Package: ecostate (via r-universe)

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Type Package

Title State-Space Mass-Balance Model for Marine Ecosystems

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Description Fits a state-space mass-balance model for marine ecosystems, which implements dynamics derived from `Ecopath with Ecosim` (`EwE`) <https://ecopath.org/> while fiting to time-series of fishery catch, biomass indices, age-composition samples, and weight-at-age data. `Ecostate` fits biological parameters (e.g., equilibrium mass) and measurement parameters (e.g., catchability coefficients) jointly with residual variation in process errors, and can include Bayesian priors for parameters.

License GPL-3

Depends R (>= 4.1.0), RTMB (>= 1.5.0),

Imports TMB, MASS, checkmate, ggplot2, ggnetwork, igraph

Suggests knitr, rmarkdown

Roxygen list(markdown = TRUE)

RoxygenNote 7.3.2

VignetteBuilder knitr

LazyData true

URL https://james-thorson-noaa.github.io/ecostate/

BugReports https://github.com/James-Thorson-NOAA/ecostate/issues

Config/pak/sysreqs libglpk-dev libxml2-dev

Repository https://james-thorson-noaa.r-universe.dev

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abm3pc_sys Adams-Ba

Adams-Bashford-Moulton for system of equations

Description

Third-order Adams-Bashford-Moulton predictor-corrector method.

Usage

```
abm3pc_sys(f, a, b, y0, n, Pars, ...)
```

| f | function in the differential equation $y' = f(x, y)$; defined as a function $R \times R^m \to R^m$, where m is the number of equations. |
|------|---|
| а | starting time for the interval to integrate |
| b | ending time for the interval to integrate. |
| уØ | starting values at time a |
| n | number of steps |
| Pars | named list of parameters passed to f |
| | additional inputs to function f |

Details

Combined Adams-Bashford and Adams-Moulton (or: multi-step) method of third order with corrections according to the predictor-corrector approach. Copied from pracma under GPL-3, with small modifications to work with RTMB

Value

List with components x for grid points between a and b and y an n-by-m matrix with solutions for variables in columns, i.e. each row contains one time stamp.

add_equilibrium Compute equilibrium values

Description

Compute equilibrium values

Usage

```
add_equilibrium(ecoparams, scale_solver, noB_i, type_i)
```

Arguments

| ecoparams | list of parameters |
|--------------|---|
| scale_solver | Whether to solve for ecotrophic efficiency EE given biomass B (scale_solver="simple") or solve for a combination of EE and B values |
| noB_i | Boolean vector indicating which taxa have no B value |
| type_i | character vector indicating whether a taxon is "hetero", "auto", or "detritus" |

Details

Replaces NA values in ecotrophic efficiency and/or biomass with equilibrium solution, and then calculates equilibrium consumption, natural mortality, and other rates.

Value

the list of parameters with missing values in ecoparams\$B_i and/or ecoparams\$EE_i filled in, as well as additional values Qe_ij, m0_i, and GE_i

compute_nll

Description

Compute negative log-likelihood for EcoState model

Usage

```
compute_nll(
 p,
  taxa,
 years,
 noB_i,
  type_i,
 n_species,
 project_vars,
 DC_ij,
 Bobs_ti,
 Cobs_ti,
 Nobs_ta_g2,
 Wobs_ta_g2,
  log_prior,
  fit_eps,
  fit_nu,
  stanza_data,
  settings,
  control
)
```

| р | list of parameters |
|--------------|--|
| taxa | Character vector of taxa included in model. |
| years | Integer-vector of years included in model |
| noB_i | Boolean vector indicating which taxa have no B value |
| type_i | character vector indicating whether a taxon is "hetero", "auto", or "detritus" |
| n_species | number of species |
| project_vars | function to integrate differential equation |
| DC_ij | Diet projections matrix |
| Bobs_ti | formatted matrix of biomass data |
| Cobs_ti | formatted matrix of catch data |
| Nobs_ta_g2 | formatted list of age-comp data |
| Wobs_ta_g2 | formatted list of weight-at-age data |

| log_prior | A user-provided function that takes as input the list of parameters out\$obj\$env\$parList() where out is the output from ecostate(), and returns a numeric vector where |
|-------------|--|
| | the sum is the log-prior probability. For example log_prior = function(p) |
| | dnorm(p\$logq_i[1], mean=0, sd=0.1, log=TRUE) specifies a lognormal prior |
| | probability for the catchability coefficient for the first taxa with logmean of zero and logsd of 0.1 |
| fit_eps | Character-vector listing taxa for which the model should estimate annual process errors in dB/dt |
| fit_nu | Character-vector listing taxa for which the model should estimate annual process errors in consumption Q_{ij} |
| stanza_data | output from make_stanza_data |
| settings | Output from stanza_settings(), used to define age-structured dynamics (called stanza-groups). |
| control | output from ecostate_control |

