Package ‘mrds’

May 12, 2020

Imports  optimx (>= 2013.8.6), mgcv, numDeriv, Rsolnp
Maintainer  Laura Marshall <lhm@st-andrews.ac.uk>
License  GPL (>= 2)
Title  Mark-Recapture Distance Sampling
LazyLoad  yes
Author  Jeff Laake <jeff.laake@noaa.gov>, David Borchers
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        Miller <dave@ninepointeightone.net> and Jon Bishop
Description  Animal abundance estimation via conventional, multiple covariate
    and mark-recapture distance sampling (CDS/MCDS/MRDS). Detection function
    fitting is performed via maximum likelihood. Also included are diagnostics
    and plotting for fitted detection functions. Abundance estimation is via a
    Horvitz-Thompson-like estimator.
Version  2.2.2
BugReports  https://github.com/DistanceDevelopment/mrds/issues
Depends  R (>= 3.0)
Suggests  testthat
RoxygenNote  7.1.0
NeedsCompilation  no
Repository  CRAN
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Examples of distance sampling analyses are available at http://examples.distancesampling.org/.

For help with distance sampling and this package, there is a Google Group https://groups.google.com/forum/#!forum/distance-sampling.

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Description

Add a line or lines to a plot of the detection function which correspond to a given covariate combination. These can be particularly useful when there is a small number of factor levels or if quantiles of a continuous covariate are specified.

Usage

```r
add_df_covar_line(ddf, data, ndist = 250, ...)
```

Arguments

- `ddf`: a fitted detection function object
- `data`: a `data.frame` with the covariate combination you want to plot
- `ndist`: number of points to evaluate the detection function at
- `...`: extra arguments to give to `lines` (lty, lwd, col)

Details

All covariates must be specified in `data`. Plots can become quite busy when this approach is used. It may be useful to fix some covariates at their median level and plot set values of a covariate of interest. For example setting weather (e.g., Beaufort) to its median and plotting levels of observer, then creating a second plot for a fixed observer with levels of weather.

Arguments to `lines` are supplied in ... and aesthetics like line type (lty), line width (lwd) and colour (col) are recycled. By default lty is used to distinguish between the lines. It may be useful to add a `legend` to the plot (lines are plotted in the order of `data`).

Value

invisibly, the values of detectability over the truncation range

Author(s)

David L Miller

Examples

```r
## Not run:
# fit an example model
data(book.tee.data)
egdata <- book.tee.data$book.tee.dataframe
result <- ddf(dsmodel = ~mcds(key = "hn", formula = ~sex),
  data = egdata[egdata$observer==1, ], method = "ds",
  meta.data = list(width = 4))
```
# make a base plot, showpoints=FALSE makes the plot less busy
plot(result, showpoints=FALSE)

# add lines for sex one at a time
add_df_covar_line(result, data.frame(sex=0), lty=2)
add_df_covar_line(result, data.frame(sex=1), lty=3)

# add a legend
legend(3, 1, c("Average", "sex==0", "sex==1"), lty=1:3)

# alternatively we can add both at once
# fixing line type and varying colour
plot(result, showpoints=FALSE)
add_df_covar_line(result, data.frame(sex=c(0,1)), lty=1,
                 col=c("red", "green"))

# add a legend
legend(3, 1, c("Average", "sex==0", "sex==1"), lty=1,
       col=c("black", "red", "green"))

## End(Not run)

---

**adj.check.order**

*Check order of adjustment terms*

**Description**

'adj.check.order' checks that the Cosine, Hermite or simple polynomials are of the correct order.

**Usage**

```
adj.check.order(adj.series, adj.order, key)
```

**Arguments**

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**Details**

Only even functions are allowed as adjustment terms. Also Hermite polynomials must be of degree at least 4 and Cosine of order at least 3. Finally, also checks that order of the terms >1 for half-normal/hazard-rate, as per p.47 of Buckland et al (2001). If incorrect terms are supplied then an error is throw via stop.

**Value**

Nothing! Just calls stop if something goes wrong.
AIC.ds

Author(s)

David Miller

References


See Also

adjfct.cos, adjfct.poly, adjfct.herm, detfct, mcds, cds

AIC.ds  Akaike's An Information Criterion for detection functions

Description

Extract the AIC from a fitted detection function.

Usage

## S3 method for class 'ds'
AIC(object, ..., k = 2)

Arguments

  object a fitted detection function object
  ... optionally more fitted model objects.
  k penalty per parameter to be used; the default k = 2 is the "classical" AIC

Author(s)

David L Miller
apex.gamma  
Get the apex for a gamma detection function

Description
Get the apex for a gamma detection function

Usage
apex.gamma(ddfobj)

Arguments
  ddfobj       ddf object

Value
the distance at which the gamma peaks

Author(s)
Jeff Laake

assign.default.values  Assign default values to list elements that have not been already assigned

Description
Assigns default values for argument in list x from argument=value pairs in ...if x$argument doesn’t already exist

Usage
assign.default.values(x, ...)

Arguments
x         generic list
...       unspecified list of argument=value pairs that are used to assign values

Value
x - list with filled values

Author(s)
Jeff Laake
average.line  

*Average detection function line for plotting*

**Description**

For models with covariates the detection probability for each observation can vary. This function computes an average value for a set of distances to plot an average line to graphically represent the fitted model in plots that compare histograms and the scatter of individual estimated detection probabilities. Averages are calculated over the observed covariate combinations.

**Usage**

```r
average.line(finebr, obs, model)
```

**Arguments**

- `finebr`: set of fine breaks in distance over which detection function values are averaged and plotted
- `obs`: value of observer for averaging (1-2 individual observers; 3 duplicates; 4 pooled observation team)
- `model`: ddf model object

**Value**

list with 2 elements

- `xgrid`: vector of gridded distance values
- `values`: vector of average detection function values at the `xgrid` values

**Note**

Internal function called from plot functions for ddf objects

**Author(s)**

Jeff Laake

---

average.line.cond  

*Average conditional detection function line for plotting*

**Description**

For models with covariates the detection probability for each observation can vary. This function computes an average value for a set of distances to plot an average line to graphically represent the fitted model in plots that compare histograms and the scatter of individual estimated detection probabilities.
Usage

average.line.cond(finebr, obs, model)

Arguments

finebr  set of fine breaks in distance over which detection function values are averaged and plotted
obs    value of observer for averaging (1-2 individual observers)
model  ddf model object

Value

list with 2 elements:

  xgrid  vector of gridded distance values
  values vector of average detection function values at the xgrid values

Note

Internal function called from plot functions for ddf objects

Author(s)

Jeff Laake

---

book.tee.data  Golf tee data used in chapter 6 of Advanced Distance Sampling examples

---

Description

Double platform data collected in a line transect survey of golf tees by 2 observers at St. Andrews. Field sex was actually colour of the golf tee: 0 - green; 1 - yellow. Exposure was either low (0) or high(1) depending on height of tee above the ground. size was the number of tees in an observed cluster.

Format

The format is: List of 4 $ book.tee.dataframe:'data.frame': 324 obs. of 7 variables: ..$ object : num [1:324] 1 1 2 2 3 3 4 4 5 5 ... ..$ observer: Factor w/ 2 levels "1","2": 1 2 1 2 1 2 1 2 1 2 ... ..$ detected: num [1:324] 1 0 1 0 1 0 1 0 1 0 ... ..$ distance: num [1:324] 2.68 2.68 3.33 3.33 0.34 0.34 2.53 2.53 1.46 1.46 ... ..$ size : num [1:324] 2 2 2 2 1 1 2 2 2 2 ... ..$ sex : num [1:324] 1 1 1 1 0 0 1 1 1 1 ... ..$ exposure: num [1:324] 1 1 0 0 0 0 1 1 0 0 ... $ book.tee.region :'data.frame': 2 obs. of 2 variables: ..$ Region.Label: Factor w/ 2 levels "1","2": 1 2 ..$ Area : num [1:2] 1040 640 $ book.tee.samples :'data.frame': 11 obs. of 3 variables: ..$ Sample.Label: num [1:11] 1 2 3 4 5 6 7
calc.se.Np

Find se of average p and N

Description

Find se of average p and N

Usage

calc.se.Np(model, avgp, n, average.p)

Arguments

model a ddf model object
avgp average p function
n sample size
average.p the average probability of detection for the model

Author(s)

David L. Miller

cdf.ds

Cumulative distribution function (cdf) for fitted distance sampling detection function

Description

Computes cdf values of observed distances from fitted distribution. For a set of observed x it returns the integral of f(x) for the range= (inner, x), where inner is the innermost distance which is observable (either 0 or left if left truncated). In terms of g(x) this is the integral of g(x) over range divided by the integral of g(x) over the entire range of the data (inner, W).

Usage

cdf.ds(model, newdata = NULL)

Arguments

model fitted distance sampling model
newdata new data values if computed for values other than the original observations
Value
vector of cdf values for each observation

Note
This is an internal function that is not intended to be invoked directly. It is called by \texttt{qqplot.ddf} to compute values for Kolmogorov-Smirnov and Cramer-von Mises tests and the Q-Q plot.

Author(s)
Jeff Laake

See Also
\texttt{qqplot.ddf}

cds \hfill \textit{CDS function definition}

Description
Creates model formula list for conventional distance sampling using values supplied in call to \texttt{ddf}

Usage
\begin{verbatim}
cds(
  key = NULL,
  adj.series = NULL,
  adj.order = NULL,
  adj.scale = "width",
  adj.exp = FALSE,
  formula = ~1,
  shape.formula = ~1
)
\end{verbatim}

Arguments
\begin{itemize}
  \item \textbf{key} \hspace{1cm} string identifying key function (currently either "hn" (half-normal),"hr" (hazard-rate), "unif" (uniform) or "gamma" (gamma distribution))
  \item \textbf{adj.series} \hspace{1cm} string identifying adjustment functions cos (Cosine), herm (Hermite polynomials), poly (simple polynomials) or NULL
  \item \textbf{adj.order} \hspace{1cm} vector of order of adjustment terms to include
  \item \textbf{adj.scale} \hspace{1cm} whether to scale the adjustment terms by "width" or "scale"
  \item \textbf{adj.exp} \hspace{1cm} if TRUE uses \texttt{exp(adj)} for adjustment to keep \texttt{f(x)>0}
  \item \textbf{formula} \hspace{1cm} formula for scale function (included for completeness only only formula=\texttt{~1} for \texttt{cds})
  \item \textbf{shape.formula} \hspace{1cm} formula for shape function
\end{itemize}
check.bounds

Value

A formula list used to define the detection function model

fct string "cds"
key key function string
adj.series adjustment function string
adj.order adjustment function orders
adj.scale adjustment function scale type
formula formula for scale function
shape.formula formula for shape function

Author(s)

Jeff Laake; Dave Miller

Description

Simple internal function to check that the optimisation didn’t hit bounds. Based on code that used to live in detfct.fit.opt.

Usage

check.bounds(lt, lowerbounds, upperbounds, ddfobj, showit, setlower, setupper)

Arguments

lt optimisation object
lowerbounds current lower bounds
upperbounds current upper bounds
ddfobj ddf object
showit debug level
setlower were lower bounds set by the user
setupper were upper bounds set by the user

Value

TRUE if bounded (ie parameters close to bound), else FALSE

Author(s)

Dave Miller; Jeff Laake
Check that a detection function is monotone non-increasing.

Usage

```r
check.mono(
  df,
  strict = TRUE,
  n.pts = 100,
  tolerance = 1e-06,
  plot = FALSE,
  max.plots = 6
)
```

Arguments

- `df`: a fitted detection function object
- `strict`: if TRUE (default) the detection function must be "strictly" monotone, that is that \( g(x[i]) \leq g(x[i-1]) \) over the whole range (left to right truncation points).
- `n.pts`: number of equally-spaced points between left and right truncation at which to evaluate the detection function (default 100)
- `tolerance`: numerical tolerance for monotonicity checks (default 1e-6)
- `plot`: plot a diagnostic highlighting the non-monotonic areas (default FALSE)
- `max.plots`: when plot=TRUE, what is the maximum number of plots of non-monotone covariate combinations that should be plotted? Plotted combinations are a random sample of the non-monotonic subset of evaluations. No effect for non-covariate models.

