# **R** documentation

# of all in 'PBSawatea'

# July 11, 2011

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allEqual

Are All Values Equal to the First?

# Description

A short-cut function for all (x==x[1]), which asks are all values in x equl to the first value in x

# Usage

Index

```
allEqual(x)
```

# Arguments

X

vector of values.

# Value

TRUE or FALSE

```
all, clearAll, clipVector
```

calc.projExpect 3

```
calc.projExpect Calculate Expectations and Probabilities
```

### **Description**

Calculate the expectation of projection to reference, and probability of being greater than reference.

#### Usage

```
calc.projExpect ( obj, projObj, refYrs )
calc.projExpect2( obj, projObj, refList )
calc.projProbs ( obj, projObj, refYrs )
calc.projProbs2 ( obj, projObj, refList )
calc.refProbs ( projObj=currentProj$B, refPlist=refPointsList )
```

### **Arguments**

```
obj matrix of biomass MCMCs.

projObj matrix of biomass projections.

refYrs numeric vector of reference years

refList list of reference years (numeric vectors).

refPList list of reference points.
```

#### **Details**

#### Value

Decision tables

### Author(s)

Andrew Edwards, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC

```
calc.refVal
```

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calc.refVal

Calculate Reference Value for Performance Measure

# Description

Calculate the reference value for performance measures.

### Usage

```
calc.refVal(obj, refYrs, fun=mean)
```

### **Arguments**

obj scape Biomass matrix with n rows and m columns,

where n = number of MCMC samples, and m = number of years.

refYrs numeric years in reference period.

fun the function to apply to reference period i.

### Value

Returns a vector of length nrow (obj) reference values.

### Author(s)

Andrew Edwards, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC

### See Also

```
calc.projExpect, findTarget
```

close.allWin

Close All Open Devices

# Description

Close all open devices.

# Usage

```
close.allWin()
```

# See Also

closeWin

cquantile 5

cquantile

Running Quantile

# Description

Creates a set of running quantiles from MCMC traces. (Uses subfunction found in **coda**'s function cumuplot.)

# Usage

```
cquantile(z, probs)
cquantile.vec(z, prob)
```

# Arguments

```
z an MCMC object.
probs vector of quantiles.
prob single quantile.
```

# Value

```
cquantile.....running quantile matrix cquantile.vec...running quantile vector
```

### Note

Arni Magnusson describes a running quantile as: "the evolution of the sample quantiles as a function of the number of iterations"

# Author(s)

Andrew Edwards, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC

```
cumuplot, plotTracePOP
```

6 findTarget

findTarget

Find Time to Achieve a Target Reference Point

### **Description**

Find the time (years) to achieve a recovery target (including a moving target) with a given confidence. Produce decision tables showing the probability of exceeding the reference point.

### Usage

```
findTarget(Vmat, yrU=as.numeric(dimnames(Vmat)[[2]]), yrG=90,
    ratio=0.5, target=B0.MCMC, conf=0.95, plotit=FALSE, retVal="N")
```

### **Arguments**

Vmat	matrix of projected biomass values $B_{Nt}$ , where $N =$ number of MCMCs and $t =$ projection year.
yrU	user-specified projection years.
yrG	number of years $G$ for a moving target window (e.g., 3 YMR generations = 90y); might not work for all possibilities.
ratio	recovery target ratio $R$ .
target	recovery target values $T_N$ = B0 . MCMC for ratios of $B_0$ ; = Bmsy . MCMC for ratios of $B_{MSY}$ ; = Bt . MCMC for moving window of $B_{N,t-G}$ .
conf	confidence level $C$ required.
plotit	logical: if TRUE, plot the probability $p_t$ of exceeding target reference point.
retVal	<pre>character name of object to return: retVal="N" : creates global object "Ttab" (see below); retVal="p.hi" : creates global object "Ptab" (see below).</pre>

### **Details**

As this function uses Bayesian output, there are N (e.g., 1000) values of some target  $T_N$ , which can remain fixed  $(B_0, B_{MSY})$  or move forward in time G years before the projection year t (that is  $T_{N,t-G}$ ). For simplification, we'll just call all targets  $T_N$ .

The probability of exceeding a target ratio R is:

$$p_t = \frac{1}{N} \sum_{t=0}^{N} \left[ \frac{B_{Nt}}{T_N} \ge R \right],$$

where R = target ratio of the reference point (e.g.,  $0.4B_{MSY}$  (R=0.4),  $0.2B_0$  (R=0.2),  $0.5B_{t-G}$  (R=0.5)).

get.resFile 7

At a glance, we can see for any given projection year t whether the probability of achieving a target ratio is greater than the confidence required:

$$p_t \geq C$$
,

where C is the confidence level acceptable.

#### Value

If retVal="N" then the function returns a data frame object called "Ttab" in the user's global environment. This table reports the number of years to achieve the target reference point at various catch levels with a specified confidence.

If retVal="p.hi" then the function returns a list object called "Ptab" in the user's global environment. This list contains data frames (tables) that report the probability of achieving various reference points at specified catch levels.

Any other retVal will return a list of the specified object, if it exists in the function.

### Author(s)

Rowan Haigh, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC

#### See Also

runSweaveMCMC

get.resFile

Get Awatea Results Files for Menu

# Description

A function that retrieves the names of Awatea results files (.res\$) for use in the mainMenu command. When choice is made, the function loads the results file and assigns it to the global environment as currentRes.

#### Usage

```
get.resFile(resFile=NULL)
```

# **Arguments**

resFile

supposedly the name of a results file, but the code suggests that argument is ignored.

#### Value

A results file chosen from a menu.

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### Note

AME: made changes so that options are compatible with those in load.allResFiles. Previously, trouble occurred when overwriting.

#### See Also

```
mainMenu,importCol2
```

getYrIdx

Select a Subset of Years for Plotting

# Description

Select a subset of years for which many years are available. The default is to select 5-year increments.

### Usage

```
getYrIdx(yrNames, mod=5)
```

### **Arguments**

yrNames vector (character or numeric) of years.

mod select the years modulo mod.

### Value

Subset of input years that are modulo mod.

# Author(s)

Andrew Edwards, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC

```
findPat, pad0
```

graphics 9

graphics	Open a Windows Device
91400	open a mina ma zerree

### **Description**

Open a windows device in portrait or landscape mode.

# Usage

```
graphics(view = "portrait")
```

# **Arguments**

```
view if "portrait", set width = 8.5 in and height = 11 in if "landscape", set width = 11 in and height = 8.5 in
```

### See Also

```
resetGraph
```

importCol2 Import Coleraine Model Results (AME version)	
---	--

# Description

Import Coleraine model results from .res file, and rearrange into a standard format suitable for plotting.

### Usage

res.file	name of Coleraine model results file to import.
info	optional string containing information to store with model results.
Dev	logical: whether recruitment deviates were estimated in model.
CPUE	logical: whether model was fitted to catch-per-unit-effort data.
Survey	logical: whether model was fitted to survey abundance index data.
CAc	logical: whether model was fitted to commercial catch-at-age data.
CAs	logical: whether model was fitted to survey catch-at-age data.
CLc	logical: whether model was fitted to commercial catch-at-length data.

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CLs	logical: whether model was fitted to survey catch-at-length data.
LA	logical: whether model was fitted to length-at-age data.
quiet	logical: whether to report progress while parsing file.
extra	logical: if TRUE, import likelihoods, parameters, priors, and recruitment residuals.

### **Details**

This function was modified from the original importCol function in the **scape** package to grab extra data.

#### Value

A list of class scape containing at least N, B, and Sel. The other elements may or may not be included in the list, depending on how importRes was called:

N	predicted numbers at age
В	predicted biomass, recruitment, and observed landings (year things)
Sel	predicted selectivity and observed maturity (age things)
Dev	predicted recruitment deviates from the stock-recruitment curve
CPUE, Survey	commercial and survey abundance index and fit
CAc, CAs	commercial and survey C@A (catch at age) and fit
CLc, CLs	commercial and survey C@L (catch at length) and fit
LA	observed L@A and fit

### Note

This import function is implemented for the Coleraine statistical catch-at-age software, and can serve as a template for **scape** users who would like to implement import functions for specific stock assessment software.

```
The functions 11 (package gdata) and head are recommended for browsing model results, e.g. 11 (x.cod); 11 (x.cod\$N); head (x.cod\$N).
```

### References

Hilborn, R., Maunder, M., Parma, A., Ernst, B., Payne, J., and Starr, P. (2003) Coleraine: A generalized age-structured stock assessment model. User's manual version 2.0. *University of Washington Report* **SAFS-UW-0116**. Available at:

```
http://fish.washington.edu/research/coleraine/pdf/coleraine.pdf.
```

```
importRes, read.table, readLines, and scan to import any data. scape-package gives an overview of the package scape.
```

importMCMC.ddiff 11

```
importMCMC.ddiff Import Functions for PJS Delay Difference Model
```

### **Description**

Make a **scapeMCMC** object identical in format to the result of importMCMC (or importProj) from PJS delay difference model output.