Details

Given a list of parameters, calculates the joint negative log-likelihood, where the Laplace approximation is then used to marginalize across random effects to calculate the log-marginal likelihood of fixed effects. The joint likelihood includes the fit to fishery catches, biomass indices, agecomposition data, weight-at-age data, priors, and the distribution for random effects.

Value

The joint negative log-likelihood including contribution of priors and fit to data.

| compute_tracer | Calculate tracers, e.g., trophic level | |
|----------------|--|--|
|----------------|--|--|

Description

Calculate how a tracer propagates through consumption.

Usage

```
compute_tracer(
   Q_ij,
   inverse_method = c("Penrose_moore", "Standard"),
   type_i,
   tracer_i = rep(1, nrow(Q_ij))
)
```

| Q_ij | Consumption of each prey i by predator j in units biomass. |
|----------------|--|
| inverse_method | whether to use pseudoinverse or standard inverse |
| type_i | character vector indicating whether a taxon is "hetero", "auto", or "detritus" |
| tracer_i | an indicator matrix specifying the traver value |

Details

Trophic level y_i for each predator *i* is defined as:

$$\mathbf{y} = \mathbf{l}\mathbf{Q}^* + \mathbf{1}$$

where \mathbf{Q}_* is the proportion consumption for each predator (column) of different prey (rows). We identify primary producers as any taxa with no consumption (a column of 0s), and assign them as the first trophic level.

More generically, a tracer might be used to track movement of biomass through consumption. For example, if we have a tracer x_i that is 1 for the base of the pelagic food chain, and 0 otherwise, then we can calculate the proportion of pelagic vs. nonpelagic biomass for each taxon:

```
\mathbf{y} = \mathbf{l} \mathbf{Q}^* + \mathbf{x}
```

This then allows us to separate alternative components of the foodweb.

Value

The vector

yi resulting from tracer $\mathbf{x_i}$

dBdt

Dynamics from EcoSim

Description

Compute system of differential equations representing EcoState dynamics derived from EcoSim.

Usage

```
dBdt(
   Time,
   State,
   Pars,
   type_i,
   n_species,
   F_type = "integrated",
   what = "dBdt"
)
```

ddirmult

Arguments

| Time | not used |
|-----------|---|
| State | vector of state variables to integrate |
| Pars | list of parameters governing the ODE |
| type_i | type for each taxon |
| n_species | number of species |
| F_type | whether to integrate catches along with biomass ("integrated") or calculate catches from the Baranov catch equation applied to average biomass ("averaged") |
| what | what output to produce |

Details

This function has syntax designed to match pracma solvers.

Value

An object (list) of ranges. Elements include:

- G_i Biomass growth per time
- **g_i** Biomass growth per time per biomass
- M2_i Consumptive mortality per time
- $m2_i$ Consumptive mortality per time per biomass
- M_i Natural mortality per time
- **m_i** Natural mortality per time per biomass (i.e., m2_i + m0_i)
- Q_ij Consumption per time for prey (rows) by predator (columns)

ddirmult

Dirichlet-multinomial

Description

Allows data-weighting as parameter

Usage

```
ddirmult(x, prob, ln_theta, log = TRUE)
```

| х | numeric vector of observations across categories |
|----------|--|
| prob | numeric vector of category probabilities |
| ln_theta | logit-ratio of effective and input sample size |
| log | whether to return the log-probability or not |

Value

The log-likelihood resulting from the Dirichlet-multinomial distribution

Examples

```
library(RTMB)
prob = rep(0.1,10)
x = rmultinom( n=1, prob=prob, size=20 )[,1]
f = function( ln_theta ) ddirmult(x, prob, ln_theta)
f( 0 )
F = MakeTape(f, 0)
F$jacfun()(0)
```

eastern_bering_sea eastern Bering Sea ecosystem data

Description

Data used to demonstrate a Model of Intermediate Complexity (MICE) for the eastern Bering Sea. data(eastern_bering_sea) loads a list that includes four components:

