xlim=c(0,18), ylim=c(0,100),
        ylab="len", col="mistyrose")
points(len~age, otoliths_had, xlim=c(0,18), ylim=c(0,100),
      pch=16, col="#0080a010")
lines(x, Lhat, lwd=2, col=2)
lines(lower~age, band, lty=1, lwd=0.5, col=2)
lines(upper~age, band, lty=1, lwd=0.5, col=2)
# Model 2: Fit to skipjack otoliths and tags
# Explore initial parameter values
plot(len~age, otoliths_skj, xlim=c(0,4), ylim=c(0,100))
x < - seq(0, 4, 0.1)
points(lenRel~I(lenRel/60), tags_skj, col=4)
points(lenRec~I(lenRel/60+liberty), tags_skj, col=3)
lines(x, vonberto_curve(x, Linf=80, k=0.8, t0=-0.5), lty=2)
legend("bottomright", c("otoliths","tag releases","tac recaptures",
       "initial curve"), col=c(1,4,3,1), pch=c(1,1,1,NA), lty=c(0,0,0,2),
      lwd=c(1.2,1.2,1.2,1), bty="n", inset=0.02, y.intersp=1.25)
# Prepare parameters and data
init <- list(log_Linf=log(80), log_k=log(0.8), t0=-0.5,
            log_sigma_min=log(3), log_sigma_max=log(3),
            log_age=log(tags_skj$lenRel/60))
dat <- list(Aoto=otoliths_skj$age, Loto=otoliths_skj$len,</pre>
           Lrel=tags_skj$lenRel, Lrec=tags_skj$lenRec,
           liberty=tags_skj$liberty)
vonberto_objfun(init, dat)
# Fit model
model <- vonberto(init, dat)</pre>
fit <- nlminb(model$par, model$fn, model$gr,</pre>
              control=list(eval.max=1e4, iter.max=1e4))
report <- model$report()</pre>
sdreport <- sdreport(model)</pre>
# Plot results
plot(len~age, otoliths_skj, xlim=c(0,4), ylim=c(0,100))
points(report$age, report$Lrel, col=4)
points(report$age+report$liberty, report$Lrec, col=3)
Lhat <- with(report, vonberto_curve(x, Linf, k, t0))
lines(x, Lhat, lwd=2)
legend("bottomright", c("otoliths","tag releases","tac recaptures",
       "model fit"), col=c(1,4,3,1), pch=c(1,1,1,NA), lty=c(0,0,0,1),
      lwd=c(1.2,1.2,1.2,2), bty="n", inset=0.02, y.intersp=1.25)
```

```
# Model summary
report[c("Linf", "k", "t0", "sigma_min", "sigma_max")]
fit[-1]
head(summary(sdreport), 5)
# Model 3: Fit to skipjack otoliths only
init <- list(log_Linf=log(80), log_k=log(0.8), t0=-0.5,</pre>
           log_sigma_min=log(3), log_sigma_max=log(3))
dat <- list(Aoto=otoliths_skj$age, Loto=otoliths_skj$len)</pre>
model <- vonberto(init, dat)</pre>
fit <- nlminb(model$par, model$fn, model$gr,</pre>
            control=list(eval.max=1e4, iter.max=1e4))
model$report()[c("Linf", "k", "t0", "sigma_min", "sigma_max")]
# Model 4: Fit to skipjack otoliths only,
# but now estimating constant sigma instead of sigma varying by length
# We do this by omitting log_sigma_max
init <- list(log_Linf=log(80), log_k=log(0.8), t0=-0.5,
           log_sigma_min=log(3))
dat <- list(Aoto=otoliths_skj$age, Loto=otoliths_skj$len)</pre>
model <- vonberto(init, dat)</pre>
fit <- nlminb(model$par, model$fn, model$gr,</pre>
            control=list(eval.max=1e4, iter.max=1e4))
model$report()[c("Linf", "k", "t0", "sigma_min")]
# Model 5: Fit to skipjack tags only
init <- list(log_Linf=log(80), log_k=log(0.8), t0=-0.5,</pre>
           log_sigma_min=log(3), log_sigma_max=log(3),
           log_age=log(tags_skj$lenRel/60))
dat <- list(Lrel=tags_skj$lenRel, Lrec=tags_skj$lenRec,</pre>
          liberty=tags_skj$liberty)
model <- vonberto(init, dat)</pre>
fit <- nlminb(model$par, model$fn, model$gr,</pre>
            control=list(eval.max=1e4, iter.max=1e4))
model$report()[c("Linf", "k", "t0", "sigma_min", "sigma_max")]
```

Index

```
fishgrowth (fishgrowth-package), 2
fishgrowth-package, 2
gcm, 2, 3, 10, 15, 18–20, 23, 28, 33, 37, 39, 44
gcm_curve (gcm), 3
gcm_objfun (gcm), 3
gompertz, 2, 5, 8, 14, 15, 18-20, 23, 28, 33,
         37, 39, 44
gompertz_curve (gompertz), 8
gompertz_objfun (gompertz), 8
gompertzo, 2, 5, 9, 10, 13, 18-20, 23, 28, 33,
         37, 39, 44
gompertzo_curve (gompertzo), 13
gompertzo_objfun (gompertzo), 13
MakeADFun, 3, 4, 8, 9, 13, 14, 22, 23, 26, 27,
         31, 32, 37, 38, 42, 43
otoliths_had, 2, 5, 10, 15, 18, 19, 23, 28, 33,
         37, 39, 44
otoliths_skj, 2, 5, 10, 15, 18, 19, 23, 28, 33,
         37, 39, 44
pred_band, 2, 5, 10, 15, 20, 23, 28, 33, 39, 44
richards, 2, 5, 10, 14, 15, 18-20, 21, 28, 33,
         37, 39, 44
richards_curve (richards), 21
richards_objfun (richards), 21
richardso, 2, 5, 18-20, 23, 26, 33, 37, 39
richardso_curve (richardso), 26
richardso_objfun (richardso), 26
RTMB, 2
schnute3, 2, 5, 10, 15, 18–20, 23, 28, 31, 37,
         39, 44
schnute3_curve (schnute3), 31
schnute3_objfun (schnute3), 31
tags_skj, 3, 5, 10, 15, 18, 19, 23, 28, 33, 36,
         39, 44
```

```
vonbert, 2, 5, 10, 15, 18–20, 23, 28, 33, 37, 37, 43, 44 vonbert_curve (vonbert), 37 vonbert_objfun (vonbert), 37 vonberto, 2, 5, 10, 15, 18–20, 23, 28, 33, 37–39, 42 vonberto_curve (vonberto), 42 vonberto_objfun (vonberto), 42
```