The difference is that B is biomass defined by a delay difference model.

### Usage

```
importMCMC.ddiff()
importProj.ddiff()
```

### Value

```
importMCMC.ddiff returns a list object containing:
L...likelihood MCMCs,
P...parameter MCMCs,
B...spawning biomass MCMCs,
R...recruitment MCMCs.
importProj.ddiff returns a list object containing:
B...projected biomass,
Y...projected yields.
```

### Note

```
Get the biomass projection – PJS does 1 year ahead projection. The column "X" appears as the last column because trailing ", " exist in the mcmcprojbiom.csv file.

Note also that "cat=" in .csv file becomes "cat." in read.table.
```

### Author(s)

Andrew Edwards, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC

```
importMCMC, importProj, read.table
```

12 importProjRec

importProjRec	Import Projected Recruitment

### **Description**

Import the projected recruitments (actually, the values are random normals N(0,1)).

### Usage

```
importProjRec(dir, info="", coda=FALSE, quiet=TRUE)
```

### **Arguments**

dir	directory where MCMC projections reside.
info	user-supplied information, if desired.
coda	logical: if TRUE, use the function $memc$ in the package $coda$ to generate MCMCs.
quiet	logical: if TRUE, print progress messages to the R console.

### **Details**

The values saved by the Awatea code are random normals N(0,1), which for a particular MCMC sample are the same for all the catch strategies.

### Value

A list object comprising:

```
B data frame of spawning biomass (dim = MCMC samples by projected years)
Y data frame of yield (dim = MCMC samples by projected years)
eps data frame of \epsilon_t (dim = MCMC samples by projected years)
```

### Note

The function import Proj does not import recruitment residuals.

This function grabs the tempdev values from Awatea, which are just N(0,1) values, then multiplies them by  $\sigma_R$  to yield  $\epsilon_t \sim N(0,\sigma_R^2)$ .

The parameter value for  $\sigma_R$  can be found in currentRes\$extra\$residuals\$p\_log\_RecDev[6].

### Author(s)

Andrew Edwards, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC

```
importProj
coda: mcmc
```

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importRes	Import Coleraine Model Results (RH version)	
-----------	---	--

### **Description**

Import Coleraine model results from .res file, and rearrange into a standard format suitable for plotting.

# Usage

### **Arguments**

res.file	name of Coleraine model results file to import.
info	optional string containing information to store with model results.
Dev	logical: whether recruitment deviates were estimated in model.
CPUE	logical: whether model was fitted to catch-per-unit-effort data.
Survey	logical: whether model was fitted to survey abundance index data.
CAc	logical: whether model was fitted to commercial catch-at-age data.
CAs	logical: whether model was fitted to survey catch-at-age data.
CLc	logical: whether model was fitted to commercial catch-at-length data.
CLs	logical: whether model was fitted to survey catch-at-length data.
LA	logical: whether model was fitted to length-at-age data.
quiet	logical: whether to report progress while parsing file.
extra	logical: if $\ensuremath{\mathtt{TRUE}}$ , import likelihoods, parameters, priors, and recruitment residuals.
sep	the field separator character (usually " " or "\t").

### **Details**

This function was modified from the original importCol function in the **scape** package to grab extra data and to deal with anomalous characters in Coleraine results files.

### Value

A list of class list containing at least N, B, and Sel. The other elements may or may not be included in the list, depending on how importRes was called:

N	predicted numbers at age
В	predicted biomass, recruitment, and observed landings (year things)

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Sel	predicted selectivity and observed maturity (age things)
Dev	predicted recruitment deviates from the stock-recruitment curve
CPUE, Survey	commercial and survey abundance index and fit
CAc, CAs	commercial and survey C@A (catch at age) and fit
CLc, CLs	commercial and survey C@L (catch at length) and fit
LA	observed L@A and fit

### Note

This import function is implemented for the Coleraine statistical catch-at-age software, and can serve as a template for **scape** users who would like to implement import functions for specific stock assessment software.

```
The functions 11 (package gdata) and head are recommended for browsing model results, e.g. 11 (x.cod); 11 (x.cod\$N); head (x.cod\$N).
```

### References

Hilborn, R., Maunder, M., Parma, A., Ernst, B., Payne, J., and Starr, P. (2003) Coleraine: A generalized age-structured stock assessment model. User's manual version 2.0. *University of Washington Report* **SAFS-UW-0116**. Available at:

http://fish.washington.edu/research/coleraine/pdf/coleraine.pdf.

### See Also

```
runADMB, readAD, reweight, importCol2 read.table, readLines, and scan to import any data. scape-package gives an overview of the package scape.
```

```
load.allResFiles Load All Awatea .res Files
```

### Description

Load all Awatea . res files in the working directory into a list object.

### Usage

```
load.allResFiles(resList = NULL)
```

### **Arguments**

resList AME: sets directory to path above current and sets the pattern to "results.dat\$"; probably deprecated.

### Value

List of multiple calls to importCol2.

MAfun2

### Note

If deprecated, remove from package **PBSawatea**.

### Author(s)

Andrew Edwards, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC

#### See Also

```
importCol2
```

MAfun2

Calculate Mean Age by Year

### **Description**

Calculate mean ages from proportions-at-age (modified from a subfunction in runADMB).

### Usage

```
MAfun2 (padata, brks=NULL)
```

### **Arguments**

padata proportion-at-age data CAc or CAs from a call to importCol2.

brks breaks specified as numeric years to split the commercial data up into regimes

that may account for index discontinuities (not used).

# **Details**

Mean age function supplied by Chris Francis (2011).

padata has fields:

Series...series identifier Year....numeric year Age.....age bin

Obs....observed proportions

Fit.....predicted (fitted) proportions SS.....sample size (effective N)

### Value

List object of observed and expected mean ages, variance of expected ages, and a few bits and bobs.

### Author(s)

Rowan Haigh, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC.

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### References

Francis, R.I.C.C. (2011, in press) Data weighting in statistical fisheries stock assessment models. *Canadian Journal of Fisheries and Aquatic Sciences*.

### See Also

```
runADMB, importCol2
```

mainMenu

Create a Menu of Options and Actions

### **Description**

From a main menu, the user can choose various options and actions.

We tend to do everything from the command line so the menu functionality fosters the warning: *CAVEAT EMPTOR*.

### Usage

```
mainMenu()
loadMenu()
mpdMenu()
mcmcMenu()
utilMenu()
```

### **Details**

### Main menu items:

```
Import files
MPD plots
Plot all MPD graphs
Save all MPD plots to PNG
MCMC plots
Plot all MCMC plots
Save all MCMC plots to PNG
Close all graphics windows
Help & Utilities
```

### Load menu items:

```
Get Awatea res file
Get Awatea MCMC file
Get Awatea projection file
Load all res files in working directory
Get PJS Delay Difference MCMC+Projection
```

#### **MPD** menu items:

```
Plot biomass, recruitment, catch Plot numbers at age
```

makeErrMat 17

```
Plot selectivity and maturity
Plot commercial catch—at—age results
Plot survey catch—at—length results
Plot survey catch—at—length results
Plot abundance index
All residual plots
Plot multi—panel biomass, recruitment, catch
Plot multi—panel exploitation rate
Plot alternative numbers at age
```

#### MCMC menu items:

Plot biomass and projections by policy
Probability of projection biomass > reference
Expectation of projection biomass / reference
Plot biomass posterior densities (plotDens)
Plot recruitment posterior densities (plotDens)
Plot parameter posterior densities (plotDens)
Plot cumulative quantiles (plotCumu)
Plot traces (plotTrace)
Plot PJS traces (plt.allTraces)

#### **Utils menu items:**

scape Help
scapeMCMC Help
Portrait graphsheet
Landscape graphsheet

### See Also

```
get.resFile,importCol2
```

makeErrMat

Make Ageing Error Matrix for Awatea

### **Description**

Make a simple ageing error matrix for Awatea.

### Usage

```
makeErrMat(N=60, ondiag=0.8, offdiag=0.1, corner=0.9)
```

N	numeric scalar indicating number of age classes, which determines the dimension of the matrix.
ondiag	numeric value to appear along the matrix diagonal.
offdiag	numeric value to appear one cell to the left and right of the matrix diagonal.
corner	numeric value to appear in the top left and bottom right corners of the matrix.

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### Value

Simple symmetric ageing error matrix.

### Author(s)

Rowan Haigh, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC

#### See Also

```
plotProp, weightBio
```

msyCalc

Calculate the Maximum Sustainable Yield

## **Description**

Load in MSY.out and calculate the MSY (maximum sustainable yield).

## Usage

```
msyCalc(dir = getwd(), error.rep = 1)
```

### **Arguments**

```
dir working directory.
error.rep numeric: if 1, report errors (reaching bounds), if 0 do not.
```

### Value

```
Returns a list object containing: yield...maximum sustainable yield,
```

u.....exploitation rate at MSY, VB.....vulnerable biomass at MSY,

B..... spawning biomass at MSY,

nProj...numnber of projections needed to reach MSY.