- Survey is a long-form data-frame with three columns, providing the Year, Mass (in relative units for most taxa, and million metric tons for Pollock, Cod, Arrowtooth, and NFS), and Taxon for each year with available data
- Catch is a long-form data-frame with three columns, providing the Year, Mass (in million metric tons), and Taxon for each year with available data
- P_over_B is a numeric vector with the unitless ratio of biomass production to population biomass for each taxon
- Q_over_B is a numeric vector with the unitless ratio of biomass consumption to population biomass for each taxon
- Diet_proportions is a numeric matrix where each column lists the proportion of biomass consumed that is provided by each prey (row)

Usage

```
data(eastern_bering_sea)
```

Details

The data compiled come from a variety of sources:

• Northern fur seal (NFS) survey is an absolute index, corrected for proportion of time spent in the eastern Bering Sea. NFS QB is developed from a bioenergetic model and also corrected for seasonal residency. Both are provided by Elizabeth McHuron. It is post-processed in a variety of ways, and not to be treated as an index of abundance for NFS for other uses.

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- Pollock, cod, and arrowtooth surveys are from a bottom trawl survey, and cod and arrowtooth are treated as an absolute index.
- Copepod and other zooplankton are from an oblique tow bongo net survey, with data provided by Dave Kimmel. It is then post-processed to account for spatially and seaonally imbalanced data.
- Other P_over_B, Q_over_B and Diet_proportions values are derived from Rpath models, provided by Andy Whitehouse.
- Primary producers is an annual index of relative biomass, developed from monthly satellite measurements and provided by Jens Nielsen. See Thorson et al. (In review) for more details regarding data standardization and sources

ecostate

fit EcoState model

Description

Estimate parameters for an EcoState model

Usage

```
ecostate(
  taxa,
 years,
 catch = data.frame(Year = numeric(0), Mass = numeric(0), Taxon = numeric(0)),
 biomass = data.frame(Year = numeric(0), Mass = numeric(0), Taxon = numeric(0)),
 agecomp = list(),
 weight = list(),
 ΡB,
  QB,
 Β,
 DC.
 EE,
  Х,
  type,
 U,
  fit_B = vector(),
  fit_Q = vector(),
  fit_B0 = vector(),
  fit_EE = vector(),
  fit_PB = vector(),
  fit_QB = vector(),
  fit_eps = vector(),
  fit_nu = vector(),
  log_prior = function(p) 0,
  settings = stanza_settings(taxa = taxa),
  control = ecostate_control()
)
```

| taxa | Character vector of taxa included in model. |
|---------|--|
| years | Integer-vector of years included in model |
| catch | long-form data frame with columns Mass, Year and Taxon |
| biomass | long-form data frame with columns Mass, Year and Taxon, where Mass is as- sumed to have the same units as catch |
| agecomp | a named list, with names corresponding to stanza_groups, where each list- element is a matrix with rownames for years and colnames for integer ages, where NA excludes the entry from inclusion and the model computes the likeli- hood across included ages in a given year, and the rowsum is the input-sample size for a given year |
| weight | a named list, with names corresponding to stanza_groups, where each list- element is a matrix with rownames for years and colnames for integer ages, where NA excludes the entry from inclusion and the model computes the log- normal likelihood for weight-at-age in each specified age-year combination |
| PB | numeric-vector with names matching taxa, providing the ratio of production to biomass for each taxon |
| QB | numeric-vector with names matching taxa, providing the ratio of consumption to biomass for each taxon |
| В | numeric-vector with names matching taxa, providing the starting (or fixed) value for equilibrium biomass for each taxon |
| DC | numeric-matrix with rownames and colnames matching taxa, where each col- umn provides the diet proportion for a given predator |
| EE | numeric-vector with names matching taxa, providing the proportion of propor- tion of production that is subsequently modeled (termed ecotrophic efficiency) |
| Х | numeric-matrix with rownames and colnames matching taxa, where each ele- ment gives the vulnerability parameter for a given interaction. |
| type | character-vector with names matching taxa and elements c("auto", "hetero", "detritus"), indicating whether each taxon is a primary producer, consumer/predator, or de- tritus, respectively. |
| U | numeric-vector with names matching taxa, providing the proportion of con- sumption that is unassimilated and therefore exported to detritus |
| fit_B | Character-vector listing taxa for which equilibrium biomass is estimated as a fixed effect |
| fit_Q | Character-vector listing taxa for which the catchability coefficient is estimated as a fixed effect |
| fit_B0 | Character-vector listing taxa for which the ratio of initial to equilibrium biomass is estimated as a fixed effect |
| fit_EE | Character-vector listing taxa for which ecotrophic efficiency is estimated. |
| fit_PB | Character-vector listing taxa for which equilibrium production per biomass is estimated. Note that it is likely a good idea to include a prior for any species for which this is estimated. |