Details

Evaluates a series of points over the range of the detection function (left to right truncation) then determines:

1. If the detection function is always less than or equal to its value at the left truncation point \( g(x) \leq g(\text{left}) \), or usually \( g(x) \leq g(0) \).
2. (Optionally) The detection function is always monotone decreasing \( g(x[i]) \leq g(x[i-1]) \). This check is only performed when strict=TRUE (the default).
3. The detection function is never less than 0 \( g(x) \geq 0 \).
4. The detection function is never greater than 1 \( g(x) \leq 1 \).

For models with covariates in the scale parameter of the detection function is evaluated at all observed covariate combinations.

Currently covariates in the shape parameter are not supported.
Value

TRUE if the detection function is monotone, FALSE if it's not. Warnings are issued to warn the user that the function is non-monotonic.

Author(s)

David L. Miller

---

**Description**

Extract coefficients and provide a summary of parameters and estimates from the output of `ddf` model objects.

**Usage**

```r
## S3 method for class 'ds'
coef(object,...)
## S3 method for class 'io'
coef(object,...)
## S3 method for class 'io.fi'
coef(object,...)
## S3 method for class 'trial'
coef(object,...)
## S3 method for class 'trial.fi'
coef(object,...)
## S3 method for class 'rem'
coef(object,...)
## S3 method for class 'rem.fi'
coef(object,...)
```

**Arguments**

- `object` - `ddf` model object of class `ds`, `io`, `io.fi`, `trial`, `trial.fi`, `rem`, or `rem.fi`.
- `...` - unspecified arguments that are unused at present

**Value**

For `coef.ds` List of data frames for coefficients (scale and exponent (if hazard))

- `scale` - dataframe of scale coefficient estimates and standard errors
- `exponent` - dataframe with exponent estimate and standard error if hazard detection function

For all others Data frame containing each coefficient and standard error
compute.Nht

Note
These functions are called by the generic function coef for any ddf model object. It can be called directly by the user, but it is typically safest to use coef which calls the appropriate function based on the type of model.

Author(s)
Jeff Laake

compute.Nht  Horvitz-Thompson estimates 1/p_i or s_i/p_i

Description
Compute individual components of Horvitz-Thompson abundance estimate in covered region for a particular subset of the data depending on value of group = TRUE (do group abundance); FALSE (do individual abundance)

Usage
compute.Nht(pdots, group = TRUE, size = NULL)

Arguments
pdots  vector of estimated detection probabilities
group if TRUE (do group abundance); FALSE (do individual abundance)
size  vector of group size values for clustered populations

Value
vector of H-T components for abundance estimate

Note
Internal function called by covered.region.dht

Author(s)
Jeff Laake
**covered.region.dht**

Covered region estimate of abundance from Horvitz-Thompson-like estimator

**Description**

Computes H-T abundance within covered region by sample.

**Usage**

```
covered.region.dht(obs, samples, group)
```

**Arguments**

- `obs` observations table
- `samples` samples table
- `group` if TRUE compute abundance of group otherwise abundance of individuals

**Value**

`Nhat.by.sample` - dataframe of abundance by sample

**Note**

Internal function called by dht and related functions

**Author(s)**

Jeff Laake

---

**create.bins**

Create bins from a set of binned distances and a set of cutpoints.

**Description**

This is an internal routine and shouldn’t be necessary in normal analyses.

**Usage**

```
create.bins(data, cutpoints)
```

**Arguments**

- `data` data.frame with at least the column distance.
- `cutpoints` vector of cutpoints for the bins
create.model.frame

**Value**

data data with two extra columns distbegin and distend.

**Author(s)**

David L. Miller

---

create.model.frame  
*Create a model frame for ddf fitting*

**Description**

Creates a model.frame for distance detection function fitting. It includes some pre-specified and computed variables with those included in the model specified by user (formula).

**Usage**

create.model.frame(xmat, scale.formula, meta.data, shape.formula = NULL)

**Arguments**

- **xmat** — dataframe for ddf
- **scale.formula** — user specified formula for scale of distance detection function
- **meta.data** — user-specified meta.data (see ddf)
- **shape.formula** — user specified formula for shape parameter of distance detection function

**Details**

The following fields are always included: detected, observer, binned, and optionally distance (unless null), timesdetected (if present in data). If the distance data were binned, include distbegin and distend point fields. If the integration width varies also include int.begin and int.end and include an offset field for an iterative glm, if used. Beyond these fields only fields used in the model formula are included.

**Value**

model frame for analysis

**Note**

Internal function and not called by user

**Author(s)**

Jeff Laake
create.varstructure  

*Description*

Creates structures needed to compute abundance and variance

*Usage*

```r
create.varstructure(model, region, sample, obs)
```

*Arguments*

- **model**: fitted ddf object
- **region**: region table
- **sample**: sample table
- **obs**: table of object #'s and links to sample and region table

*Details*

The function performs the following tasks: 1) tests to make sure that region labels are unique, 2) merges sample and region tables into a samples table and issue a warning if not all samples were used, 3) if some regions have no samples or if some values of Area were not valid areas given then issue error and stop, then an error is given and the code stops, 4) creates a unique region/sample label in samples and in obs, 5) merges observations with sample and issues a warning if not all observations were used, 6) sorts regions by its label and merges the values with the predictions from the fitted model based on the object number and limits it to the data that is appropriate for the fitted detection function.

*Value*

List with 2 elements:

- **samples**: merged dataframe containing region and sample info - one record per sample
- **obs**: merged observation data and links to region and samples

*Note*

Internal function called by `dht`

*Author(s)*

Jeff Laake
Distance Detection Function Fitting

Description

Generic function for fitting detection functions for distance sampling with single and double observer configurations. Independent observer, trial and dependent observer (removal) configurations are included. This is a generic function which does little other than to validate the calling arguments and methods and then calls the appropriate method specific function to do the analysis.

Usage

```r
ddf(
    dsmodel = call(),
    mrmodel = call(),
    data,
    method = "ds",
    meta.data = list(),
    control = list()
)
```

Arguments

- `dsmodel`: distance sampling model specification
- `mrmodel`: mark-recapture model specification
- `data`: dataframe containing data to be analyzed
- `method`: analysis method
- `meta.data`: list containing settings controlling data structure
- `control`: list containing settings controlling model fitting

Details

The fitting code has certain expectations about `data`. It should be a dataframe with at least the following fields named and defined as follows:

- `object`: object number
- `observer`: observer number (1 or 2) for double observer; only 1 if single observer
- `detected`: 1 if detected by the observer and 0 if missed; always 1 for single observer
- `distance`: perpendicular distance

If the data are for clustered objects, the dataframe should also contain a field named `size` that gives the observed number in the cluster. If the data are for a double observer survey, then there are two records for each observation and each should have the same object number. The code assumes the observations are listed in the same order for each observer such that if the data are subsetted by
observer there will be the same number of records in each and each subset will be in the same object order. In addition to these predefined and pre-named fields, the dataframe can have any number and type of fields that are used as covariates in the dsmodel and mrmodel. At present, discrepancies between observations in distance, size and any user-specified covariates cannot be assimilated into the uncertainty of the estimate. The code presumes the values for those fields are the same for both records (observer=1 and observer=2) and it uses the value from observer 1. Thus it makes sense to make the values the same for both records in each pair even when both detect the object or when observer 1 doesn’t detect the object the data would have to be taken from observer 2 and would not be consistent.

Five different fitting methods are currently available and these in turn define whether dsmodel and mrmodel need to be defined.

<table>
<thead>
<tr>
<th>Method</th>
<th>Single/Double</th>
<th>dsmodel</th>
<th>mrmodel</th>
</tr>
</thead>
<tbody>
<tr>
<td>ds</td>
<td>Single</td>
<td>yes</td>
<td>no</td>
</tr>
<tr>
<td>io</td>
<td>Double</td>
<td>yes</td>
<td>yes</td>
</tr>
<tr>
<td>io.fi</td>
<td>Double</td>
<td>no</td>
<td>yes</td>
</tr>
<tr>
<td>trial</td>
<td>Double</td>
<td>yes</td>
<td>yes</td>
</tr>
<tr>
<td>trial.fi</td>
<td>Double</td>
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<td>yes</td>
</tr>
<tr>
<td>rem</td>
<td>Double</td>
<td>yes</td>
<td>yes</td>
</tr>
<tr>
<td>rem.fi</td>
<td>Double</td>
<td>no</td>
<td>yes</td>
</tr>
</tbody>
</table>

Methods with the suffix ".fi" use the assumption of full independence and do not use the distance sampling portion of the likelihood which is why a dsmodel is not needed. An mrmodel is only needed for double observer surveys and thus is not needed for method ds.

The dsmodel specifies the detection function g(y) for the distance sampling data and the models restrict g(0)=1. For single observer data g(y) is the detection function for the single observer and if it is a double observer survey it is the relative detection function (assuming g(0)=1) of both observers as a team (the unique observations from both observers). In double observer surveys, the detection function is p(y)=p(0)g(y) such that p(0)<1. The detection function g(y) is specified by dsmodel and p(0) estimated from the conditional detection functions (see mrmodel below). The value of dsmodel is specified using a hybrid formula/function notation. The model definition is prefixed with a ~ and the remainder is a function definition with specified arguments. At present there are two different functions, cds and mcds, for conventional distance sampling and multi-covariate distance sampling. Both functions have the same required arguments (key,formula). The first specifies the key function this can be half-normal ("hn"), hazard-rate ("hr"), gamma ("gamma") or uniform ("unif"). The argument formula specifies the formula for the log of the scale parameter of the key function (e.g., the equivalent of the standard deviation in the half-normal). The variable distance should not be included in the formula because the scale is for distance. See Marques, F.F.C. and S.T. Buckland (2004) for more details on the representation of the scale formula. For the hazard rate and gamma functions, an additional shape.formula can be specified for the model of the shape parameter. The default will be ~1. Adjustment terms can be specified by setting adj.series which can have the values: "none", "cos" (cosine), "poly" (polynomials), and "herm" (Hermite polynomials). One must also specify a vector of orders for the adjustment terms (adj.order) and a scaling (adj.scale) which may be "width" or "scale" (for scaling by the scale parameter). Note that the uniform key can only be used with adjustments (usually cosine adjustments for a Fourier-type analysis).
The mrmodel specifies the form of the conditional detection functions (i.e., probability it is seen by observer j given it was seen by observer 3-j) for each observer (j=1,2) in a double observer survey. The value is specified using the same mix of formula/function notation but in this case the functions are glm and gam. The arguments for the functions are formula and link. At present, only glm is allowed and it is restricted to link=logit. Thus, currently the only form for the conditional detection functions is logistic as expressed in eq 6.32 of Laake and Borchers (2004). In contrast to dsmodel, the argument formula will typically include distance and all other covariates that affect detection probability. For example, mrmodel=~glm(formula=~distance+size+sex) constructs a conditional detection function based on the logistic form with additive factors, distance, size, and sex. As another example, mrmodel=~glm(formula=~distance*size+sex) constructs the same model with an added interaction between distance and size.

The argument meta.data is a list that enables various options about the data to be set. These options include:

- point if TRUE the data are from point counts and FALSE (default) implies line transect data
- width distance specifying half-width of the transect
- left distance specifying inner truncation value
- binned TRUE or FALSE to specify whether distances should be binned for analysis
- breaks if binned=TRUE, this is a required sequence of break points that are used for plotting/gof. They should match distbegin, distend values if bins are fixed
- int.range an integration range for detection probability; either a vector of 2 or matrix with 2 columns
- mono constrain the detection function to be weakly monotonically decreasing (only applicable when there are no covariates in the detection function)
- mono.strict when TRUE constrain the detection function to be strictly monotonically decreasing (again, only applicable when there are no covariates in the detection function)

Using meta.data=list(int.range=c(1,10)) is the same as meta.data=list(left=1,width=10). If meta.data=list(binned=TRUE) is used, the dataframe needs to contain the fields distbegin and distend for each observation which specify the left and right hand end points of the distance interval containing the observation. This is a general data structure that allows the intervals to change rather than being fixed as in the standard distance analysis tools. Typically, if the intervals are changing so is the integration range. For example, assume that distance bins are generated using fixed angular measurements from an aircraft in which the altitude is varying. Because all analyses are truncated (i.e., the last interval does not go to infinity), the transect width (and the left truncation point if there is a blindspot below the aircraft) can potentially change for each observation. The argument int.range can also be entered as a matrix with 2 columns (left and width) and a row for each observation.