### Note

See msyTestCreating.r for full details when figuring this out.

### Author(s)

Andrew Edwards, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC

```
findTarget
```

out.pmTables 19

out.pmTables

Write Decision Tables to Comma-Delimited Files

# Description

Write decision tables to comma-delimited text files (.csv).

## Usage

```
out.pmTables(obj, fileName="pm", dec=3)
```

### **Arguments**

obj list object containing tables (matrices or data frames).

fileName prefix for output file names.

dec number of decimal places to retain.

#### Value

Comma-delimited text files (.csv).

#### Author(s)

Andrew Edwards, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC

### See Also

```
write.table, writeList
```

panLab

Write Text to Figure in Relative (0:1) Coordinates

# Description

```
Write text to a figure by first setting the coordinate space to lie between 0 and 1: par(usr=c(0,1,0,1)).
```

# Usage

```
panLab(x, y, txt, ...)
```

### **Arguments**

x relative x-coordinate.

y relative y-coordinate.

txt text to add to figure.

 $\dots$  additional arguments sent to function text.

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### Note

Currently, this function does not reset the coordinate space to the original. Use addLabel instead.

### See Also

```
addLabel, addLegend
```

panLegend

Place a Legend in a Figure using Relative (0:1) Coordinates

# Description

```
Place a legend in a figure by first setting the coordinate space to lie between 0 and 1: par(usr=c(0,1,0,1)).
```

# Usage

```
panLegend(x, y, legTxt, ...)
```

# Arguments

```
    x relative x-coordinate.
    y relative y-coordinate.
    legTxt legend text to add to figure.
    additional arguments sent to function legend.
```

### Note

Currently, this function does not reset the coordinate space to the original. Use addLegend instead.

```
addLabel, addLegend
```

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PBSawatea

PBS Awatea

### Description

**PBSawatea** contains the code used for the modelling of populations of Pacific Ocean Perach (*Sebastes alutus*) and Yellowmouth Rockfish (*S. reedi*) along the British Columbia (BC) coast.

Implementation is done using a modified version of the Coleraine statistical catch-at-age software (Hilborn *et al.* 2003) called Awatea (Alan Hicks, NOAA, pers. comm.). Awatea is a platform for implementing the AD (Automatic Differentiation) Model Builder software (Otter Research 1999), which provides (a) maximum posterior density estimates using a function minimiser and automatic differentiation, and (b) an approximation of the posterior distribution of the parameters using the Markov Chain Monte Carlo (MCMC) method, specifically using the Hastings-Metropolis algorithm (Gelman *et al.* 2004).

Running of Awatea is streamlined using code written in R (R Development Core Team 2009), rather than the original Microsoft Excel implementation. Figures and tables of output are automatically produced through R using code adapted from the R packages **scape** (Magnusson 2009) and **scapeMCMC** (Magnusson and Stewart 2007). We use the R function Sweave (Leisch 2008) in the package **utils** to automatically collate, via LATEX, the large amount of figures and tables into a single portable document file (.pdf) for each model run.

We provide master Sweave files used in folder .../library/PBSawatea/snw to build the .pdf document. The user must copy these to a local working directory if they are not already there.

#### References

Gelman, A., Carlin, J.B., Stern, H.S, and Rubin, D.B. (2004) Bayesian data analysis, 2nd edition. Chapman and Hall/CRC, New York, 668 p.

Hilborn, R., Maunder, M., Parma, A., Ernst, B. Payne, J., and Starr, P. (2003) Coleraine: a generalized age-structured stock assessment model. *School of Aquatic and Fishery Sciences*, University of Washington, 54 p.

Leisch, F. (2008) Sweave, R package.

Magnusson, A. (2009) Scape – statistical catch-at-age plotting environment, R package.

Magnusson, A. and Stewart, I. (2007) MCMCscape – MCMC diagnostic plots. R package.

Otter Research Ltd. (1999) An introduction to AD Model Builder for use nonlinear modeling and statistics. Otter Research Ltd., British Columbia. 194 p.

R Development Core Team (2011) R: A language and environment for statistical computing. R Foundation for Statistical Computing, Vienna, Austria.

ISBN 3-900051-07-0

http://www.R-project.org

plotB2

plotB2	Plot Biomass, Recruitment, and Landings (AME Version)

### **Description**

Plot scape model predicted biomass, stock recruitment, and landings.

AME: This is an alteration of Arni Magnussons plotB function to accommodate PJS's request not to show biomass prior to fishery and survey indices period.

# Usage

```
plotB2 (model, what="d", series=NULL, years=NULL, axes=TRUE, div=1,
    legend="bottom", main="", xlab="", ylab="", cex.main=1.2,
    cex.legend=1, cex.lab=1, cex.axis=0.8, las=1,
    tck=c(1,what=="d")/2, tick.number=5, lty.grid=3, col.grid="white",
    pch=16, cex.points=0.8, col.points="black", lty.lines=1:3,
    lwd.lines=2, col.lines="black", ratio.bars=3, col.bars="grey",
    plot=TRUE, ...)
```

model	fitted scape model.
what	what to plot: "d"[efault], "s"[tock recruitment], or "l"[andings].
series	vector of strings indicating which column names in model\$B data frame to plot (all by default).
years	vector of numbers indicating which years to include (all by default).
axes	whether to plot axis values.
div	denominator to shorten values on the y axis, or a vector with two elements referring to x and y axis.
legend	<pre>legend location: "bottom", "left", "top", "right", or "" to suppress legend.</pre>
main	main title.
xlab	x-axis label.
ylab	y-axis label.
cex.main	size of main title.
cex.legend	size of legend text.
cex.lab	size of axis labels.
cex.axis	size of tick labels.
las	orientation of tick labels: 0=parallel, 1=horizontal, 2=perpendicular, 3=vertical.
tck	tick mark length.
tick.number	number of tick marks.

plotB2

```
line type of gridlines.
lty.grid
col.grid
                 color of gridlines.
                 symbol for points.
pch
cex.points
                 size of points.
col.points
                 color of points.
lty.lines
                 line type of main lines.
lwd.lines
                 line width of main lines.
col.lines
                 color of main lines.
ratio.bars
                 width of bars.
col.bars
                 color of bars.
                 whether to draw plot.
plot
                 passed to xyplot and panel.superpose.
```

### **Details**

The "d"[efault] plot shows spawning biomass and vulnerable biomass as lines, and landings as bars, on the same scale.

### Value

When plot=TRUE, a trellis plot is drawn and a data frame is returned, containing the data used for plotting. When plot=FALSE, a trellis object is returned.

### Note

The Args function from the **gdata** package is recommended for reviewing the arguments, instead of args.

```
xyplot, panel.barchart, and panel.superpose are the underlying drawing functions.
plotCA, plotCL, plotIndex, plotIndex2 and plotLA plot model fit and data.
plotB, plotN, and plotSel plot derived quantities.
scape-package gives an overview of the scapeMCMC package.
```

24 plotBmcmcPOP

plotBmcmcPOP Plot Spa	wning and Vulnerable Biomass
-----------------------	------------------------------

### **Description**

Plot spawning and vulnerable biomass from posterior as boxplots, and add catch bars on same graph.

### Usage

```
MCMC data frame of B (currentMCMC$B).
obj
                 list/scape object created by importCol2.
currentRes1
                 quantiles to use in quantBox.
хуТуре
                 type of plot (currently only uses quantBox).
                 line types to use in quantBox.
lineType
refLines
                 reference lines to add to plot.
xLim
                 limits of the x-axis.
                 limits of the y-axis.
yLim
                 not used
userPrompt
                 not used
save
xLab
                 x-coordinates for labels.
                 y-coordinates for labels.
yLab
                 text labels to display on plot.
textLab
                 increments along the y-axis to place tick marks.
yaxis.by
tcl.val
                 tick length.
                 additional arguments passed to the function rect.
```

plotChains 25

### Note

Combines ideas from plt.quantBio and plotB2. Don't need lattice, just one figure, no panels. Vulnerable biomass has no posterior saved, which must be why it's not been done before. Hmmm... still worth seeing spawning though?

```
Taking what is needed from plt.quantBio, this basically works: plt.quantBio(currentMCMC$B, xyType=rpType), though it creates 2x3 plots. The object should be the specific MCMC posterior by year (so just a data frame), e.g., currentMCMC$B. currentRes1 is local currentRes.
```

### Author(s)

Andrew Edwards, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC

#### See Also

```
plotVBcatch, plotB2
```

plotChains

Plot Cumulative Frequency of MCMC Chains

# **Description**

Plot cumulative fequency of n chains by partitioning one trace. (Modified from the function plotTracePOP.)