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| fit_QB | Character-vector listing taxa for which equilibrium consumption per biomass is estimated. Note that it is likely a good idea to include a prior for any species for which this is estimated. |
|-----------|--|
| fit_eps | Character-vector listing taxa for which the model should estimate annual process errors in dB/dt |
| fit_nu | Character-vector listing taxa for which the model should estimate annual pro- cess errors in consumption Q_ij |
| log_prior | A user-provided function that takes as input the list of parameters out\$obj\$env\$parList() where out is the output from ecostate(), and returns a numeric vector where the sum is the log-prior probability. For example log_prior = function(p) dnorm(p\$logq_i[1], mean=0, sd=0.1, log=TRUE) specifies a lognormal prior probability for the catchability coefficient for the first taxa with logmean of zero and logsd of 0.1 |
| settings | Output from <pre>stanza_settings()</pre> , used to define age-structured dynamics (called stanza-groups). |
| control | Output from ecostate_control(), used to define user settings. |

Details

All taxa must be included in QB, PB, B, and DC, but additional taxa can be in QB, PB, B, and DC that are not in taxa. So taxa can be used to redefine the set of modeled species without changing other inputs

Value

An object (list) of S3-class ecostate. Elements include:

obj RTMB object from MakeADFun

tmb_inputs The list of inputs passed to MakeADFun

opt The output from nlminb

sdrep The output from sdreport

interal Objects useful for package function, i.e., all arguments passed during the call

rep report file, including matrix B_ti for biomass in each year t and taxon i, g_ti for growth rate per biomass, and see dBdt for other quantities reported by year

derived derived quantity estimates and standard errors, for rep objects as requested

- call function call record
- run_time Total runtime

This S3 class then has functions summary, print, and logLik

References

Introducing the state-space mass-balance model:

Thorson, J. Kristensen, K., Aydin, K., Gaichas, S., Kimmel, D.G., McHuron, E.A., Nielsen, J.N., Townsend, H., Whitehouse, G.A (In press). The benefits of hierarchical ecosystem models: demonstration using a new state-space mass-balance model EcoState. Fish and Fisheries.

Description

Define a list of control parameters.

Usage

```
ecostate_control(
  nlminb_loops = 1,
  newton_loops = 0,
  eval.max = 1000,
  iter.max = 1000,
  getsd = TRUE,
  silent = getOption("ecostate.silent", TRUE),
  trace = getOption("ecostate.trace", 0),
  verbose = getOption("ecostate.verbose", FALSE),
  profile = c("logF_ti", "log_winf_z", "s50_z", "srate_z"),
  random = c("epsilon_ti", "alpha_ti", "nu_ti", "phi_tg2"),
  tmb_par = NULL,
  map = NULL,
  getJointPrecision = FALSE,
  integration_method = c("ABM", "RK4", "ode23", "rk4", "lsoda"),
  process_error = c("epsilon", "alpha"),
  n_{steps} = 10,
  F_type = c("integrated", "averaged"),
  derived_quantities = c("h_g2", "B_ti", "B0_i"),
  scale_solver = c("joint", "simple"),
  inverse_method = c("Standard", "Penrose_moore"),
  tmbad.sparse_hessian_compress = 1,
  start_tau = 0.001
)
```

| nlminb_loops | Integer number of times to call stats::nlminb(). |
|--------------|---|
| newton_loops | Integer number of Newton steps to do after running stats::nlminb(). |
| eval.max | Maximum number of evaluations of the objective function allowed. Passed to control in stats::nlminb(). |
| iter.max | Maximum number of iterations allowed. Passed to control in stats::nlminb(). |
| getsd | Boolean indicating whether to call TMB::sdreport() |
| silent | Disable terminal output for inner optimizer? |
| trace | Parameter values are printed every trace iteration for the outer optimizer. Passed to control in stats::nlminb(). |