The argument control is a list that enables various analysis options to be set. It is not necessary to set any of these for most analyses. They were provided so the user can optionally see intermediate fitting output and to control fitting if the algorithm doesn’t converge which happens infrequently. The list values include:

- showit Integer (0-3, default 0) controls the (increasing)amount of information printed during fitting. 0 - none, >=1 - information about refitting and bound changes is printed, >=2 - information about adjustment term fitting is printed, ==3 -per-iteration parameter estimates and log-likelihood printed.
estimate if FALSE fits model but doesn’t estimate predicted probabilities
refit if TRUE the algorithm will attempt multiple optimizations at different starting values if it
doesn’t converge
nrefits number of refitting attempts
initial a named list of starting values for the parameters (e.g. $scale, $shape, $adjustment)
lowerbounds a vector of lowerbounds for the parameters
upperbounds a vector of upperbounds for the parameters
limit if TRUE restrict analysis to observations with detected=1
debug if TRUE, if fitting fails, return an object with fitting information
nofit if TRUE don’t fit a model, but use the starting values and generate an object based on those
values
optimx.method one (or a vector of) string(s) giving the optimisation method to use. If more than
one is supplied, the results from one are used as the starting values for the next. See optimx
optimx.maxit maximum number of iterations to use in the optimisation.
silent silences warnings within ds fitting method (helpful for running many times without gener-
ating many warning/error messages).

Examples of distance sampling analyses are available at http://examples.distancesampling.org/.

Value

model object of class=(method, "ddf")

Author(s)

Jeff Laake

References


See Also

ddf.ds, ddf.io, ddf.io.fi, ddf.trial, ddf.trial.fi, ddf.rem, ddf.rem.fi, mrds-opt
Examples

# load data
data(book.tee.data)
region <- book.tee.data$book.tee.region
egdata <- book.tee.data$book.tee.dataframe
samples <- book.tee.data$book.tee.samples
obs <- book.tee.data$book.tee.obs

# fit a half-normal detection function
result <- ddf(dsmodel=~mcds(key="hn", formula=~1), data=egdata, method="ds",
             meta.data=list(width=4))

# fit an independent observer model with full independence
result.io.fi <- ddf(mrmodel=~glm(~distance), data=egdata, method="io.fi",
                   meta.data=list(width = 4))

# fit an independent observer model with point independence
result.io <- ddf(dsmodel=~cds(key = "hn"), mrmodel=~glm(~distance),
                 data=egdata, method="io", meta.data=list(width=4))

## Not run:

# simulated single observer point count data (see ?ptdata.single)
data(ptdata.single)
ptdata.single$distbegin <- (as.numeric(cut(ptdata.single$distance,10*(0:10)))-1)*10
ptdata.single$distend <- (as.numeric(cut(ptdata.single$distance,10*(0:10))))*10
model <- ddf(data=ptdata.single, dsmodel=~cds(key="hn"),
             meta.data=list(point=TRUE,binned=TRUE,breaks=10*(0:10)))

summary(model)
plot(model,main="Single observer binned point data - half normal")

model <- ddf(data=ptdata.single, dsmodel=~cds(key="hr"),
             meta.data=list(point=TRUE, binned=TRUE, breaks=10*(0:10)))

summary(model)
plot(model,main="Single observer binned point data - hazard rate")

dev.new()

# simulated double observer point count data (see ?ptdata.dual)
# setup data
data(ptdata.dual)
ptdata.dual$distbegin <- (as.numeric(cut(ptdata.dual$distance,10*(0:10)))-1)*10
ptdata.dual$distend <- (as.numeric(cut(ptdata.dual$distance,10*(0:10))))*10
model <- ddf(method="io", data=ptdata.dual, dsmodel=~cds(key="hn"),
             mrmmodel=~glm(formula=~distance*observer),
             meta.data=list(point=TRUE, binned=TRUE, breaks=10*(0:10)))

summary(model)
```
plot(model, main="Dual observer binned point data", new=FALSE, pages=1)

model <- ddf(method="io", data=ptdata.dual,
        dsmodel=~cds(key="unif", adj.series="cos", adj.order=1),
        mrmodel=~glm(formula=~distance*observer),
        meta.data=list(point=TRUE, binned=TRUE, breaks=10*(0:10)))

summary(model)

par(mfrow=c(2,3))
plot(model,main="Dual observer binned point data",new=FALSE)

## End(Not run)
```

ddf.ds  

_CDS/MCDS Distance Detection Function Fitting_

**Description**

Fits a conventional distance sampling (CDS) (likelihood eq 6.6 in Laake and Borchers 2004) or multi-covariate distance sampling (MCDS) (likelihood eq 6.14 in Laake and Borchers 2004) model for the detection function of observed distance data. It only uses key functions and does not incorporate adjustment functions as in CDS/MCDS analysis engines in DISTANCE (Marques and Buckland 2004). Distance can be grouped (binned), ungrouped (unbinned) or mixture of the two. This function is not called directly by the user and is called from ddf, ddf.io, or ddf.trial.

**Usage**

```r
## S3 method for class 'ds'
ddf(model, data, meta.data = list(), control = list(), call, method = "ds")
```

**Arguments**

- **model**: model list with key function and scale formula if any
- **data**: analysis dataframe
- **meta.data**: list containing settings controlling data structure
- **control**: list containing settings controlling model fitting
- **call**: original function call if this function not called directly from ddf (e.g., called via ddf.io
- **method**: analysis method; only needed if this function called from ddf.io or ddf.trial

**Details**

For a complete description of each of the calling arguments, see _ddf_. The argument _model_ in this function is the same as _dsmodel_ in _ddf_. The argument _dataname_ is the name of the dataframe specified by the argument _data_ in _ddf_. The arguments _control, meta.data, and method_ are defined the same as in _ddf_.

Value

result: a ds model object

Note

If mixture of binned and unbinned distance, width must be set to be >= largest interval endpoint; this could be changed with a more complicated analysis; likewise, if all binned and bins overlap, the above must also hold; if bins don’t overlap, width must be one of the interval endpoints; same holds for left truncation Although the mixture analysis works in principle it has not been tested via simulation.

Author(s)

Jeff Laake

References


See Also

flnl, summary.ds, coef.ds, plot.ds, gof.ds

Examples

# ddf.ds is called when ddf is called with method="ds"

data(book.tee.data)
region <- book.tee.data$book.tee.region
egdata <- book.tee.data$book.tee.dataframe
samples <- book.tee.data$book.tee.samples
obs <- book.tee.data$book.tee.obs
result <- ddf(dsmodel = ~mcds(key = "hn", formula = ~1),
              data = egdata[egdata$observer==1, ], method = "ds",
              meta.data = list(width = 4))
summary(result, se=TRUE)
plot(result, main="cds - observer 1")
print(dht(result, region, samples, obs, options=list(varflag=0, group=TRUE),
         se=TRUE))
print(ddf.gof(result))
Goodness of fit tests for distance sampling models

Description

Generic function that computes chi-square goodness of fit test for detection function models with binned data and Cramer-von Mises and Kolmogorov-Smirnov (if ks=TRUE) tests for exact distance data. By default a Q-Q plot is generated for exact data (and can be suppressed using the qq=FALSE argument).

Usage

```r
ddf.gof(
  model,
  breaks = NULL,
  nc = NULL,
  qq = TRUE,
  nboot = 100,
  ks = FALSE,
  ...
)
```

Arguments

- `model`: model object
- `breaks`: Cutpoints to use for binning data
- `nc`: Number of distance classes
- `qq`: Flag to indicate whether quantile-quantile plot is desired
- `nboot`: number of replicates to use to calculate p-values for the Kolmogorov-Smirnov goodness of fit test statistics
- `ks`: perform the Kolmogorov-Smirnov test (this involves many bootstraps so can take a while)
- `...`: Graphics parameters to pass into `qqplot` function

Details

Formal goodness of fit testing for detection function models using Kolmogorov-Smirnov and Cramer-von Mises tests. Both tests are based on looking at the quantile-quantile plot produced by `qqplot.ddf` and deviations from the line x=y.

The Kolmogorov-Smirnov test asks the question "what's the largest vertical distance between a point and the y=x line?" It uses this distance as a statistic to test the null hypothesis that the samples (EDF and CDF in our case) are from the same distribution (and hence our model fits well). If the deviation between the y=x line and the points is too large we reject the null hypothesis and say the model doesn’t have a good fit.
Rather than looking at the single biggest difference between the y=x line and the points in the Q-Q plot, we might prefer to think about all the differences between line and points, since there may be many smaller differences that we want to take into account rather than looking for one large deviation. Its null hypothesis is the same, but the statistic it uses is the sum of the deviations from each of the point to the line.

Value

List of class `ddf.gof` containing

- **chi-square**: Goodness of fit test statistic
- **df**: Degrees of freedom associated with test statistic
- **p-value**: Significance level of test statistic

Details

Note that a bootstrap procedure is required for the Kolmogorov-Smirnov test to ensure that the p-values from the procedure are correct as the we are comparing the cumulative distribution function (CDF) and empirical distribution function (EDF) and we have estimated the parameters of the detection function. The `nboot` parameter controls the number of bootstraps to use. Set to 0 to avoid computing bootstraps (much faster but with no Kolmogorov-Smirnov results, of course).

Author(s)

Jeff Laake

See Also

- `qqplot.ddf`

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**Mark-Recapture Distance Sampling (MRDS) IO - PI**

Description

Mark-Recapture Distance Sampling (MRDS) Analysis of Independent Observer Configuration and Point Independence

Usage

```r
## S3 method for class 'io'
ddf(dsmodel, mrmodel, data, meta.data = list(), control = list(), call = "")
```
**Arguments**

- **dsmodel**: distance sampling model specification; model list with key function and scale formula if any
- **mrmodel**: mark-recapture model specification; model list with formula and link
- **data**: analysis dataframe
- **meta.data**: list containing settings controlling data structure
- **control**: list containing settings controlling model fitting
- **call**: original function call used to call `ddf`

**Details**

MRDS analysis based on point independence involves two separate and independent analyses of the mark-recapture data and the distance sampling data. For the independent observer configuration, the mark-recapture data are analysed with a call to `ddf.io.fi` (see likelihood eq 6.8 and 6.16 in Laake and Borchers 2004) to fit conditional distance sampling detection functions to estimate \( p(0) \), detection probability at distance zero for the independent observer team based on independence at zero (eq 6.22 in Laake and Borchers 2004). Independently, the distance data, the union of the observations from the independent observers, are used to fit a conventional distance sampling (CDS) (likelihood eq 6.6) or multi-covariate distance sampling (MCDS) (likelihood eq 6.14) model for the detection function, \( g(y) \), such that \( g(0)=1 \). The detection function for the observer team is then created as \( p(y)=p(0)\cdot g(y) \) (eq 6.28 of Laake and Borchers 2004) from which predictions are made. `ddf.io` is not called directly by the user and is called from `ddf` with `method``io``.``

For a complete description of each of the calling arguments, see `ddf`. The argument `dataname` is the name of the dataframe specified by the argument `data` in `ddf`. The arguments `dsmodel`, `mrmodel`, `control` and `meta.data` are defined the same as in `ddf`.

**Value**

- **result**: an io model object which is composed of io.fi and ds model objects

**Author(s)**

Jeff Laake

**References**


**See Also**

- `ddf.io.fi`, `ddf.ds`, `summary.io`, `coef.io`, `plot.io`, `gof.io`
Description

Mark-Recapture Analysis of Independent Observer Configuration with Full Independence

Usage

```r
## S3 method for class 'io.fi'
ddf(model, data, meta.data = list(), control = list(), call = "", method)
```

Arguments

- `model`: mark-recapture model specification
- `data`: analysis dataframe
- `meta.data`: list containing settings controlling data structure
- `control`: list containing settings controlling model fitting
- `call`: original function call used to call `ddf`
- `method`: analysis method; only needed if this function called from `ddf.io`

Details

The mark-recapture data derived from an independent observer distance sampling survey can be used to derive conditional detection functions \( p_j(y) \) for both observers \( j=1,2 \). They are conditional detection functions because detection probability for observer \( j \) is based on seeing or not seeing observations made by observer \( 3-j \). Thus, \( p_1(y) \) is estimated by \( p_{1|2}(y) \).