### Usage

```
plotChains(mcmc, nchains=3, pdisc=0.1, axes=FALSE, same.limits=FALSE,
    between=list(x=axes,y=axes), div=1, span=1/4, log=FALSE,
    base=10, main=NULL, xlab=NULL, ylab=NULL, cex.main=1.2,
    cex.lab=1, cex.strip=0.8, cex.axis=0.8,
    las=0, tck=0.5, tick.number=5, lty.trace=1, lwd.trace=1,
    col.trace="grey", lty.median=1, lwd.median=1,
    col.median="black", lty.quant=2, lwd.quant=1,
    col.quant="black", plot=TRUE, probs=c(0.025, 0.5, 0.975), ...)
```

mcmc	MCMC chain(s) as a vector, data frame or mcmc object.
nchains	number of chains to create from one trace.
pdisc	proportion of the initial trace to discard before creating chains.
axes	whether axis values should be plotted.
same.limits	whether panels should have same x-axis limits.
between	list with $x$ and $y$ indicating panel spacing.

26 plotChains

denominator to shorten values on the y axis. div smoothness parameter (not used). span whether values should be log-transformed. log logarithm base. base main main title. xlab x-axis title. ylab y-axis title. cex.main size of main title. cex.lab size of axis labels. cex.strip size of strip labels. size of tick labels. cex.axis orientation of tick labels: 0=parallel, 1=horizontal, 2=perpendicular, 3=vertical. las tick mark length. tck tick.number number of tick marks. lty.trace line type of trace. lwd.trace line width of trace. colour of trace. col.trace lty.median line type of median. lwd.median line width of median. colour of median. col.median line type of quantile trace. lty.quant lwd.quant line width of quantile trace. colour of quantile trace. col.quant whether to draw plot. plot quantile values for quantile trace. probs passed to panel.trace (not used). . . .

### Value

When plot=TRUE, a trellis plot is drawn and a data frame is returned, containing the data used for plotting. When plot=FALSE, a trellis object is returned.

### Note

This idea stemmed from a discussion with PJS.

#### Author(s)

Rowan Haigh, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC

```
plotTracePOP, plotDensPOP
```

plotCPUE 27

plotCPUE	Plot CPUE and Add Error Bars

# Description

Plot CPUE and fit with error bars.

# Usage

```
plotCPUE(obj, main="", save=NULL, bar=1.96, yLim=NULL, ...)
```

# Arguments

	not used
yLim	limits of the y-axis.
bar	standard deviation of the normal distribution (1.96 is the approximate value of the 97.5 percentile point).
save	not used
main	title for figure
obj	data frame of CPUE indices from Awatea's results file (e.g., currentRes\$CPUE).

# Value

### A postscript file:

```
CPUEser.eps...CPUE indices with error bars.
```

### Note

Copied code from plotIndexNotLattice.

# Author(s)

Andrew Edwards, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC

```
plotIndexNotLattice
```

28 plotDensPOP

plotDensPOP

Plot MCMC Density (AME Version)

### **Description**

Plot Markov-chain Monte Carlo density. This is an approximation of the posterior probability density function.

### Usage

```
plotDensPOP(mcmc, probs=c(0.025,0.975), points=FALSE, axes=TRUE,
     same.limits=FALSE, between=list(x=axes,y=axes), div=1,
     log=FALSE, base=10, main=NULL, xlab=NULL, ylab=NULL,
     cex.main=1.2, cex.lab=1, cex.strip=0.8, cex.axis=0.7,
     las=0, tck=0.5, tick.number=5,
     lty.density=1, lwd.density=3, col.density="black",
     lty.median=2, lwd.median=1, col.median="darkgrey",
     lty.outer=3, lwd.outer=1, col.outer="darkgrey", pch="|",
     cex.points=1, col.points="black", plot=TRUE, MPD.height=0.04,...)
plotDensPOPpars (mcmc, probs=c(0.025,0.975), points=FALSE,
     axes=TRUE, same.limits=FALSE, between=list(x=axes,y=axes),
     div=1, log=FALSE, base=10, main=NULL, xlab=NULL, ylab=NULL,
     cex.main=1.2, cex.lab=1, cex.strip=0.8, cex.axis=0.7,
     las=0, tck=0.5, tick.number=5,
     lty.density=1, lwd.density=3, col.density="black",
     lty.median=2, lwd.median=1, col.median="darkgrey",
     lty.outer=3, lwd.outer=1, col.outer="darkgrey", pch="|",
     cex.points=1, col.points="black", plot=TRUE, MPD.height=0.04,...)
plotDensPOPparsPrior(mcmc, probs=c(0.025,0.975), points=FALSE,
     axes=TRUE, same.limits=FALSE, between=list(x=axes,y=axes),
     div=1, log=FALSE, base=10, main=NULL, xlab=NULL, ylab=NULL,
     cex.main=1.2, cex.lab=1, cex.strip=0.8, cex.axis=0.7,
     las=0, tck=0.5, tick.number=5,
     lty.density=1, lwd.density=3, col.density="black",
     lty.median=2, lwd.median=1, col.median="darkgrey",
     lty.outer=3, lwd.outer=1, col.outer="darkgrey", pch="|",
     cex.points=1, col.points="black", plot=TRUE, MPD.height=0.04,...)
```

mcmc	MCMC chain(s) as a vector, data frame or mcmc object.
probs	vector of outer quantiles to draw, besides the median.
points	whether data points should be plotted along the x axis.
axes	whether axis values should be plotted.

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```
whether panels should have same x-axis limits.
same.limits
                 list with x and y indicating panel spacing.
between
                  denominator to shorten values on the x axis.
div
                  whether values should be log-transformed.
log
                 logarithm base.
base
                 main title.
main
                 x-axis label.
xlab
ylab
                 y-axis label.
cex.main
                 size of main title.
                 size of axis labels.
cex.lab
cex.strip
                 size of strip labels.
cex.axis
                 size of tick labels.
                 orientation of tick labels: 0=parallel, 1=horizontal, 2=perpendicular, 3=vertical.
las
                 tick mark length.
tck
tick.number
                 number of tick marks.
lty.density
                 line type of density curve.
lwd.density
                 line width of density curve.
                 colour of density curve.
col.density
lty.median
                 line type of median.
lwd.median
                 line width of median.
col.median
                 colour of median.
lty.outer
                 line type of outer quantiles.
lwd.outer
                 line width of outer quantiles.
col.outer
                 colour of outer quantiles.
                 symbol for data points.
pch
cex.points
                 size of data points.
col.points
                 colour of data points.
                 whether to draw plot.
plot
                 how far up to put MPD.
MPD.height
                 passed to densityplot and panel.densityplot.
. . .
```

### **Details**

The function plotDensPOPpars differs from plotDensPOP only by a few tweaks to the internal list object myscales.

The function plotDensPOPparsPrior adds the priors automatically.

#### Value

When plot=TRUE, a trellis plot is drawn and a data frame is returned, containing the data used for plotting. When plot=FALSE, a trellis object is returned.

30 plotIndex2

#### Note

The Args function from the gdata package is recommended for reviewing the arguments, instead of args.

AME: edited plotDens function to have less whitesapce, not repeat x-axis labels, and make yaxes the same scales. Cannot just do it through the options.

For Recruits and Biomass, use plotDensPOPpars for parameters. Tried y-axes the same scales, but 1973-1975 are so narrow that they make all the others really small: same.limits=TRUE, ylim=c(0, 0.0005).

### See Also

xyplot and panel.densityplot are the underlying drawing functions, and densplot is a similar non-trellis plot.

```
plotTrace, plotAuto, plotCumu, and plotSplom are diagnostic plots.
```

```
plotDens and plotQuant are posterior plots.
```

scapeMCMC-package gives an overview of the package.

plotIndex2

Plot Abundance Index (AME Version)

### Description

Plot scape model fit to abundance index data.

Revised version of Arni's function to confine plotting to data region.

### Usage

```
plotIndex2(model, what="c", series=NULL, axes=TRUE, same.limits=FALSE,
          between=list(x=axes,y=axes), ylim=NULL, q=1, bar=1, log=FALSE,
          base=10, main="", xlab="", ylab="", cex.main=1.2, cex.lab=1,
          cex.strip=0.8, cex.axis=0.8, las=1, tck=c(1,0)/2,
          tick.number=5, lty.grid=3, col.grid="white", pch=16,
          cex.points=1.2, col.points="black", lty.lines=1, lwd.lines=4,
          col.lines="dimgrey", lty.bar=1, plot=TRUE, ...)
```

model	fitted scape model containing element CPUE and/or Survey.
what	what to plot: "c"[ommercial] or "s"[urvey] abundance index.
series	vector of strings indicating which gears or surveys to plot (all by default).
axes	whether to plot axis values.
same.limits	whether panels should have same y-axis limits.
between	list with $x$ and $y$ indicating panel spacing.

plotIndex2 31

ylim	vector with lower and upper y-axis limits.
q	denominator to scale the y axis, e.g. to vulnerable biomass. Similar to the div
	argument in plotN and plotB.
bar	extent of error bars relative to standard error.
log	whether to log-transform values.
base	logarithm base.
main	main title.
xlab	x-axis label.
ylab	y-axis label.
cex.main	size of main title.
cex.lab	size of axis labels.
cex.strip	size of strip labels.
cex.axis	size of tick labels.
las	orientation of tick labels: 0=parallel, 1=horizontal, 2=perpendicular, 3=vertical.
tck	tick mark length.
tick.number	number of tick marks.
lty.grid	line type of gridlines.
col.grid	color of gridlines.
pch	symbol for points.
cex.points	size of points.
col.points	color of points and error bars.
lty.lines	line type of main lines.
lwd.lines	line width of main lines.
col.lines	color of main lines.
lty.bar	line type of error bars.
plot	whether to draw plot.
	<pre>passed to xyplot, panel.xyplot, and panel.xYplot.</pre>

### Value

When plot=TRUE, a trellis plot is drawn and a data frame is returned, containing the data used for plotting. When plot=FALSE, a trellis object is returned.

# Note

The Args function from the **gdata** package is recommended for reviewing the arguments, instead of args.