| verbose | Output additional messages about model steps during fitting? |
|-------------------------------|--|
| profile | parameters that are profiled across, passed to MakeADFun |
| random | parameters that are treated as random effects, passed to MakeADFun |
| tmb_par | list of parameters for starting values, with shape identical to tiny VAST() $internal$ |
| map | list of mapping values, passed to RTMB::MakeADFun |
| getJointPrecis | ion |
| | whether to get the joint precision matrix. Passed to sdreport. |
| integration_me | thod |
| | What numerical integration method to use. "ABM" uses a native-R versions of Adam-Bashford, "RK4" uses a native-R version of Runge-Kutta-4, and "ode23" uses a native-R version of adaptive Runge-Kutta-23, where all are adapted from pracma functions. "rk4" and lsoda use those methods from deSolve::ode as implemented by RTMBode::ode |
| process_error | Whether to include process error as a continuous rate (i.e., an "innovation" parameterization, process_error="epsilon") or as a discrete difference be- tween expected and predicted biomass (i.e., a "state-space" parameterization), process_error="alpha"The former is more interpretable, whereas the latter is much more computationally efficient. |
| n_steps | number of steps used in the ODE solver for biomass dynamics |
| F_type | whether to integrate catches along with biomass ("integrated") or calculate catches from the Baranov catch equation applied to average biomass ("averaged") |
| derived_quanti | ties |
| | character-vector listing objects to ADREPORT |
| scale_solver | Whether to solve for ecotrophic efficiency EE given biomass B (scale_solver="simple") or solve for a combination of EE and B values |
| inverse_method | whether to use pseudoinverse or standard inverse |
| tmbad.sparse_hessian_compress | |
| | passed to TMB::config(), and enabling an experimental feature to save memory when first computing the inner Hessian matrix. Using tmbad.sparse_hessian_compress=1 seems to have no effect on the MLE (although users should probably confirm this), and hugely reduces memory use in both small and large models. Using tmbad.sparse_hessian_compress=1 seems to hugely speed up the model- fitting with a large model but results in a small decrease in speed for model- fitting with a small model. |
| start_tau | Starting value for the standard deviation of process errors |

Value

An S3 object of class "ecostate_control" that specifies detailed model settings, allowing user specification while also specifying default values

ginv

Description

Extend MASS: ginv to work with RTMB

Usage

ginv(x)

Arguments

х

Matrix used to compute pseudoinverse

logLik.ecostate Marginal log-likelihood

Description

Extract the (marginal) log-likelihood of a ecostate model

Usage

```
## S3 method for class 'ecostate'
logLik(object, ...)
```

Arguments

| object | Output from ecostate |
|--------|----------------------|
| | Not used |

Value

object of class logLik with attributes

| val | log-likelihood |
|-----|----------------------|
| df | number of parameters |

Returns an object of class logLik. This has attributes "df" (degrees of freedom) giving the number of (estimated) fixed effects in the model, abd "val" (value) giving the marginal log-likelihood. This class then allows AIC to work as expected.

ode23

Description

Runge-Kutta (2, 3)-method with variable step size, resp

Usage

ode23(f, a, b, y0, n, Pars, rtol = 0.001, atol = 1e-06)

Plot foodweb

Arguments

| f | function in the differential equation $y' = f(x, y)$; defined as a function $R \times R^m \to R^m$, where m is the number of equations. |
|------|---|
| а | starting time for the interval to integrate |
| b | ending time for the interval to integrate. |
| у0 | starting values at time a |
| n | Not used |
| Pars | named list of parameters passed to f |
| rtol | relative tolerance. |
| atol | absolute tolerance. |
| | |

Details

Copied from pracma under GPL-3, with small modifications to work with RTMB. This can be used to simulate dynamics, but not during estimation

Value

List with components t for time points between a and b and y an n-by-m matrix with solutions for variables in columns, i.e. each row contains one time stamp.

plot_foodweb

Description

Plot consumption as a directed graph including all taxa (vertices) and biomass consumed (arrows). Taxa are located using tracers, where by default the y-axis is trophic level. #'

Usage

```
plot_foodweb(
    Q_ij,
    type_i,
    xtracer_i,
    ytracer_i = rep(1, nrow(Q_ij)),
    B_i = rep(1, nrow(Q_ij)),
    taxa_labels = letters[1:nrow(Q_ij)],
    xloc,
    yloc
)
```

Arguments

| Q_ij | Consumption of each prey i by predator j in units biomass. |
|-------------|--|
| type_i | character vector indicating whether a taxon is "hetero", "auto", or "detritus" |
| xtracer_i | tracer to use when computing x-axis values |
| ytracer_i | tracer to use when computing y-axis values |
| B_i | biomass to use when weighting taxa in plot |
| taxa_labels | character vector of labels to use for each taxon |
| xloc | x-axis location (overrides calculation using xtracer_i) |
| yloc | y-axis location (overrides calculation using ytracer_i) |

Details

Trophic level l_i for each predator *i* is defined as:

 $\mathbf{l-1} = \mathbf{lQ}^*$

where \mathbf{Q}_* is the proportion consumption for each predator (column) of different prey (rows). We identify primary producers as any taxa with no consumption (a column of 0s), and assign them as the first trophic level.