If detections by the observers are independent (full independence) then \( p_1(y)=p_{1|2}(y), p_2(y)=p_{2|1}(y) \) and for the union, full independence means that \( p(y)=p_1(y) + p_2(y) - p_{1|2}(y) \) for each distance \( y \). In fitting the detection functions the likelihood given by eq 6.8 and 6.16 in Laake and Borchers (2004) is used. That analysis does not require the usual distance sampling assumption that perpendicular distances are uniformly distributed based on line placement that is random relative to animal distribution. However, that assumption is used in computing predicted detection probability which is averaged based on a uniform distribution (see eq 6.11 of Laake and Borchers 2004).

For a complete description of each of the calling arguments, see `ddf`. The argument `model` in this function is the same as `mrmodel` in `ddf`. The argument `dataname` is the name of the dataframe specified by the argument `data` in `ddf`. The arguments `control`, `meta.data`, and `method` are defined the same as in `ddf`.

Value

- `result`: an io.fi model object

Author(s)

Jeff Laake
Mark-Recapture Distance Sampling (MRDS) Removal - PI

Description
Mark-Recapture Distance Sampling (MRDS) Analysis of Removal Observer Configuration and Point Independence

Usage
```r
## S3 method for class 'rem'
ddf(dsmodel, mrmodel, data, meta.data = list(), control = list(), call = "")
```

Arguments
- `dsmodel`: distance sampling model specification; model list with key function and scale formula if any
- `mrmodel`: mark-recapture model specification; model list with formula and link
- `data`: analysis dataframe
- `meta.data`: list containing settings controlling data structure
- `control`: list containing settings controlling model fitting
- `call`: original function call used to call `ddf`

Details
MRDS analysis based on point independence involves two separate and independent analyses of the mark-recapture data and the distance sampling data. For the removal observer configuration, the mark-recapture data are analysed with a call to `ddf.rem.fi` (see Laake and Borchers 2004) to fit conditional distance sampling detection functions to estimate \( p(0) \), detection probability at distance zero for the primary observer based on independence at zero (eq 6.22 in Laake and Borchers 2004). Independently, the distance data, the observations from the primary observer, are used to fit a conventional distance sampling (CDS) (likelihood eq 6.6) or multi-covariate distance sampling (MCDS) (likelihood eq 6.14) model for the detection function, \( g(y) \), such that \( g(0)=1 \). The detection function for the primary observer is then created as \( p(y)=p(0)*g(y) \) (eq 6.28 of Laake and Borchers 2004) from which predictions are made. `ddf.rem` is not called directly by the user and is called from `ddf` with method="rem".
For a complete description of each of the calling arguments, see `ddf`. The argument `data` is the dataframe specified by the argument `data` in `ddf`. The arguments `dsmodel`, `mrmodel`, `control` and `meta.data` are defined the same as in `ddf`.

**Value**

result: an `rem` model object which is composed of `rem.fi` and `ds` model objects

**Author(s)**

Jeff Laake

**References**


**See Also**

`ddf.rem.fi`, `ddf.ds`

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**Description**

Mark-Recapture Distance Sampling (MRDS) Analysis of Removal Observer Configuration with Full Independence

**Usage**

```r
## S3 method for class 'rem.fi'
ddf(model, data, meta.data = list(), control = list(), call = "", method)
```

**Arguments**

- `model`: mark-recapture model specification
- `data`: analysis dataframe
- `meta.data`: list containing settings controlling data structure
- `control`: list containing settings controlling model fitting
- `call`: original function call used to call `ddf`
- `method`: analysis method; only needed if this function called from `ddf.io`
Details

The mark-recapture data derived from an removal observer distance sampling survey can only derive conditional detection functions \( p_j(y) \) for both observers \( (j=1) \) because technically it assumes that detection probability does not vary by occasion (observer in this case). It is a conditional detection function because detection probability for observer 1 is conditional on the observations seen by either of the observers. Thus, \( p_1(y) \) is estimated by \( p_{1|2}(y) \).

If detections by the observers are independent (full independence) then \( p_1(y) = p_{1|2}(y) \) and for the union, full independence means that \( p(y) = p_1(y) + p_2(y) - p_1(y)*p_2(y) \) for each distance \( y \). In fitting the detection functions the likelihood from Laake and Borchers (2004) are used. That analysis does not require the usual distance sampling assumption that perpendicular distances are uniformly distributed based on line placement that is random relative to animal distribution. However, that assumption is used in computing predicted detection probability which is averaged based on a uniform distribution (see eq 6.11 of Laake and Borchers 2004).

For a complete description of each of the calling arguments, see `ddf`. The argument `model` in this function is the same as `mrmodel` in `ddf`. The argument `dataname` is the name of the dataframe specified by the argument `data` in `ddf`. The arguments `control`, `meta.data`, and `method` are defined the same as in `ddf`.

Value

result: an rem.fi model object

Author(s)

Jeff Laake

References


See Also

ddf.io, rem.glm
Arguments

dsmode | distance sampling model specification; model list with key function and scale formula if any
mrmodel | mark-recapture model specification; model list with formula and link
data | analysis data.frame
meta.data | list containing settings controlling data structure
control | list containing settings controlling model fitting
call | original function call used to call ddf

Details

MRDS analysis based on point independence involves two separate and independent analyses of the mark-recapture data and the distance sampling data. For the trial configuration, the mark-recapture data are analysed with a call to ddf.trial.fi (see likelihood eq 6.12 and 6.17 in Laake and Borchers 2004) to fit a conditional distance sampling detection function for observer 1 based on trials (observations) from observer 2 to estimate $p_1(0)$, detection probability at distance zero for observer 1. Independently, the distance data from observer 1 are used to fit a conventional distance sampling (CDS) (likelihood eq 6.6) or multi-covariate distance sampling (MCDS) (likelihood eq 6.14) model for the detection function, $g(y)$, such that $g(0)=1$. The detection function for observer 1 is then created as $p_1(y)=p_1(0)g(y)$ (eq 6.28 of Laake and Borchers 2004) from which predictions are made. ddf.trial is not called directly by the user and is called from ddf with method="trial".

For a complete description of each of the calling arguments, see ddf. The argument dataname is the name of the dataframe specified by the argument data in ddf. The arguments dsmode, mrmodel, control and meta.data are defined the same as in ddf.

Value

result: a trial model object which is composed of trial.fi and ds model objects

Author(s)

Jeff Laake

References


See Also

ddf.trial.fi, ddf.ds, summary.trial, coef.trial, plot.trial, gof.trial
Description

Mark-Recapture Analysis of Trial Observer Configuration with Full Independence

Usage

```r
## S3 method for class 'trial.fi'
ddf(model, data, meta.data = list(), control = list(), call = "", method)
```

Arguments

- `model`: mark-recapture model specification
- `data`: analysis dataframe
- `meta.data`: list containing settings controlling data structure
- `control`: list containing settings controlling model fitting
- `call`: original function call used to call `ddf`
- `method`: analysis method; only needed if this function called from `ddf.trial`

Details

The mark-recapture data derived from a trial observer distance sampling survey can be used to derive a conditional detection function ($p_1(y)$) for observer 1 based on trials (observations) from observer 2. It is a conditional detection function because detection probability for observer 1 is based on seeing or not seeing observations made by observer 2. Thus, $p_1(y)$ is estimated by $p_{1|2}(y)$. If detections by the observers are independent (full independence) then $p_1(y) = p_{1|2}(y)$ for each distance $y$. In fitting the detection functions the likelihood given by eq 6.12 or 6.17 in Laake and Borchers (2004) is used. That analysis does not require the usual distance sampling assumption that perpendicular distances are uniformly distributed based on line placement that is random relative to animal distribution. However, that assumption is used in computing predicted detection probability which is averaged based on a uniform distribution (see eq 6.13 of Laake and Borchers 2004).

For a complete description of each of the calling arguments, see `ddf`. The argument `model` in this function is the same as `mrmodel` in `ddf`. The argument `data.name` is the name of the dataframe specified by the argument `data` in `ddf`. The arguments `control,meta.data,method` are defined the same as in `ddf`.

Value

- `result`: a `trial.fi` model object

Author(s)

Jeff Laake
References


See Also
ddf.trial, summary.trial.fi, coef.trial.fi, plot.trial.fi, gof.trial.fi

DeltaMethod

**Numeric Delta Method approximation for the variance-covariance matrix**

**Description**

Computes delta method variance-covariance matrix of results of any generic function `fct` that computes a vector of estimates as a function of a set of estimated parameters `par`.

**Usage**

```r
DeltaMethod(par, fct, vcov, delta, ...)
```

**Arguments**

- `par`: vector of parameter values at which estimates should be constructed
- `fct`: function that constructs estimates from parameters `par`
- `vcov`: variance-covariance matrix of the parameters
- `delta`: proportional change in parameters used to numerically estimate first derivative with central-difference formula
- `...`: any additional arguments needed by `fct`

**Details**

The delta method (aka propagation of errors is based on Taylor series approximation - see Seber’s book on Estimation of Animal Abundance). It uses the first derivative of `fct` with respect to `par` which is computed in this function numerically using the central-difference formula. It also uses the variance-covariance matrix of the estimated parameters which is derived in estimating the parameters and is an input argument.

The first argument of `fct` should be `par` which is a vector of parameter estimates. It should return a single value (or vector) of estimate(s). The remaining arguments of `fct` if any can be passed to `fct` by including them at the end of the call to `DeltaMethod` as name=value pairs.
**det.tables**

**Value**

- a list with values
  - variance: estimated variance-covariance matrix of estimates derived by `fct`
  - partial: matrix (or vector) of partial derivatives of `fct` with respect to the parameters `par`

**Note**

This is a generic function that can be used in any setting beyond the `mrds` package. However this is an internal function for `mrds` and the user does not need to call it explicitly.

**Author(s)**

Jeff Laake

---

**det.tables**  
*Observation detection tables*

**Description**

Creates a series of tables for dual observer data that shows the number missed and detected for each observer within defined distance classes.

**Usage**

```r
det.tables(model, nc = NULL, breaks = NULL)
```

**Arguments**

- `model`: fitted model from `ddf`
- `nc`: number of equal-width bins for histogram
- `breaks`: user defined breakpoints

**Value**

- list object of class "det.tables"
  - Observer1: table for observer 1
  - Observer2: table for observer 2
  - Duplicates: histogram counts for duplicates
  - Pooled: histogram counts for all observations by either observer
  - Obs1_2: table for observer 1 within subset seen by observer 2
  - Obs2_1: table for observer 2 within subset seen by observer 1
Author(s)

Jeff Laake

Examples

data(book.tee.data)
region<-book.tee.data$book.tee.region
egdata<-book.tee.data$book.tee.dataframe
samples<-book.tee.data$book.tee.samples
obs<-book.tee.data$book.tee.obs
xx=ddf(mrmodel=~glm(formula=~distance*observer),
    dsmodel = ~mcds(key = "hn", formula = "sex"), data = egdata, method = "io",
    meta.data = list(width = 4))
tabs=det.tables(xx,breaks=c(0,.5,1,2,3,4))
par(mfrow=c(2,2))
plot(tabs,new=FALSE,which=c(1,2,5,6))

detfct.fit

Fit detection function using key-adjustment functions

Description

Fit detection function to observed distances using the key-adjustment function approach. If adjustment functions are included it will alternate between fitting parameters of key and adjustment functions and then all parameters much like the approach in the CDS and MCDS DISTANCE FORTRAN code. To do so it calls detfct.fit.opt which uses the R optim function which does not allow non-linear constraints so inclusion of adjustments does allow the detection function to be non-monotone.