```
xyplot, panel.xyplot, and panel.xYplot are the underlying drawing functions.
plotCA, plotCL, plotIndex, and plotLA plot model fit and data.
plotB, plotB2, plotN, and plotSel plot derived quantities.
scape-package gives an overview of the package.
```

32 plotIndexNotLattice

```
plotIndexNotLattice

Plot Survey Indices
```

# Description

Plot index series with error bars. Create postscript files automatically.

# Usage

```
plotIndexNotLattice(obj, objCPUE, main="", save=NULL, bar=1.96, ...)
```

# Arguments

	not used
bar	standard deviation of the normal distribution (1.96 is the approximate value of the 97.5 percentile point).
save	not used
main	title for figure
objCPUE	data frame of CPUE indices from Awatea's results file (e.g., $currentRes\$CPUE$ ).
obj	$\label{thm:continuous} \mbox{data frame of survey indices from Awatea's results file (\it{e.g.}, \texttt{currentRes}\$Survey).}$

### Value

### Four postscript files:

```
survIndSer.eps...each survey panel focuses on the years of the survey; survIndSer2.eps...each panel uses a fixed set of years that span all surveys; survIndSer3.eps...one panel showing all series normalised to their means; survIndSer4.eps...compares normalised GIG series with CPUE series.
```

### Author(s)

Andrew Edwards, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC

```
plt.idx
```

plotRmcmcPOP 33

plotRmcmcPOP	Plot Recruitment Posterior Quantiles

# Description

Plot recruitment posteriors quantiles as one graph over time.

### Usage

# Arguments

obj	MCMC data frame of $R$ (currentMCMC\$R).	
р	quantiles to use in quantBox.	
xyType	type of plot (currently only uses quantBox).	
lineType	line types to use in quantBox.	
refLines	reference lines to add to plot.	
xLim	limits of the x-axis.	
yLim	limits of the y-axis.	
userPrompt	not used	
save	not used	
tcl.val	tick length.	
yaxis.by	increments along the y-axis to place tick marks.	
yLab	label for the y-axis.	
	additional arguments passed to the function rect.	

#### Note

AME: Plot recruitment posteriors quantiles as one graph over time.

Already have the full posterior densities done.

Using plotBmcmcPOP as template, but will be simpler as no extra stuff. Probably not simplifying down as much as could due to time constraints.

Adding yLab and then using for exploitation plot also.

### Author(s)

Andrew Edwards, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC

34 plotSnail

### See Also

```
plotBmcmcPOP, plotB2
```

plotSnail

Plot Snail Trails of Exploitation vs. Biomass

# Description

Plot the historical progression of the ratio  $u_t/u_{MSY}$  against  $B_t/B_{MSY}$ .

### Usage

### **Arguments**

BoverBmsy numeric matrix of  $B_t$  over  $B_{MSY}$ .

UoverUmsy numeric matrix of  $u_t$  over  $u_{MSY}$ .

p quantiles to show the bulk of the distribution.

xLim limits of the x-axis. yLim limits of the y-axis.

Lwd line width of the snail trail.

### **Details**

The graph attempts to show the time history of the exploitation rate compared to the spawning biomass using a precautionary framework recast in Bayesian terms.

# Note

The term snail trail comes from PJS.

### Author(s)

Andrew Edwards, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC

```
plotBVBnorm
```

plotTracePOP 35

plotTracePOP Plot MCMC Traces (AME Version)

### **Description**

Plot Markov-chain Monte Carlo traces. This is a diagnostic plot for deciding whether a chain shows unwanted trends by showing the trace of moving quantiles. (Modified from the **scapeMCMC** function plotTrace.)

### Usage

```
plotTracePOP(mcmc, axes=FALSE, same.limits=FALSE,
    between=list(x=axes,y=axes), div=1, span=1/4, log=FALSE,
    base=10, main=NULL, xlab=NULL, ylab=NULL, cex.main=1.2,
    cex.lab=1, cex.strip=0.8, cex.axis=0.8,
    las=0, tck=0.5, tick.number=5, lty.trace=1, lwd.trace=1,
    col.trace="grey", lty.median=1, lwd.median=1,
    col.median="black", lty.quant=2, lwd.quant=1,
    col.quant="black", plot=TRUE, probs=c(0.025, 0.5, 0.975), ...)
```

mcmc	MCMC chain(s) as a vector, data frame or meme object.	
axes	whether axis values should be plotted.	
same.limits	whether panels should have same x-axis limits.	
between	list with $x$ and $y$ indicating panel spacing.	
div	denominator to shorten values on the y axis.	
span	smoothness parameter (not used).	
log	whether values should be log-transformed.	
base	logarithm base.	
main	main title.	
xlab	x-axis title.	
ylab	y-axis title.	
cex.main	size of main title.	
cex.lab	size of axis labels.	
cex.strip	size of strip labels.	
cex.axis	size of tick labels.	
las	orientation of tick labels: 0=parallel, 1=horizontal, 2=perpendicular, 3=vertical.	
tck	tick mark length.	
tick.number	number of tick marks.	
lty.trace	line type of trace.	

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lwd.trace	line width of trace.
col.trace	colour of trace.
lty.median	line type of median.
lwd.median	line width of median.
col.median	colour of median.
lty.quant	line type of quantile trace.
lwd.quant	line width of quantile trace.
col.quant	colour of quantile trace.
plot	whether to draw plot.
probs	quantile values for quantile trace.
	passed to panel.trace (not used).

### Value

When plot=TRUE, a trellis plot is drawn and a data frame is returned, containing the data used for plotting. When plot=FALSE, a trellis object is returned.

### Note

The Args function from the **gdata** package is recommended for reviewing the arguments, instead of args.

### See Also

```
\tt xyplot and \tt panel.loess are the underlying drawing functions, and \tt traceplot is a similar non-trellis plot.
```

```
plotTracePOP, plotAuto, plotCumu, and plotSplom are diagnostic plots. plotDensPOP, plotDens, plotQuant, and plotChains are posterior plots. scapeMCMC-package gives an overview of the scapeMCMC package.
```

Plot Vulnerable Biomass	
	Plot Vulnerable Biomass

# Description

Plot vulnerable biomass from posterior as boxplots, and add catch bars on same graph.

plotVBcatch 37

# Usage

# **Arguments**

obj	MCMC data frame of $B$ (currentMCMC\$B).
currentRes1	list/scape object created by importCol2.
р	quantiles to use in quantBox.
хуТуре	type of plot (currently only uses quantBox).
lineType	line types to use in quantBox.
refLines	reference lines to add to plot.
xLim	limits of the x-axis.
yLim	limits of the y-axis.
userPrompt	not used
save	not used
xLab	x-coordinates for labels.
yLab	y-coordinates for labels.
textLab	text labels to display on plot.
yaxis.by	increments along the y-axis to place tick marks.
tcl.val	tick length.

# Note

AME: This function is essentially a tweak of plotBmcmcPOP.

## Author(s)

Andrew Edwards, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC

additional arguments passed to the function rect.

# See Also

```
plotBmcmcPOP, plotB2
```

38 plotVBnorm

plotVBnorm

Plot Spawning and Vulnerable Biomass Relative to Virgin

# Description

Plot spawning and vulnerable biomass boxplots relative to virgin levels  $B_0$  and  $V_0$ , respectively.

## Usage

```
plotVBnorm( mcmcObj,
            p=c(0.025, 0.25, 0.5, 0.75, 0.975),
            xyType="quantBox",
            lineType=c(3,2,1,2,3),
            refLines=NULL, xLim=NULL, yLim=NULL,
            userPrompt=FALSE, save=T, xLeq=0.7, yLeq=0.9,
            yaxis.by=0.02, tcl.val=-0.2,
            B.col="black", VB.col="black", ...)
plotBVBnorm(mcmcObj,
            p=c(0.025, 0.25, 0.5, 0.75, 0.975),
            xyType="quantBox",
            lineType=c(3, 2, 1, 2, 3),
            refLines=NULL, xLim=NULL, yLim=NULL,
            userPrompt=FALSE, save=T, xLeg=0.7, yLeg=0.9,
            yaxis.by=0.02, tcl.val=-0.2,
            B.col="black", VB.col="black", ...)
```

# **Arguments**

mcmcObj

```
quantiles to use in quantBox.
                  type of plot (currently only uses quantBox).
xyType
lineType
                  line types to use in quantBox.
refLines
                  reference lines to add to plot.
xLim
                 limits of the x-axis.
yLim
                 limits of the y-axis.
userPrompt
                 not used
                  not used
save
xLeg
                  x-coordinate for legend.
                  y-coordinate for legend.
yLeg
yaxis.by
                  increments along the y-axis to place tick marks.
tcl.val
                 tick length.
B.col
                  colour for spawning biomass.
VB.col
                  colour for vulnerable biomass.
                  not used
. . .
```

MCMC list object (currentMCMC).

plt.ageResidsPOP 39

#### Note

AME: tried in separate file, but then changed that to lattice and wouldn't be good format for Arni's boxplots.