Value

invisibly return ggplot object for foodweb

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print.ecostate Print fitted ecostate object

Description

Prints output from fitted ecostate model

Usage

S3 method for class 'ecostate'
print(x, ...)

Arguments

| х | Output from ecostate |
|---|----------------------|
| | Not used |

Value

No return value, called to provide clean terminal output when calling fitted object in terminal.

print_ecopars Print EcoSim parameters

Description

Prints parameters defining EcoSim dynamics

Usage

print_ecopars(x, silent = FALSE)

Arguments

| х | Output from ecostate |
|--------|------------------------------|
| silent | whether to print to terminal |

Value

invisibly returns table printed

rk4sys

Description

Classical Runge-Kutta of order 4.

Usage

rk4sys(f, a, b, y0, n, Pars, ...)

Arguments

| f | function in the differential equation $y' = f(x, y)$; defined as a function $R \times R^m \to R^m$, where m is the number of equations. |
|------|---|
| а | starting time for the interval to integrate |
| b | ending time for the interval to integrate. |
| y0 | starting values at time a |
| n | the number of steps from a to b. |
| Pars | named list of parameters passed to f |
| | additional inputs to function f |
| | |

Details

Classical Runge-Kutta of order 4 for (systems of) ordinary differential equations with fixed step size. Copied from pracma under GPL-3, with small modifications to work with RTMB

Value

List with components x for grid points between a and b and y an n-by-m matrix with solutions for variables in columns, i.e. each row contains one time stamp.

stanza_settings Detailed control for stanza structure

Description

Define a list of control parameters.

stanza_settings

Usage

```
stanza_settings(
  taxa,
  stanza_groups,
 Κ,
 d,
 Wmat,
 Amax,
 SpawnX,
 Leading,
 fit_K = c(),
 fit_d = c(),
 fit_phi = vector(),
 Amat = NULL,
 Wmatslope,
 STEPS_PER_YEAR = 1,
 comp_weight = c("multinom", "dir", "dirmult")
)
```

| taxa | Character vector of taxa included in model. |
|---------------|---|
| stanza_groups | character-vector with names corresponding to taxa and elements specifying the multi-stanza group (i.e., age-structured population) for a given taxa |
| К | numeric-vector with names matching unique(stanza_groups), providing the von Bertalanffy growth coefficient for length |
| d | numeric-vector with names matching unique(stanza_groups), providing the von Bertalanffy allometric consumption-at-weight (default is 2/3) |
| Wmat | numeric-vector with names matching unique(stanza_groups), providing the weight-at-maturity relative to asymptotic weight |
| Amax | numeric-vector with names matching names(stanza_groups), providing the maximum age (in units years) for a given taxon (and the oldest taxon for a given stanza_group is treated as a plus-group) |
| SpawnX | numeric-vector with names matching unique(stanza_groups), providing the larval vulnerability (density dependence) parameter |
| Leading | Boolean vector with names matching names(stanza_groups), with TRUE for the taxon for which scale (B or EE) is specified or estimated, where this is then calculated deterministically for other taxa for a given stanza_group |
| fit_K | Character-vector listing stanza_groups for which K is estimated |
| fit_d | Character-vector listing stanza_groups for which d is estimated (note that this currently does not work) |
| fit_phi | Character-vector listing stanza_groups for which the model should estimate annual recruitment deviations, representing nonconsumptive variation in larval survival (e.g., oceanographic advection) |

| Amat | numeric-vector with names matching unique(stanza_groups), providing the integer age-at-maturity (in units years) |
|----------------|---|
| Wmatslope | numeric-vector with names matching unique(stanza_groups), providing the slope at 0.5 maturity for a logistic maturity-at-weight ogive |
| STEPS_PER_YEAR | integer number of Euler steps per year for calculating integrating individual weight-at-age |
| comp_weight | method used for weighting age-composition data |

Value

An S3 object of class "stanza_settings" that specifies detailed model settings related to age-structured dynamics (e.g., stanzas), allowing user specification while also specifying default values

whitehouse_2021 Full rpath inputs for eastern Bering Sea

Description

All Rpath inputs from Whitehouse et al. 2021

Usage

data(whitehouse_2021)

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