Usage

detfct.fit(ddfobj, optim.options, bounds, misc.options)

Arguments

ddfobj detection function object
optim.options control options for optim
bounds bounds for the parameters
misc.options miscellaneous options
Value

fitted detection function model object with the following list structure

- **par**: final parameter vector
- **value**: final negative log likelihood value
- **counts**: number of function evaluations
- **convergence**: see codes in optim
- **message**: string about convergence
- **hessian**: hessian evaluated at final parameter values
- **aux**: a list with 20 elements
  - maxit: maximum number of iterations allowed for optimization
  - lower: lower bound values for parameters
  - upper: upper bound values for parameters
  - setlower: TRUE if they are user set bounds
  - setupper: TRUE if they are user set bounds
  - point: TRUE if point counts and FALSE if line transect
  - int.range: integration range values
  - showit: integer value that determines information printed during iteration
  - silent: option to silence errors from detfct.fit.opt
  - integral.numeric if TRUE compute logistic integrals numerically
  - breaks: breaks in distance for defined fixed bins for analysis
  - maxiter: maximum iterations used
  - refit: if TRUE, detection function will be fitted more than once if parameters are at a boundary or when convergence is not achieved
  - nrefits: number of refittings
  - mono: if TRUE monotonicity will be enforced
  - mono.strict: if TRUE, then strict monotonicity is enforced; otherwise weak
  - width: radius of point count or half-width of strip
  - standardize: if TRUE, detection function is scaled so g(0)=1
  - ddfobj: distance detection function object; see `create.ddfobj`
  - bounded: TRUE if parameters ended up a boundary (I think)
  - model: list of formulas for detection function model (probably can remove this)

Author(s)

Dave Miller; Jeff Laake
**detfct.fit.opt**

*Fit detection function using key-adjustment functions*

**Description**

Fit detection function to observed distances using the key-adjustment function approach. If adjustment functions are included it will alternate between fitting parameters of key and adjustment functions and then all parameters much like the approach in the CDS and MCDS Distance FORTRAN code. This function is called by the driver function `detfct.fit`, then calls `optimx` function.

**Usage**

`detfct.fit.opt(ddfobj, optim.options, bounds, misc.options, fitting = "all")`

**Arguments**

- **ddfobj**: detection function object
- **optim.options**: control options for `optim`
- **bounds**: bounds for the parameters
- **misc.options**: miscellaneous options
- **fitting**: character string with values "all","key","adjust" to determine which parameters are allowed to vary in the fitting

**Value**

fitted detection function model object with the following list structure

- **par**: final parameter vector
- **value**: final negative log likelihood value
- **counts**: number of function evaluations
- **convergence**: see codes in `optim`
- **message**: string about convergence
- **hessian**: hessian evaluated at final parameter values
- **aux**: a list with 20 elements
  - maxit: maximum number of iterations allowed for optimization
  - lower: lower bound values for parameters
  - upper: upper bound values for parameters
  - setlower: TRUE if they are user set bounds
  - setupper: TRUE if they are user set bounds
  - point: TRUE if point counts and FALSE if line transect
  - int.range: integration range values
  - showit: integer value that determines information printed during iteration
  - integral.numeric if TRUE compute logistic integrals numerically
• breaks: breaks in distance for defined fixed bins for analysis
• maxiter: maximum iterations used
• refit: if TRUE, detection function will be fitted more than once if parameters
  are at a boundary or when convergence is not achieved
• nrefits: number of refittings
• mono: if TRUE, monotonicity will be enforced
• mono.strict: if TRUE, then strict monotonicity is enforced; otherwise weak
• width: radius of point count or half-width of strip
• standardize: if TRUE, detection function is scaled so g(0)=1
• ddfobj: distance detection function object; see create.ddfobj
• bounded: TRUE if parameters ended up a boundary (I think)
• model: list of formulas for detection function model (probably can remove
  this)

**Author(s)**

Dave Miller; Jeff Laake; Lorenzo Milazzo

---

**dht**

*Density and abundance estimates and variances*

**Description**

Compute density and abundance estimates and variances based on Horvitz-Thompson-like estim-

**Usage**

```r
  dht(
    model,
    region.table,
    sample.table,
    obs.table = NULL,
    subset = NULL,
    se = TRUE,
    options = list()
  )
```

**Arguments**

- `model` ddf model object
- `region.table` data.frame of region records. Two columns: Region.Label and Area.
obs.table: a data.frame of observation records with fields: object, Region.Label, and Sample.Label which give links to sample.table, region.table and the data records used in model. Not necessary if the data.frame used to create the model contains Region.Label, Sample.Label columns.

subset: a subset statement to create obs.table

se: if TRUE computes standard errors, coefficient of variation and confidence intervals (based on log-normal approximation). See "Uncertainty" below.

options: a list of options that can be set, see "dht options", below.

Details

Density and abundance within the sampled region is computed based on a Horvitz-Thompson-like estimator for groups and individuals (if a clustered population) and this is extrapolated to the entire survey region based on any defined regional stratification. The variance is based on replicate samples within any regional stratification. For clustered populations, \( E(s) \) and its standard error are also output.

Abundance is estimated with a Horvitz-Thompson-like estimator (Huggins 1989, 1991; Borchers et al 1998; Borchers and Burnham 2004). The abundance in the sampled region is simply \( \frac{1}{p_1} + \frac{1}{p_2} + \ldots + \frac{1}{p_n} \), where \( p_i \) is the estimated detection probability for the \( i \)th detection of \( n \) total observations. It is not strictly a Horvitz-Thompson estimator because the \( p_i \) are estimated and not known. For animals observed in tight clusters, that estimator gives the abundance of groups (\( group=TRUE \) in options) and the abundance of individuals is estimated as \( \frac{s_1}{p_1} + \frac{s_2}{p_2} + \ldots + \frac{s_n}{p_n} \), where \( s_i \) is the size (e.g., number of animals in the group) of each observation (\( group=FALSE \) in options).

Extrapolation and estimation of abundance to the entire survey region is based on either a random sampling design or a stratified random sampling design. Replicate samples (lines) are specified within regional strata region.table, if any. If there is no stratification, region.table should contain only a single record with the Area for the entire survey region. The sample.table is linked to the region.table with the Region.Label. The obs.table is linked to the sample.table with the Sample.Label and Region.Label. Abundance can be restricted to a subset (e.g., for a particular species) of the population by limiting the list the observations in obs.table to those in the desired subset. Alternatively, if Sample.Label and Region.Label are in the data.frame used to fit the model, then a subset argument can be given in place of the obs.table. To use the subset argument but include all of the observations, use subset=1==1 to avoid creating an obs.table.

In extrapolating to the entire survey region it is important that the unit measurements be consistent or converted for consistency. A conversion factor can be specified with the convert.units variable in the options list. The values of Area in region.table, must be made consistent with the units for Effort in sample.table and the units of distance in the data.frame that was analyzed. It is easiest to do if the units of Area is the square of the units of Effort and then it is only necessary to convert the units of distance to the units of Effort. For example, if Effort was entered in kilometres and Area in square kilometres and distance in metres then using options=list(convert.units=0.001) would convert metres to kilometres, density would be expressed in square kilometres which would then be consistent with units for Area. However, they can all be in different units as long as the appropriate composite value for convert.units is chosen. Abundance for a survey region can be expressed as: \( A*N/a \) where \( A \) is Area for the survey region, \( N \) is the abundance in the covered (sampled) region, and \( a \) is the area of the sampled region and is in units of Effort * distance. The sampled region \( a \) is multiplied by convert.units,
so it should be chosen such that the result is in the same units of Area. For example, if Effort was entered in kilometres, Area in hectares (100m x 100m) and distance in metres, then using options=list(convert.units=10) will convert a to units of hectares (100 to convert metres to 100 metres for distance and .1 to convert km to 100m units).

The argument options is a list of variable=value pairs that set options for the analysis. All but one of these has been described so far. The remaining variable pdelta should not need to be changed but was included for completeness. It controls the precision of the first derivative calculation for the delta method variance.

**Value**

list object of class dht with elements:

- clusters: result list for object clusters
- individuals: result list for individuals
- Expected.S: data.frame of estimates of expected cluster size with fields Region, Expected.S and se.Expected.S If each cluster size=1, then the result only includes individuals and not clusters and Expected.S.

The list structure of clusters and individuals are the same:

- bysample: data.frame giving results for each sample; Nhat is the estimated abundance within the sample and Nhat is scaled by surveyed area/covered area within that region
- summary: data.frame of summary statistics for each region and total
- N: data.frame of estimates of abundance for each region and total
- D: data.frame of estimates of density for each region and total
- average.p: average detection probability estimate
- cormat: correlation matrix of regional abundance/density estimates and total (if more than one region)
- vc: list of 3: total variance-covariance matrix, detection function component of variance and encounter rate component of variance. For detection the v-c matrix and partial vector are returned

Nhat.by.sample: another summary of Nhat by sample used by dht.se

**Uncertainty**

If the argument se=TRUE, standard errors for density and abundance is computed. Coefficient of variation and log-normal confidence intervals are constructed using a Satterthwaite approximation for degrees of freedom (Buckland et al. 2001 p. 90). The function dht.se computes the variance and interval estimates.

The variance has two components:

- variation due to uncertainty from estimation of the detection function parameters;
- variation in abundance due to random sample selection;
The first component (model parameter uncertainty) is computed using a delta method estimate of variance (Huggins 1989, 1991, Borchers et al. 1998) in which the first derivatives of the abundance estimator with respect to the parameters in the detection function are computed numerically (see \texttt{DeltaMethod}).

The second component (encounter rate variance) can be computed in one of several ways depending on the form taken for the encounter rate and the estimator used. To begin with there are three possible values for \texttt{varflag} to calculate encounter rate:

- \( 0 \) uses a binomial variance for the number of observations (equation 13 of Borchers et al. 1998). This estimator is only useful if the sampled region is the survey region and the objects are not clustered; this situation will not occur very often;
- \( 1 \) uses the encounter rate \( n/L \) (objects observed per unit transect) from Buckland et al. (2001) pg 78-79 (equation 3.78) for line transects (see also Fewster et al, 2009 estimator R2). This variance estimator is not appropriate if size or a derivative of size is used in the detection function;
- \( 2 \) is the default and uses the encounter rate estimator \( \hat{N}/L \) (estimated abundance per unit transect) suggested by Innes et al (2002) and Marques & Buckland (2004).

In general if any covariates are used in the models, the default \( \texttt{varflag}=2 \) is preferable as the estimated abundance will take into account variability due to covariate effects. If the population is clustered the mean group size and standard error is also reported.

For options 1 and 2, it is then possible to choose one of the estimator forms given in Fewster et al (2009) for line transects: "R2", "R3", "R4", "S1", "S2", "O1", "O2" or "O3" by specifying the \texttt{ervar=} option (default "R2"). For points estimator "P3" is the only option. See \texttt{varn} and Fewster et al (2009) for further details on these estimators.

dht \texttt{options}

Several options are available to control calculations and output:

- \texttt{ci.width} Confidence interval width, expressed as a decimal between 0 and 1 (default 0.95, giving a 95% CI)
- \texttt{pdelta} delta value for computing numerical first derivatives (Default: 0.001)
- \texttt{varflag} 0,1,2 (see "Uncertainty") (Default: 2)
- \texttt{convert.units} multiplier for width to convert to units of length (Default: 1)
- \texttt{ervar} encounter rate variance type (see "Uncertainty" and type argument of \texttt{varn}). (Default: "R2" for lines and "P3" for points)

\textbf{Author(s)}

Jeff Laake, David L Miller

\textbf{References}


See Also

print.dht dht.se

dht.deriv

Computes abundance estimates at specified parameter values using Horvitz-Thompson-like estimator

Description

Computes abundance at specified values of parameters for numerical computation of first derivative with respect to parameters in detection function. An internal function called by DeltaMethod which is invoked by dht.se

Usage

dht.deriv(par, model, obs, samples, options = list())

Arguments

par  detection function parameter values
model  ddf model object
obs  observations table
samples  samples table
options  list of options as specified in dht
Value

vector of abundance estimates at values of parameters specified in par

Note

Internal function; not intended to be called by user

Author(s)

Jeff Laake

See Also

dht, dht.se, DeltaMethod

dht.se

Variance and confidence intervals for density and abundance estimates

Description

Computes standard error, CV, and log-normal confidence intervals for abundance and density within each region (if any) and for the total of all the regions. It also produces the correlation matrix for regional and total estimates.