```
Based on plotVBcatch (tweaking some). currentRes1 is local currentRes.
```

#### Author(s)

Andrew Edwards, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC

#### See Also

```
plotVBcatch, plotBmcmcPOP
```

# Description

Plot model residuals as boxplots or qq-plots.

## Usage

```
plt.ageResidsPOP (obj, ages=c(2,60), pct=c(5,25,50,75,95), main=NULL)
plt.ageResidsqqPOP(obj, ages=c(2,60), pct=c(5,25,50,75,95), main=NULL)
plt.yearResidsPOP (obj, ages=c(2,60), pct=c(5,25,50,75,95), main=NULL, fill.in=TRUE, ...)
plt.cohortResids (obj, ages=c(2,59), pct=c(5,25,50,75,95), main=NULL)
```

## **Arguments**

```
obj output from stdRes.CA.

ages age classes to plot.

pct quantiles to show in boxplot or qq-plot.

main title for plot if desired.

fill.in logical: if TRUE, add missing years to boxplot.

additional arguments for boxplot.
```

## **Details**

```
plt.ageResidsPOP....plot age class residuals as boxplots.
plt.ageResidsqqPOP...plot age class residuals as qq-plot.
plt.yearResidsPOP....plot age residuals by year as boxplots.
plt.cohortResids....plot age residuals by cohort as boxplots.
```

40 plt.allTraces

## Note

Some trouble noted adding text and legend.

# Author(s)

Andrew Edwards, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC

#### See Also

```
plt.allTraces,plt.expRate,plt.idx,plt.mcmcGraphs,plotIndexNotLattice,
plotChains,plotCPUE
```

plt.allTraces

Plot MCMC Traces

# **Description**

Plot traces from MCMC samples.

# Usage

```
plt.allTraces(obj, bioYrList=NULL, recYrList=NULL, save=TRUE)
```

# **Arguments**

obj vector of MCMC samples.

bioYrList years to plot spawning biomass traces.

recYrList years to plot recruitment traces.

save logical: if TRUE, save figure to a raster file (.jpg).

#### Note

Appears to be some figure requested by PJS.

## Author(s)

Andrew Edwards, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC

## See Also

```
plotChains
```

plt.expRate 41

nlt	.expRate	
$\rho_{\perp}$	• CAPITALL	

Plot Exploitation Rate

## **Description**

Plot exploitation rate against year.

# Usage

```
plt.expRate(obj, yLim=c(0,0.5), xLim=c(1954,2005))
```

# Arguments

```
obj an object from load.allResFiles. yLim limits of the y-axis.
```

xLim limits of the x-axis.

#### **Details**

Simple points and lines plot.

# Author(s)

Andrew Edwards, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC

# See Also

```
plt.idx
```

plt.idx

Plot Survey Index Residuals

# **Description**

Plot the survey index residuals as a quantile-quantile plot (see qqnorm).

# Usage

```
plt.idx(obj, main="Residuals", save=NULL, ...)
```

# **Arguments**

```
obj a data frame with columns Year, stdRes, and Fit.
main title for the plot.
save logical: if TRUE, save the figure to a raster file (.png).
```

... not used

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## **Details**

QQ-plots show sample quantiles vs. theoretical quantiles.

#### Note

The save option has been disabled for some reason.

## Author(s)

Andrew Edwards, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC

## See Also

```
plt.stdResids,plotIndexNotLattice
```

plt.mpdGraphs

Plot a Set of Figures for MPD and MCMC

# **Description**

Plot a set of postscript figures .eps for the MPD (mode of the posterior distribution) and MCMC (Monte Carlo Markoff Chain) results.

## Usage

```
plt.mpdGraphs(obj, save = FALSE)
plt.mcmcGraphs(mcmcObj, projObj, save=FALSE, xlimrec=c(0, 2e+05))
```

# **Arguments**

```
an Awatea results object (e.g., currentRes).

mcmcObj an Awatea MCMC object (e.g., currentMCMC).

projObj an Awatea projected biomass object (e.g., currentProj).

save not used

xlimrec range for recruitments (not used).
```

# **Details**

Creates a whole heap o postscript files.

plt.mpdGraphs 43

#### Value

```
plt.mpdGraphs creates the following postscript files:
exploit.eps.....annual exploitation rate,
recruits.eps.....annual recruitment at age 1,
selectivity.eps.....selectivity curves for commercial gear(s) and survey(s),
ageComm.eps......fits to annual commercial age composition (panel plots),
ageSurv.eps............fits to annual survey age composition (panel plots),
survIndSer.eps......four figures of survey indices (calls plotIndexNotLattice),
commAgeResids.eps.....standardised residuals for commercial gear,
survAgeResidsSer.eps...standardised residuals for surveys,
meanAge.eps......mean age for catch and surveys,
stockRecruit.eps.....stock recruitment function.
plt.mcmcGraphs creates the following postscript files:
recruitsMCMC.eps.....boxplots of annual MCMC recruitment,
exploitMCMC.eps.....boxplots of annual MCMC exploitation rate,
pdfBiomass.eps......density panel plots of annual female spawning biomass,
pdfRecruitment.eps....density panel plots of annual recuitment,
selectivityMCMC.eps....not currently implemented,
traceRecruits.eps.....panel plots of annual recruitmwent traces with running quantiles,
traceBiomass.eps.....panel plots of annual spawning biomass with running quantiles,
traceParams.eps.....panel plots of parameter traces with running quantiles,
splitChain.eps.....panel plots of cumulative parameter estimate chains,
VBcatch.eps.....boxplots of annual vulnerable biomass and barplots of catch,
BVBnorm.eps.....spawning and vulnerable biomass relative to their virgin levels,
Bproj.eps.....boxplots of spawning biomass – MCMCs and projections,
```

# Note

The function plt.mpdGraphs needs some cleaning up. For example, there appears to be extensive reliance on the object currentRes whereas the function should be using obj, which is the actual results object within the function.

The same is true for plt.mcmcGraphs where the global objects currentMCMC and currentProj are use within the function rather than using the local objects mcmcObj and projObj.

#### Author(s)

Andrew Edwards, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC

snail.eps.....time series of  $u_t/u_{MSY}$  vs.  $B_t/B_{MSY}$ , pairs [1, 2, 3].eps.....pairs plot of parameter MCMC samples.

## See Also

plt.mcmcGraphs, plotIndexNotLattice, plotCPUE, plotChains, plotSnail

44 plt.quantBio

plt.numR

Plot Numbers at Age at Equilibrium

## Description

Plot numbers at age at equilibrium. Plot recruitment (age 1).

# Usage

```
plt.numR(obj, minYr = NULL)
```

# **Arguments**

obj an Awatea results object from load.allResFiles.
minYr minimum year to display in plot.

#### Note

Not sure what this plot is used for (RH).

## Author(s)

Andrew Edwards, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC

## See Also

```
plotRmcmcPOP
```

plt.quantBio

Plot Quantile Boxes of MCMC and Projected Biomass

## Description

Plots MCMC and projected biomass as quantile boxes, the former in black, the latter in red.

## Usage

```
plt.quantBio(obj, projObj=NULL, policy=NULL,
    p=c(0.025,0.25,0.5,0.75,0.975), xyType="lines",
    lineType=c(3,2,1,2,3), refLines=NULL,
    xLim=NULL, yLim=NULL, userPrompt=FALSE, save=T)

plt.quantBioBB0(obj, projObj=NULL, policy=NULL,
    p=c(0.025,0.25,0.5,0.75,0.975), xyType="lines",
    lineType=c(3,2,1,2,3), refLines=NULL,
    xLim=NULL, yLim=NULL, userPrompt=FALSE, save=T,
```

plt.quantBio 45

```
main="", cex.main="", tcl.val=-0.2,
xaxis.by=1, yaxis.by=10000, xaxis.lab="Year",
yaxis.lab="Spawning biomass"
```

## **Arguments**

obj	an Awatea MCMC object (e.g., currentMCMC).
projObj	an Awatea projected biomass object (e.g., currentProj).
policy	numeric vector specifying catch policy.
р	quantiles to use from the biomass samples.
хуТуре	string specifying type of plot.
lineType	line types for the quantiles if xyType="lines".
refLines	reference points.
xLim	limits of the x-axis.
yLim	limits of the y-axis.
userPrompt	logical: if TRUE prompts user before figure is drawn.
save	logical: if TRUE save figure as a raster file .png.
main	character string specifying a title for the plot.
cex.main	font size for figure title.
tcl.val	tick length.
xaxis.by	tick mark intervals for x-axis.
yaxis.by	tick mark intervals for y-axis.
xaxis.lab	label for x-axis.
yaxis.lab	label for y-axis.

# Value

List of the reconstructed (MCMC) and projected results.

# Note

plt.quantBioBBO performs similarly as for plt.quantBio but uses  $B_t/B_0$  instead of  $B_t$ .

# Author(s)

Andrew Edwards, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC

# See Also