Usage

dht.se(
  model,
  region.table,
  samples,
  obs,
  options,
  numRegions,
  estimate.table,
  Nhat.by.sample
)

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>model</td>
<td>ddf model object</td>
</tr>
<tr>
<td>region.table</td>
<td>table of region values</td>
</tr>
<tr>
<td>samples</td>
<td>table of samples(replicates)</td>
</tr>
<tr>
<td>obs</td>
<td>table of observations</td>
</tr>
<tr>
<td>options</td>
<td>list of options that can be set (see dht)</td>
</tr>
<tr>
<td>numRegions</td>
<td>number of regions</td>
</tr>
<tr>
<td>estimate.table</td>
<td>table of estimate values</td>
</tr>
<tr>
<td>Nhat.by.sample</td>
<td>estimated abundances by sample</td>
</tr>
</tbody>
</table>
Details

The variance has two components:

- variation due to uncertainty from estimation of the detection function parameters;
- variation in abundance due to random sample selection;

The first component (model parameter uncertainty) is computed using a delta method estimate of variance (Huggins 1989, 1991, Borchers et al. 1998) in which the first derivatives of the abundance estimator with respect to the parameters in the detection function are computed numerically (see `DeltaMethod`).

The second component (encounter rate variance) can be computed in one of several ways depending on the form taken for the encounter rate and the estimator used. To begin with there three possible values for `varflag` to calculate encounter rate:

- 0 uses a binomial variance for the number of observations (equation 13 of Borchers et al. 1998). This estimator is only useful if the sampled region is the survey region and the objects are not clustered; this situation will not occur very often;
- 1 uses the encounter rate $n/L$ (objects observed per unit transect) from Buckland et al. (2001) pg 78-79 (equation 3.78) for line transects (see also Fewster et al, 2009 estimator R2). This variance estimator is not appropriate if `size` or a derivative of `size` is used in the detection function;
- 2 is the default and uses the encounter rate estimator $\hat{N}/L$ (estimated abundance per unit transect) suggested by Innes et al (2002) and Marques & Buckland (2004).

In general if any covariates are used in the models, the default `varflag=2` is preferable as the estimated abundance will take into account variability due to covariate effects. If the population is clustered the mean group size and standard error is also reported.

For options 1 and 2, it is then possible to choose one of the estimator forms given in Fewster et al (2009). For line transects: "R2", "R3", "R4", "S1", "S2", "O1", "O2" or "O3" can be used by specifying the `ervar=` option (default "R2"). For point transects only the "P3" estimator may be used. See `varn` and Fewster et al (2009) for further details on these estimators.

Exceptions to the above occur if there is only one sample in a stratum. In that case it uses Poisson assumption ($\text{Var}(x) = x$) and it assumes a known variance so $z = 1.96$ is used for critical value.

In all other cases the degrees of freedom for the $t$-distribution assumed for the log(abundance) or log(density) is based on the Satterthwaite approximation (Buckland et al. 2001 pg 90) for the degrees of freedom (df). The df are weighted by the squared cv in combining the two sources of variation because of the assumed log-normal distribution because the components are multiplicative. For combining df for the sampling variance across regions they are weighted by the variance because it is a sum across regions.

A non-zero correlation between regional estimates can occur from using a common detection function across regions. This is reflected in the correlation matrix of the regional and total estimates which is given in the value list. It is only needed if subtotals of regional estimates are needed.

Value

List with 2 elements:

- `estimate.table` completed table with se, cv and confidence limits
- `vc` correlation matrix of estimates
Note

This function is called by dht and it is not expected that the user will call this function directly but it is documented here for completeness and for anyone expanding the code or using this function in their own code.

Author(s)

Jeff Laake

References

see dht

See Also

dht, print.dht

ds.function

Distance Sampling Functions

Description

Computes values of conditional and unconditional detection functions and probability density functions for line/point data for single observer or dual observer in any of the 3 configurations (io, trial, rem).

Usage

ds.function(
  model,
  newdata = NULL,
  obs = "All",
  conditional = FALSE,
  pdf = TRUE,
  finebr
)

Arguments

model model object
newdata dataframe at which to compute values; if NULL uses fitting data
obs 1 or 2 for observer 1 or 2, 3 for duplicates, "." for combined and "All" to return all of the values
conditional if FALSE, computes p(x) based on distance detection function and if TRUE based on mr detection function
pdf if FALSE, returns p(x) and if TRUE, returns p(x)*pi(x)/integral p(x)*pi(x)
finebr fine break values over which line is averaged
Details

Placeholder – Not functional —-

Value

List containing

- **xgrid**: grid of distance values
- **values**: average detection fct values at the xgrid values

Author(s)

Jeff Laake

---

flnl

*Log-likelihood computation for distance sampling data*

---

Description

For a specific set of parameter values, it computes and returns the negative log-likelihood for the distance sampling likelihood for distances that are unbinned, binned and a mixture of both. The function `flnl` is the function minimized using `optim` from within `ddf.ds`.

Usage

```
flnl(fpar, ddfobj, misc.options, fitting = "all")
```

Arguments

- **fpar**: parameter values for detection function at which negative log-likelihood should be evaluated
- **ddfobj**: distance sampling object
- **misc.options**: a list with the following elements: width transect width; int.range the integration range for observations; showit 0 to 3 controls level debug output; integral.numeric if TRUE integral is computed numerically rather than analytically; point is this a point transect?
- **fitting**: character "key" if only fitting key function parameters, "adjust" if fitting adjustment parameters or "all" to fit both
Details

Most of the computation is in `flt.lnl` in which the negative log-likelihood is computed for each observation. `flnlnl` is a wrapper that optionally outputs intermediate results and sums the individual log-likelihood values.

`flnlnl` is the main routine that manipulates the parameters using `getpar` to handle fitting of key, adjustment or all of the parameters. It then calls `flpt.lnl` to do the actual computation of the likelihood. The probability density function for point counts is \( f_r \) and for line transects is \( f_x \). \( f_x = g(x)/\mu \) (where \( g(x) \) is the detection function); whereas, \( f(r) = r^a g(r)/\mu \) where \( \mu \) in both cases is the normalizing constant. Both functions are in source code file for `link{detfct}` and are called from `distpdf` and the integral calculations are made with `integratepdf`.

Value

negative log-likelihood value at the parameter values specified in `fpar`

Note

These are internal functions used by `ddf.ds` to fit distance sampling detection functions. It is not intended for the user to invoke these functions but they are documented here for completeness.

Author(s)

Jeff Laake, David L Miller

See Also

`flt.var`, `detfct`

Description

Computes hessian to be used for variance-covariance matrix. The hessian is the outer product of the vector of first partials (see pg 62 of Buckland et al 2002).

Usage

```r
flt.var(ddfobj, misc.options)
```

Arguments

- `ddfobj` distance sampling object
- `misc.options` width-transect width (W); int.range-integration range for observations; showit-0 to 3 controls level of iteration printing; integral.numeric-if TRUE integral is computed numerically rather than analytically
Value

variance-covariance matrix of parameters in the detection function

Note

This is an internal function used by `ddf.ds` to fit distance sampling detection functions. It is not intended for the user to invoke this function but it is documented here for completeness.

Author(s)

Jeff Laake

References

Buckland et al. 2002

See Also

`flnl`, `flpt.lnl`, `ddf.ds`

g0

Compute value of p(0) using a logit formulation

Description

Compute value of p(0) using a logit formulation

Usage

g0(beta, z)

Arguments

beta    logistic parameters
z       design matrix of covariate values

Value

vector of p(0) values

Author(s)

Jeff Laake
getpar

Description

Extracts parameters of a particular type (scale, shape, adjustments or g0 (p(0))) from the vector of parameters in ddfobj. All of the parameters are kept in a single vector for optimization even though they have very different uses. assign.par parses them from the vector based on a known structure and assigns them into ddfobj. getpar extracts the requested types to be extracted from ddfobj.

Usage

getpar(ddfobj, fitting = "all", index = FALSE)

Arguments

  ddfobj distance sampling object (see create_ddfobj)
  fitting character string which is either "all","key","adjust" which determines which parameters are retrieved
  index logical that determines whether parameters are returned (FALSE) or starting indices in parameter vector for scale, shape, adjustment parameters

Value

  index==FALSE, vector of parameters that were requested or index==TRUE, vector of 3 indices for shape, scale, adjustment

Note

  Internal functions not intended to be called by user.

Author(s)

  Jeff Laake

See Also

  assign.par
gof.ds

Compute chi-square goodness-of-fit test for ds models

Description

Compute chi-square goodness-of-fit test for ds models

Usage

gof.ds(model, breaks = NULL, nc = NULL)

Arguments

model     ddf model object
breaks    distance cut points
nc         number of distance classes

Value

list with chi-square value, df and p-value

Author(s)

Jeff Laake

See Also

ddf.gof

---

gandalone

Integral of pdf of distances

Description

Computes the integral of distpdf with scale=1 (stdint=TRUE) or specified scale (stdint=FALSE).

Usage

`gstdin(x, ddfobj, index = NULL, select = NULL, width, standardize = TRUE,`
point = FALSE,
stdin = TRUE,
doeachint = FALSE,
left = left
)

Arguments

x lower, upper value for integration
ddfobj distance detection function specification
index specific data row index
select logical vector for selection of data values
width truncation width
standardize if TRUE, divide through by the function evaluated at 0
point logical to determine if point (TRUE) or line transect (FALSE)
stdin if TRUE, scale=1 otherwise specified scale used
doeachint if TRUE perform integration using integrate
left left truncation width

Value

vector of integral values of detection function

Note

This is an internal function that is not intended to be invoked directly.

Author(s)

Jeff Laake and David L Miller

Description

Takes bar heights (height) and cutpoints (breaks), and constructs a line-only histogram from them using the function plot() (if lineonly==FALSE) or lines() (if lineonly==TRUE).
Usage

histline(
  height,
  breaks,
  lineonly = FALSE,
  outline = FALSE,
  ylim = range(height),
  xlab = "x",
  ylab = "y",
  det.plot = FALSE,
  add = FALSE,
  ...
)

Arguments

  height        heights of histogram bars
  breaks        cutpoints for x
  lineonly      if TRUE, drawn with plot; otherwise with lines to allow addition of current plot
  outline       if TRUE, only outline of histogram is plotted
  ylim          limits for y axis
  xlab          label for x axis
  ylab          label for y axis
  det.plot      if TRUE, plot is of detection so yaxis limited to unit interval
  add           should this plot add to a previous window
  ...           Additional unspecified arguments for plot

Value

None

Author(s)

Jeff Laake and David L Miller

integratedetfct.logistic

Integrate a logistic detection function

Description

Integrates a logistic detection function; a separate function is used because in certain cases the integral can be solved analytically and also because the scale trick used with the half-normal and hazard rate doesn’t work with the logistic.
Usage

integratedetfct.logistic(x, scalemodel, width, theta1, integral.numeric, w)

Arguments

x                  logistic design matrix values
scalemodel         scale model for logistic
width               transect width
theta1             parameters for logistic
integral.numeric   if TRUE computes numerical integral value
w                   design covariates

Value

vector of integral values

Author(s)

Jeff Laake

integrate_logistic.analytic

Analytically integrate logistic detection function

Description

Computes integral (analytically) over x from 0 to width of a logistic detection function; For reference see integral #526 in CRC Std Math Table 24th ed

Usage

integrate_logistic.analytic(x, models, beta, width)

Arguments

x                    matrix of data
models               list of model formulae
beta                 parameters of logistic detection function
width                transect half-width

Author(s)

Jeff Laake
**integratepdf**

Numerically integrate pdf of observed distances over specified ranges

**Description**

Computes integral of pdf of observed distances over x for each observation. The method of computation depends on argument switches set and the type of detection function.

**Usage**

```r
integratepdf(
  ddfobj,
  select,
  width,
  int.range,
  standardize = TRUE,
  point = FALSE,
  left = 0,
  doeachint = FALSE
)
```

**Arguments**

- `ddfobj` distance detection function specification
- `select` logical vector for selection of data values
- `width` truncation width
- `int.range` integration range matrix; vector is converted to matrix
- `standardize` logical used to decide whether to divide through by the function evaluated at 0
- `point` logical to determine if point count (TRUE) or line transect (FALSE)
- `left` left truncation width
- `doeachint` calculate each integral numerically

**Value**

vector of integral values - one for each observation

**Author(s)**

Jeff Laake & Dave Miller
io.glm

Iterative offset GLM/GAM for fitting detection function

Description

Provides an iterative algorithm for finding the MLEs of detection (capture) probabilities for a two-occasion (double observer) mark-recapture experiment using standard algorithms GLM/GAM and an offset to compensate for conditioning on the set of observations. While the likelihood can be formulated and solved numerically, the use of GLM/GAM provides all of the available tools for fitting, predictions, plotting etc without any further development.

Usage

io.glm(
  datavec,  # dataframe
  fitformula,  # logit link formula
  eps = 1e-05,  # convergence criterion
  iterlimit = 500,  # maximum number of iterations allowed
  GAM = FALSE,  # uses GAM instead of GLM for fitting
  gamplot = TRUE  # set to TRUE to get a gam plot object if GAM=TRUE
)

Arguments

datavec dataframe
fitformula logit link formula
eps convergence criterion
iterlimit maximum number of iterations allowed
GAM uses GAM instead of GLM for fitting
gamplot set to TRUE to get a gam plot object if GAM=TRUE

Details

Note that currently the code in this function for GAMs has been commented out until the remainder of the mrds package will work with GAMs. This is an internal function that is used as by ddf.io.fi to fit mark-recapture models with 2 occasions. The argument mrmodel is used for fitformula.

Value

list of class("ioglm","glm","lm") or class("ioglm","gam")
glmobj GLM or GAM object
offsetvalue offsetvalues from iterative fit
plotobj gam plot object (if GAM & gamplot==TRUE, else NULL)
is.linear.logistic

Author(s)

Jeff Laake, David Borchers, Charles Paxton

References


Collection of functions for logistic detection functions

Description

These functions are used to test whether a logistic detection function is a linear function of distance (is.linear.logistic) or is constant (varies by distance but no other covariates) is.logistic.constant. Based on these tests, the most appropriate manner for integrating the detection function with respect to distance is chosen. The integrals are needed to estimate the average detection probability for a given set of covariates.

Usage

is.linear.logistic(xmat, g0model, zdim, width)

Arguments

xmat data matrix
g0model logit model
zdim number of columns in design matrix
width transect width

Details

If the logit is linear in distance then the integral can be computed analytically. If the logit is constant or only varies by distance then only one integral needs to be computed rather than an integral for each observation.

Value

Logical TRUE if condition holds and FALSE otherwise

Author(s)

Jeff Laake
is.logistic.constant  Is a logit model constant for all observations?

Description
Determines whether the specified logit model is constant for all observations. If it is constant then only one integral needs to be computed.

Usage
is.logistic.constant(xmat, g0model, width)

Arguments
xmat  data
g0model  logit model
width  transect width

Value
logical value

Author(s)
Jeff Laake

keyfct.th1  Threshold key function

Description
Threshold key function

Usage
keyfct.th1(distance, key.scale, key.shape)

Arguments
distance  perpendicular distance vector
key.scale  vector of scale values
key.shape  vector of shape values

Value
vector of probabilities
Threshold key function

**Description**
Threshold key function

**Usage**
```
keyfct.th2(distance, key.scale, key.shape)
```

**Arguments**
- `distance` : perpendicular distance vector
- `key.scale` : vector of scale values
- `key.shape` : vector of shape values

**Value**
vector of probabilities

Black-capped vireo mark-recapture distance sampling analysis

**Description**
These data represent avian point count surveys conducted at 453 point sample survey locations on the 24,000 (approx) live-fire region of Fort Hood in central Texas. Surveys were conducted by independent double observers (2 per survey occasion) and as such we had a maximum of 3 paired survey histories, giving a maximum of 6 sample occasions (see MacKenzie et al. 2006, MacKenzie and Royle 2005, and Laake et al. 2011 for various sample survey design details). At each point, we surveyed for 5 minutes (technically broken into 3 time intervals of 2, 2, and 1 minutes; not used here) and we noted detections by each observer and collected distance to each observation within a set of distance bins (0-25, 25-50, 50-75, 75-100m) of the target species (Black-capped vireo’s in this case) for each surveyor. Our primary focus was to use mark-recapture distance sampling methods to estimate density of Black-capped vireo’s, and to estimate detection rates for the mark-recapture, distance, and composite model.

**Format**
The format is a data frame with the following covariate metrics.

- **PointID** : Unique identifier for each sample location; locations are the same for both species
- **VisitNumber** : Visit number to the point
- **Species** : Species designation, either Golden-cheeked warbler (GW) or Black-capped Vireo (BV)
**Distance** Distance measure, which is either NA (representing no detection), or the median of the binned detection distances

**PairNumber** ID value indicating which observers were paired for that sampling occasion

**Observer** Observer ID, either primary(1), or secondary (2)

**Detected** Detection of a bird, either 1 = detected, or 0 = not detected

**Date** Date of survey since 15 march 2011

**Pred** Predicted occupancy value for that survey hexagon based on Farrell et al. (2013)

**Category** Region.Label categorization, see mrds help file for details on data structure

**Effort** Amount of survey effort at the point

**Day** Number of days since 15 March 2011

**ObjectID** Unique ID for each paired observations

### Details

In addition to detailing the analysis used by Collier et al. (2013, In Review), this example documents the use of mrds for avian point count surveys and shows how density models can be incorporated with occupancy models to develop spatially explicit density surface maps. For those that are interested, for the distance sampling portion of our analysis, we used both conventional distance sampling (cds) and multiple covariate distance sampling (mcds) with uniform and half-normal key functions. For the mark-recapture portion of our analysis, we tended to use covariates for distance (median bin width), observer, and date of survey (days since 15 March 2011).

We combined our mrds density estimates via a Horvitz-Thompson styled estimator with the resource selection function gradient developed in Farrell et al. (2013) and estimated density on an ~3.14ha hexagonal grid across our study area, which provided a density gradient for the Fort Hood military installation. Because there was considerable data manipulation needed for each analysis to structure the data appropriately for use in mrds, rather than wrap each analysis in a single function, we have provided both the Golden-cheeked warbler and Black-capped vireo analyses in their full detail. The primary differences you will see will be changes to model structures and model outputs between the two species.

### Author(s)

Bret Collier and Jeff Laake

### References


Examples

```r
## Not run:
data(lfbcvi)
xy=cut(lfbcvi$Pred, c(-0.0001, .1, .2, .3, .4, .5, .6, .7, .8, .9, 1),
  labels=c("1", "2", "3", "4", "5", "6", "7", "8", "9", "10"))
x=data.frame(lfbcvi, New=xy)

# Note that I scaled the individual covariate of day-helps with
# convergence issues
bird.data <- data.frame(object=x$ObjectID, observer=x$Observer,
  detected=x$Detected, distance=x$Distance,
  Region.Label=x$New, Sample.Label=x$PointID,
  Day=(x$Day/max(x$Day)))

# make observer a factor variable
bird.data$observer=factor(bird.data$observer)

# Jeff Laake suggested this snippet to quickly create distance medians
# which adds bin information to the bird.data dataframe
bird.data$distbegin=0
bird.data$distend=100
bird.data$distend[bird.data$distance==12.5]=25
bird.data$distend[bird.data$distance==37.5]=25
bird.data$distend[bird.data$distance==37.5]=50
bird.data$distbegin[bird.data$distance==62.5]=50
bird.data$distend[bird.data$distance==62.5]=75
bird.data$distbegin[bird.data$distance==87.5]=75
bird.data$distend[bird.data$distance==87.5]=100

# Removed all survey points with distance=NA for a survey event;
# hence no observations for use in ddf() but needed later
bird.data=bird.data[complete.cases(bird.data),]

# Manipulations on full dataset for various data.frame creation for
# use in density estimation using dht()
#Samples dataframe
xx=x
x=data.frame(PointID=x$PointID, Species=x$Species,
  Category=x$New, Effort=x$Effort)
x=x[!duplicated(x$PointID),]
point.num=table(x$Category)
samples=data.frame(PointID=x$PointID, Region.Label=x$Category,
  Effort=x$Effort)
final.samples=data.frame(Sample.Label=samples$PointID,
  Region.Label=samples$Region.Label,
  Effort=samples$Effort)

#obs dataframe
obs=data.frame(ObjectID=xx$ObjectID, PointID=xx$PointID)
```
#used to get Region and Sample assigned to ObjectID
obs=merge(obs, samples, by=c("PointID", "PointID"))
obs=obs[!duplicated(obs$ObjectID),]
obs=data.frame(object=obs$ObjectID, Region.Label=obs$Region.Label,
Sample.Label=obs$PointID)

region.data=data.frame(Region.Label=c(1, 2, 3, 4, 5, 6, 7, 8, 9, 10),
Area=c(point.num[1]*3.14, point.num[2]*3.14,
point.num[3]*3.14, point.num[4]*3.14,
point.num[5]*3.14, point.num[6]*3.14,
point.num[7]*3.14, point.num[8]*3.14,
point.num[9]*3.14, point.num[10]*3.14))

# Candidate Models
BV1=ddf(
   dsmodel=mcds(key="hn", formula=-1),
   mrmodel=glm(~distance),
   data=bird.data,
   method="io",
   meta.data=list(binned=TRUE, point=TRUE, width=100, breaks=c(0,50,100)))
BV1FI=ddf(
   dsmodel=mcds(key="hn", formula=-1),
   mrmodel=glm(~distance),
   data=bird.data,
   method="io.fi",
   meta.data=list(binned=TRUE, point=TRUE, width=100, breaks=c(0,50,100)))
BV2=ddf(
   dsmodel=mcds(key="hr", formula=-1),
   mrmodel=glm(~distance),
   data=bird.data,
   method="io",
   meta.data=list(binned=TRUE, point=TRUE, width=100, breaks=c(0,50,100)))
BV3=ddf(
   dsmodel=mcds(key="hn", formula=-1),
   mrmodel=glm(~distance+observer),
   data=bird.data,
   method="io",
   meta.data=list(binned=TRUE, point=TRUE, width=100, breaks=c(0,50,100)))
BV3FI=ddf(
   dsmodel=mcds(key="hn", formula=-1),
   mrmodel=glm(~distance+observer),
   data=bird.data,
   method="io.fi",
   meta.data=list(binned=TRUE, point=TRUE, width=100, breaks=c(0,50,100)))
BV4=ddf(
   dsmodel=mcds(key="hr", formula=-1),
   mrmodel=glm(~distance+observer),
   data=bird.data,
   method="io",
   meta.data=list(binned=TRUE, point=TRUE, width=100, breaks=c(0,50,100)))
BV5=ddf(
   dsmodel=mcds(key="hn", formula=-1),
   mrmodel=glm(~distance+observer),
   data=bird.data,
   method="io",
   meta.data=list(binned=TRUE, point=TRUE, width=100, breaks=c(0,50,100)))
mrmodel=glm(~distance*observer),
data=bird.data,
method="io",
meta.data=list(binned=TRUE, point=TRUE, width=100, breaks=c(0,50,100))
BV5FI=ddf(
  dsmodel=mcds(key="hn", formula=~1),
  mrmodel=glm(~distance*observer),
data=bird.data,
  method="io.fi",
  meta.data=list(binned=TRUE, point=TRUE, width=100, breaks=c(0,50,100)))
BV6=ddf(
  dsmodel=mcds(key="hr", formula=~1),
  mrmodel=glm(~distance*observer),
data=bird.data,
  method="io",
  meta.data=list(binned=TRUE, point=TRUE, width=100, breaks=c(0,50,100)))
BV7=ddf(
  dsmodel=cds(key="hn", formula=~1),
  mrmodel=glm(~distance*Day),
data=bird.data,
  method="io",
  meta.data=list(binned=TRUE, point=TRUE, width=100, breaks=c(0,50,100)))
BV7FI=ddf(
  dsmodel=cds(key="hn", formula=~1),
  mrmodel=glm(~distance*Day),
data=bird.data,
  method="io.fi",
  meta.data=list(binned=TRUE, point=TRUE, width=100, breaks=c(0,50,100)))
BV8=ddf(
  dsmodel=cds(key="hr", formula=~1),
  mrmodel=glm(~distance*Day),
data=bird.data,
  method="io",
  meta.data=list(binned=TRUE, point=TRUE, width=100, breaks=c(0,50,100)))
BV9=ddf(
  dsmodel=mcds(key="hn", formula=~1),
  mrmodel=glm(~distance*observer*Day),
data=bird.data,
  method="io",
  meta.data=list(binned=TRUE, point=TRUE, width=100, breaks=c(0,50,100)))
BV9FI=ddf(
  dsmodel=mcds(key="hn", formula=~1),
  mrmodel=glm(~distance*observer*Day),
data=bird.data,
  method="io.fi",
  meta.data=list(binned=TRUE, point=TRUE, width=100, breaks=c(0,50,100)))
BV10=ddf(
  dsmodel=mcds(key="hr", formula=~1),
  mrmodel=glm(~distance*observer*Day),
data=bird.data,
  method="io",
  meta.data=list(binned=TRUE, point=TRUE, width=100, breaks=c(0,50,100)))
#BV.DS=ddf(
dsmodel=mcds(key="hn", formula=-1),
data=bird.data,
method="ds",
meta.data=list(binned=TRUE, point=TRUE, width=100, breaks=c(0, 50, 100)))