```
plotBmcmcPOP, plotVBcatch, plotBVBnorm, plotRmcmcPOP
```

plt.stdResids

plt.ssbVbCatch

Plot Annual Spawning and Vulnerable Biomass

# **Description**

Plot MPD values of spawning biomass (SB) and vulnerable biomass(VB), as well as catch, against year.

# Usage

```
plt.ssbVbCatch(obj, x1=1966, xLim=c(1954,2005), yLim=c(0,25000))
```

#### **Arguments**

an Awatea results object from load.allResFiles.

x1 year to start plotting SB and VB lines.

xLim limits of the x-axis.

yLim limits of the y-axis.

#### Note

This analysis uses the MPD (mode of the posterior distribution) values for B and V.

## Author(s)

Andrew Edwards, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC

## See Also

```
load.allResFiles, get.resFile, plt.expRate
```

plt.stdResids

Plot Diagnostics for Standardised Residuals

# **Description**

Plot standardised residuals against year, fitted value, and theoretical residuals.

# Usage

readAD 47

# **Arguments**

obj	a data frame with columns Year, stdRes, and Fit.
pct	percentiles to display as horizontal lines on the quantile-quantile plot.
main	title for the figure.
yLim	limits of the y-axis.
xLim	limits of the x-axis.

## **Details**

Figure provides three panels of standardised residuals *vs.* (i) years, (ii) fitted or predicted values, and (iii) theoretical quantiles.

# Author(s)

Andrew Edwards, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC

# See Also

```
plt.ssbVbCatch
```

readAD

Read ADMB Input and Create AWATEA Class Object

# Description

Read the ADMB input file and create an AWATEA class object.

# Usage

```
readAD(txt)
```

# Arguments

txt string name of an Awatea input file.

#### **Details**

The Awatea input file contains headers (lines prefixed with "#") and data that are read sequentially into the model by the binary executable Awatea.exe.

48 refPoints

#### Value

```
An AWATEA class cobject with the slots:

txtnam....character: name of the input file,
input.....list: each line of the input file with a label specifying line number and
.......indicating whether the line is a Comment or Data,
dnam.....character: vector of strings specifying data contents labelled by line number,
nvars.....numeric: number of data variables,
vdesc.....character: vector of strings specifying data contents labelled by
.......variable number (e.g., v001),
vars.....list: numeric values of data labelled by variable number,
gcomm.....character: vector of comments labelled by line number,
vcomm.....character: vector of variable names labelled by line number,
vcomm.....character: vector of variable names labelled by line number,
output....list: Awatea results file imported by function importRes,
reweight...list: empty (later populated by function reweight).
```

#### Author(s)

Rowan Haigh, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC.

#### References

Hilborn, R., Maunder, M., Parma, A., Ernst, B. Payne, J., and Starr, P. (2003) Coleraine: a generalized age-structured stock assessment model. School of Aquatic and Fishery Sciences, University of Washington, 54 p.

#### See Also

```
runADMB, reweight
```

refPoints

Calculate Reference Points

## Description

Calculate reference points relative to either  $B_{MSY}$  or  $B_0$ .

## Usage

reweight 49

# **Arguments**

mcmcObj	MCMC list object (e.g., currentMCMC).
projObj	projected biomass list object (e.g., currentProj).
msyObj	MSY list object (e.g., currentMSY).
refLevels	reference levels relative to $B_{MSY}$ (or $B_0$ ).
B00bj	vector of $B_0$ MCMC values (e.g., B0 . MCMC).
refNames	names of the $B_0$ reference levels reflevels.

#### Value

List of reference points relative to either  $B_{MSY}$  or  $B_0$ .

#### Note

```
Call from Sweave as refPoints() or, in full:
refPoints(currentMCMC, currentProj, currentMSY, refLevels=c(0.4,0.8,1))
```

## Author(s)

Andrew Edwards, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC

## See Also

```
calc.refVal
```

reweight	Re	ewe	ig	ht	A	$b\iota$	ı
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*undance and Composition Data* 

# Description

Weight the abundance data by adjusting survey and CPUE CVs, and weight the composition data by adjusting the effective sample size N.

## **Usage**

```
reweight(obj, cvpro=FALSE, mean.age=TRUE, ...)
```

## **Arguments**

obj an AWATEA class object created initially by readAD.

CV process error added to CV observation error:  $c_t = \sqrt{c_o^2 + c_p^2}$ ; cvpro

if FALSE index CVs are reweighted using the standard deviation of normalized

residuals.

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mean.age logical: if TRUE, use mean-age residuals to reweight the effective N for the age composition data (see Francis 2011); if FALSE, reweight N using  $\Sigma(P(1-P))/\Sigma(O-P)^2$ , where O = observed proportions-at age and P = predicted/fitted proportions-at-age. . . . . additional arguments to reweight.

#### **Details**

For the reweight to work, a corresponding Awatea results file (.res) with the same prefix as the input file must be available in the working directory before calling readAD. This will populate the output slot with fitted data that the reweight needs.

## Value

An AWATEA class cobject with the slots outlined in readAD with the following slot populated by this function:

```
reweight...list of reweight results:
..nrwt....the number of the current reweighting,
..survey...survey indices with CV values (observed, fitted, normalised residuals, reweighted),
..cpue....if used in the model, CPUE indices with CV values (as above),
..wNcpa...reweighted effective N for commercial compositions (proportions-at-age),
..wNspa...reweighted effective N for survey compositions,
..SDNR....standard deviation of normalised residuals for abundance and composition data,
..wj....weights for composition data from a mean-age weighted calculation (Francis 2011).
```

## Author(s)

Rowan Haigh, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC.

#### References

Francis, R.I.C.C. (2011, in press) Data weighting in statistical fisheries stock assessment models. *Canadian Journal of Fisheries and Aquatic Sciences*.

## See Also

```
runADMB, readAD, stdRes.CA, stdRes.index
```

runADMB

Run AD Model Builder Code for Awatea

## Description

Run compiled AD Model Builder code called Awatea to reconstruct a population trajectory for a marine fish stock.

runADMB 51

#### Usage

```
runADMB(filename.ext, wd=getwd(), strSpp="YMR", runNo=25, rwtNo=0,
    doMPD=FALSE, N.reweight=0, cvpro=FALSE, mean.age=TRUE,
    doMCMC=FALSE, mcmc=1e+06, mcsave=1000, ADargs=NULL, verbose=FALSE,
    doMSY=FALSE, msyMaxIter=15000, msyTolConv=0.01,
    endStrat=0.301, stepStrat=0.001, ...)
```

## **Arguments**

filename.ext character file name including its extension.

wd character string specifying the working directory, if different from the current

working directory.

strSpp string 3-letter code of the species.

runNo the model run number.
rwtNo the reweight number.

dompd logical: if True, perform an MPD analysis.

N. reweight the number of reweights to perform in the MPD analysis.

cvpro CV process error added to CV observation error:  $c_t = \sqrt{c_o^2 + c_p^2}$ ;

if FALSE index CVs are reweighted using the standard deviation of normalized

residuals.

mean.age logical: if TRUE, use mean-age residuals to reweight the effective N for the age

composition data (see Francis 2011);

if FALSE, reweight N using  $\Sigma(P(1-P))/\Sigma(O-P)^2$ , where O = observed

proportions-at age and P = predicted/fitted proportions-at-age.

doMCMC logical: if TRUE, perform an MCMC analysis.

mcmc number of MCMC iterations to perform.

mcsave frequency of MCMC iterations to save.

ADargs additional arguments for a call to Awatea.

verbose logical: if TRUE, spew Awatea messages to the R console.

domsy logical: if TRUE, perform an MSY analysis.

msyMaxIter maximum iterations for the MSY calculations.

msyTolConv tolerance for convergence in the MSY calculations.

endStrat maximum fishing mortality for the MSY analysis.

stepStrat fishing mortality step size for the MSY analysis.

. . . additional arguments (not currently used for any purpose).

#### **Details**

This function is primarily used to automate MPD reweightings and to perform the MSY calculations. The MCMCs are better run from a command line console on a super computer as they generally require > 12 h to complete.

Note that once an MCMC has been created, the user can run various projections separately using awatea -ind filename.ext -mceval on the command line.

52 runMCMC

#### Author(s)

Rowan Haigh, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC.

#### References

Edwards, A.M., Starr, P.J., and Haigh, R. (2010, in revision). Stock assessment for Pacific Ocean Perch (*Sebastes alutus*) in Queen Charlotte Sound, British Columbia. *Canadian Science Advisory Secretariat, Research Document*.

Edwards, A.M., Haigh, R., and Starr, P.J. (2011, in revision). Stock assessment and recovery potential assessment for Yellowmouth Rockfish (*Sebastes reedi*) along the Pacific coast of Canada. *Canadian Science Advisory Secretariat, Research Document*.

Francis, R.I.C.C. (2011, in press) Data weighting in statistical fisheries stock assessment models. *Canadian Journal of Fisheries and Aquatic Sciences*.

Hilborn, R., Maunder, M., Parma, A., Ernst, B. Payne, J., and Starr, P. (2003) Coleraine: a generalized age-structured stock assessment model. School of Aquatic and Fishery Sciences, University of Washington, 54 p.

#### See Also

importRes, readAD, reweight, runSweave

runMCMC

Wrapper to Function <runSweaveMCMC>

## **Description**

This small utility function simply provides a wrapper to the function runSweaveMCMC so that multiple documents can be built at once.

# Usage

```
runMCMC(runs=7, rewts=0:6, cpue=FALSE, estM=TRUE)
```

# **Arguments**

runs	the run number(s).
rewts	the reweight number(s).
cpue	logical: if TRUE, retain the CPUE figures in the Sweave file.
estM	logical: if TRUE the routine will assume natural mortality $M$ was estimated in the run, and the Sweave code uses the parameters "M_1" and "M_2"; if FALSE, the function removes these parameters from the Sweave file.

## **Details**

Essentially loops through runSweaveMCMC using (i in runs) and (j in rewts).

runMPD 53

## Value

Produces multiple Sweave documents and PDF files for MCMCs.

#### Note

```
Seave files can be found in the library directory:
```

```
.../R/.../library/PBSawatea/snw/ymrrun-masterMCMC.Snw .../R/.../library/PBSawatea/snw/ymrrun-masterMCMC20yrProjs.Snw
```

# Author(s)

Rowan Haigh, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC

## See Also

```
runSweaveMCMC, runSweave, runMPD
```

runMPD

Wrapper to Function <runSweave>

# Description

This small utility function simply provides a wrapper to the function runSweave so that multiple documents can be built at once.

# Usage

```
runMPD(runs=1, rewts=0:6, cpue=FALSE)
```

#### **Arguments**

runs the run number(s).
rewts the reweight number(s).

cpue logical: if TRUE, retain the CPUE figures in the Sweave file.

#### **Details**

Essentially loops through runSweave using (i in runs) and (j in rewts).

## Value

Produces multiple Sweave documents and PDF files.

## Note

```
A recent Seave file called ymrrun-master. Snw can be found in the library directory: .../R/.../library/PBSawatea/snw
```

54 runSweave

#### Author(s)

Rowan Haigh, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC

#### See Also

```
runSweave, runSweaveMCMC, runMCMC
```

runSweave

Run Customised Sweave Files for Awatea MPD Runs

# **Description**

Create and run customised Sweave files for Awatea MPD runs.

# Usage

```
runSweave(wd=getwd(), cpue=FALSE, strSpp="YMR",
    filename="input25-ymr.txt", runNo=25, rwtNo=0,
    running.awatea=0, Nsurvey=5)
```

## **Arguments**

wd working directory in which Awatea input and master Sweave files occur.

cpue logical: if TRUE the routine will leave calls to CPUE figures in the Sweave file,

otherwise, they are removed.

strSpp three-letter code that identifies the species.

filename name of Awatea input file.

runNo the run number that identifies a unique combination of input values.

rwtNo the reweight number of the MPD run to build and collate figures.

running.awatea

numeric: if 0 load a previous .rep file, if 1 re-run  ${\tt Awatea}$ 

Nsurvey number of surveys specified in the input file; controls image creation and place-

holders.

#### **Details**

The values specified by the arguments (or derived variables) are directly substituted into the Sweave file wherever similarly named variables preceded by the @ symbol occur.

The cpue switch signals the removal of pieces of Sweave code if cpue=TRUE.

The Nsurvey argument essentially copies one Sweave line into Nsurvey lines for a set of lines identified by unique Sweave snippets.

runSweaveMCMC 55

# Value

```
A customised Sweave file for runNo and rwtNo is created in a subdirectory ./strSpprun/MPD.runNo.rwtNo from where it is run.
```

#### Note

```
A fairly recent Seave file called ymrrun-master. Snw can be found in the library directory: .../R/.../library/PBSawatea/snw
```

# Author(s)

Rowan Haigh, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC

## See Also

```
runMPD, runSweaveMCMC, runADMB
```

runSweaveMCMC

Run Customised Sweave Files for Awatea MCMC Runs

# Description

Create and run customised Sweave files for Awatea MCMC runs.

# Usage

```
runSweaveMCMC( wd=getwd(), cpue=FALSE, estM=TRUE,
    strSpp="YMR", filename="input25-ymr.txt",
    runNo=25, rwtNo=0, running.awatea=0)
```

#### **Arguments**

wd	working directory in which Awatea input and master Sweave files occur.	
cpue	logical: if $\mathtt{TRUE}$ the routine will leave calls to CPUE figures in the Sweave file, otherwise, they are removed.	
estM	logical: if TRUE the routine will assume natural mortality $M$ was estimated in the run, and the Sweave code uses the parameters "M_1" and "M_2"; if FALSE, the function removes these parameters from the Sweave file.	
strSpp	three-letter code that identifies the species.	
filename	name of Awatea input file.	
runNo	the run number that identifies a unique combination of input values.	
rwtNo	the reweight number of the MPD run to build and collate figures.	
running.awatea		
	numeric: if 0 load a previous .rep file, if 1 re-run Awatea	

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## **Details**

The values specified by the arguments (or derived variables) are directly substituted into the Sweave file wherever similarly named variables preceded by the @ symbol occur.

The cpue switch signals the removal of pieces of Sweave code if cpue=TRUE.

The estM switch signals the removal of "M\_1" and "M\_2" from the Sweave code if estM=FALSE.

#### Value

```
A customised Sweave file for runNo and rwtNo is created in a subdirectory ./strSpprun/MCMC.runNo.rwtNo from where it is run.
```

## Note

```
Seave files can be found in the library directory:
.../R/.../library/PBSawatea/snw/ymrrun-masterMCMC.Snw
.../R/.../library/PBSawatea/snw/ymrrun-masterMCMC20yrProjs.Snw
```

# Author(s)

Rowan Haigh, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC

## See Also

```
runMCMC, runSweave, runADMB
```

srFun

Stock Recruitment Function

# Description

Take a vector of spawners in year t-1 and calculate recruits in year t.

# Usage

```
srFun(spawners, h = h.mpd, R0 = R0.mpd, B0 = B0.mpd)
```

# Arguments

spawners	a vector of spawners where either: each element corresponds to spawners in year $t$ -1 or the vector calculates a single year but multiple MCMCs.
h	steepness parameter value.
R0	recruitment at $t = 0$ (virgin conditions).
в0	spawning biomasss at virgin conditons.

stdRes.CA 57

#### **Details**

(AME wording) To input a vector of spawners in year t-1 and calculate recruits in year t. Output for recruits is vector, each element corresponds to spawners the the year before, so will usually want to shift the output by 1 so that recruits in year t are based on spawners in year t-1.

Can also have each input as a vector (used when calculating a single year but multiple MCMCs, as in first year of projections is based on penultimate year of MCMC calcualtions).

#### Value

A vector of recruitments in year t.

#### Note

This function was originally a subfunction in plt.mpdGraphs.

## Author(s)

Andrew Edwards, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC

## See Also

```
plt.mpdGraphs
```

stdRes.CA

Calculate Standardised Residuals for Robust Normal Likelihood

# Description

Calculate the standardised residuals for Awatea's implementation of the Fournier *robustified* normal likelihood for proportions-at-length.

Based on PJS's summary of the CASAL document and ACH's change to length.

## Usage

```
stdRes.CA(obj, trunc=3, myLab="Age Residuals", prt=TRUE)
```

## **Arguments**

obj scape/list object of Awatea's results file (.res).

trunc maximum standardised residual; values greater than this are set to trunc.

myLab general label for the output.

prt logical: if TRUE, print the results.

## Value

List object of standardised residuals.

58 stdRes.index

## Author(s)

Andrew Edwards, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC

#### See Also

```
stdRes.index, importCol2, reweight
```

stdRes.index

Calculate Standardised Residuals for Abundance Indices

# Description

Calculate the standardised residuals for commercial and survey indices.

## Usage

```
stdRes.index(obj, label=NULL, prt=TRUE)
```

## **Arguments**

obj data frame of observed and fitted index values from Awatea's results file (.res).

label general label for the output.

prt logical: if TRUE, print the results.

## Value

Input data frame with additional column of standardised residuals.

# Author(s)

Andrew Edwards, Pacific Biological Station, Fisheries and Oceans Canada, Nanaimo BC

# See Also

```
stdRes.CA, importCol2, reweight
```

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