# AIC table building code.
AIC = c(BV1$criterion, BV1FI$criterion, BV2$criterion, BV3$criterion, BV3FI$criterion, BV4$criterion, BV5$criterion, BV5FI$criterion, BV6$criterion, BV7$criterion, BV7FI$criterion, BV8$criterion, BV9$criterion, BV9FI$criterion, BV10$criterion)

# creates a set of row names for me to check my grep() call below
rn = c("BV1", "BV1FI", "BV2", "BV3", "BV3FI", "BV4", "BV5", "BV5FI", "BV6", "BV7", "BV7FI", "BV8", "BV9", "BV9FI", "BV10")

# Number parameters
k = c(length(BV1$par), length(BV1FI$par), length(BV2$par), length(BV3$par), length(BV3FI$par), length(BV4$par), length(BV5$par), length(BV5FI$par), length(BV6$par), length(BV7$par), length(BV7FI$par), length(BV8$par), length(BV9$par), length(BV9FI$par), length(BV10$par))

# Build AIC Table
AIC.table=data.frame(AIC = AIC, rn=rn, k=k, dAIC = abs(min(AIC)-AIC), likg=exp(-.5*(abs(min(AIC)-AIC))))
row.names(AIC.table)=grep("BV", ls(), value=TRUE)
AIC.table=AIC.table[with(AIC.table, order(-likg, -dAIC, AIC, k)),]
AIC.table=data.frame(AIC.table, wi=AIC.table$likg/sum(AIC.table$likg))
AIC.table

# Model average N_hat_covered estimates
# not very clean, but I wanted to show full process, need to use
# collect.models and model.table here later on
estimate <- c(BV1$Nhat, BV1FI$Nhat, BV2$Nhat, BV3$Nhat, BV3FI$Nhat, BV4$Nhat, BV5$Nhat, BV5FI$Nhat, BV6$Nhat, BV7$Nhat, BV7FI$Nhat, BV8$Nhat, BV9$Nhat, BV9FI$Nhat, BV10$Nhat)

AIC.values=AIC

# had to use str() to extract here as Nhat.se is calculated in
# mrds:::summary.io, not in ddf(), so it takes a bit

## End(Not run)

## Not run:
# Not run
Golden-cheeked warbler mark-recapture distance sampling analysis

Description

These data represent avian point count surveys conducted at 453 point sample survey locations on the 24,000 (approx) live-fire region of Fort Hood in central Texas. Surveys were conducted by independent double observers (2 per survey occasion) and as such we had a maximum of 3 paired survey histories, giving a maximum of 6 sample occasions (see MacKenzie et al. 2006, MacKenzie and Royle 2005, and Laake et al. 2011 for various sample survey design details). At each point, we surveyed for 5 minutes (technically broken into 3 time intervals of 2, 2, and 1 minutes; not used here) and we noted detections by each observer and collected distance to each observation within a set of distance bins (0-50, 50-100m; Laake et al. 2011) of the target species (Golden-cheeked warblers in this case) for each surveyor. Our primary focus was to use mark-recapture distance
sampling methods to estimate density of Golden-cheeked warblers, and to estimate detection rates for the mark-recapture, distance, and composite model.

Format

The format is a data frame with the following covariate metrics.

- **PointID** Unique identifier for each sample location; locations are the same for both species
- **VisitNumber** Visit number to the point
- **Species** Species designation, either Golden-cheeked warbler (GW) or Black-capped Vireo (BV)
- **Distance** Distance measure, which is either NA (representing no detection), or the median of the binned detection distances
- **PairNumber** ID value indicating which observers were paired for that sampling occasion
- **Observer** Observer ID, either primary (1), or secondary (2)
- **Detected** Detection of a bird, either 1 = detected, or 0 = not detected
- **Date** Date of survey since 15 March 2011, numeric value
- **Pred** Predicted occupancy value for that survey hexagon based on Farrell et al. (2013)
- **Category** Region.Label categorization, see R package mrd$ help file for details on data structure
- **Effort** Amount of survey effort at the point
- **Day** Number of days since 15 March 2011, numeric value
- **ObjectID** Unique ID for each paired observations

Details

In addition to detailing the analysis used by Collier et al. (2013, In Review), this example documents the use of mrd$ for avian point count surveys and shows how density models can be incorporated with occupancy models to develop spatially explicit density surface maps. For those that are interested, for the distance sampling portion of our analysis, we used both conventional distance sampling (cds) and multiple covariate distance sampling (mcds) with uniform and half-normal key functions. For the mark-recapture portion of our analysis, we tended to use covariates for distance (median bin width), observer, and date of survey (days since 15 March 2011).

We combined our mrd$ density estimates via a Horvitz-Thompson styled estimator with the resource selection function gradient developed in Farrell et al. (2013) and estimated density on an ~3.14ha hexagonal grid across our study area, which provided a density gradient for Fort Hood. Because there was considerable data manipulation needed for each analysis to structure the data appropriately for use in mrd$, rather than wrap each analysis in a single function, we have provided both the Golden-cheeked warbler and Black-capped vireo analyses in their full detail. The primary differences you will see will be changes to model structures and model outputs between the two species.

Author(s)

Bret Collier and Jeff Laake
lfgcwa

References


Examples

```r
## Not run:
data(lfgcwa)
xy <- cut(lfgcwa$Pred, c(-0.0001, .1, .2, .3, .4, .5, .6, .7, .8, .9, 1), 
labels=c("1", "2", "3", "4", "5", "6", "7", "8", "9", "10"))
x <- data.frame(lfgcwa, New=xy)

# Note that I scaled the individual covariate of day-helps with
# convergence issues
bird.data <- data.frame(object=x$ObjectID, observer=x$Observer,
detected=x$Detected, distance=x$Distance,
Region.Label=x$New, Sample.Label=x$PointID,
Day=(x$Day/max(x$Day)))

# make observer a factor variable
bird.data$observer=factor(bird.data$observer)

# Jeff Laake suggested this snippet to quickly create distance medians
# which adds bin information to the \code{bird.data} dataframe
bird.data$distbegin=0
bird.data$distend=100
bird.data$distend[bird.data$distance==12.5]=50
bird.data$distbegin[bird.data$distance==37.5]=0
bird.data$distend[bird.data$distance==37.5]=50
bird.data$distbegin[bird.data$distance==62.5]=50
bird.data$distend[bird.data$distance==62.5]=100
bird.data$distbegin[bird.data$distance==87.5]=50
bird.data$distend[bird.data$distance==87.5]=100

# Removed all survey points with distance=NA for a survey event;
# hence no observations for use in \code{ddf()} but needed later
bird.data=bird.data[complete.cases(bird.data),]

# Manipulations on full dataset for various data.frame creation
# for use in density estimation using \code{dht()}

# Samples data frame
```

```
xx <- x
x <- data.frame(PointID=x$PointID, Species=x$Species,
Category=x$New, Effort=x$Effort)
x <- x[!duplicated(x$PointID),]
point.num <- table(x$Category)
samples <- data.frame(PointID=x$PointID, Region.Label=x$Category,
Effort=x$Effort)
final.samples=data.frame(Sample.Label=samples$PointID,
Region.Label=samples$Region.Label,
Effort=samples$Effort)

# obs dataframe
obs <- data.frame(ObjectID=xx$ObjectID, PointID=xx$PointID)
# used to get Region and Sample assigned to ObjectID
obs <- merge(obs, samples, by=c("PointID", "PointID"))
obs <- obs[!duplicated(obs$ObjectID),]
obs <- data.frame(object=obs$ObjectID, Region.Label=obs$Region.Label,
Sample.Label=obs$PointID)

#Region.Label dataframe
region.data <- data.frame(Region.Label=c(1,2,3,4,5,6,7,8,9),
Area=c(point.num[1]*3.14,
point.num[2]*3.14,
point.num[3]*3.14,
point.num[4]*3.14,
point.num[5]*3.14,
point.num[6]*3.14,
point.num[7]*3.14,
point.num[8]*3.14,
point.num[9]*3.14))

# Candidate Models
GW1=ddf(
  dsmodel=~cds(key="unif", adj.series="cos", adj.order=1,adj.scale="width"),
mrmodel=~glm(~distance),
data=bird.data,
method="io",
meta.data=list(binned=TRUE,point=TRUE,width=100,breaks=c(0,50,100)))
GW2=ddf(
  dsmodel=~cds(key="unif", adj.series="cos", adj.order=1,adj.scale="width"),
mrmodel=~glm(~distance+observer),
data=bird.data,
method="io",
meta.data=list(binned=TRUE,point=TRUE,width=100,breaks=c(0,50,100)))
GW3=ddf(
  dsmodel=~cds(key="unif", adj.series="cos", adj.order=1,adj.scale="width"),
mrmodel=~glm(~distance*observer),
data=bird.data,
method="io",
meta.data=list(binned=TRUE,point=TRUE,width=100,breaks=c(0,50,100)))
GW4=ddf(
  dsmodel=mcds(key="hn",formula=~1),
mrmodel=glm(~distance),
data=bird.data,
method="io",
meta.data=list(binned=TRUE,point=TRUE,width=100,breaks=c(0,50,100)))

GW4FI=dfdf(
  dsmodel=mcds(key="hn",formula=~1),
mrmodel=glm(~distance),
data=bird.data,
method="io.fi",
meta.data=list(binned=TRUE,point=TRUE,width=100,breaks=c(0,50,100)))

GW5=dfdf(
  dsmodel=mcds(key="hn",formula=~1),
mrmodel=glm(~distance+observer),
data=bird.data,
method="io",
meta.data=list(binned=TRUE,point=TRUE,width=100,breaks=c(0,50,100)))

GW5FI=dfdf(
  dsmodel=mcds(key="hn",formula=~1),
mrmodel=glm(~distance+observer),
data=bird.data,
method="io.fi",
meta.data=list(binned=TRUE,point=TRUE,width=100,breaks=c(0,50,100)))

GW6=dfdf(
  dsmodel=mcds(key="hn",formula=~1),
mrmodel=glm(~distance*observer),
data=bird.data,
method="io",
meta.data=list(binned=TRUE,point=TRUE,width=100,breaks=c(0,50,100)))

GW6FI=dfdf(
  dsmodel=mcds(key="hn",formula=~1),
mrmodel=glm(~distance*observer),
data=bird.data,
method="io.fi",
meta.data=list(binned=TRUE,point=TRUE,width=100,breaks=c(0,50,100)))

GW7=dfdf(
  dsmodel=cds(key="hn",formula=~1),
mrmodel=glm(~distance*Day),
data=bird.data,
method="io",
meta.data=list(binned=TRUE,point=TRUE,width=100,breaks=c(0,50,100)))

GW7FI=dfdf(
  dsmodel=cds(key="hn",formula=~1),
mrmodel=glm(~distance*Day),
data=bird.data,
method="io.fi",
meta.data=list(binned=TRUE,point=TRUE,width=100,breaks=c(0,50,100)))

GW8=dfdf(
  dsmodel=mcds(key="hn",formula=~1),
mrmodel=glm(~distance*observer*Day),
data=bird.data,
method="io",
meta.data=list(binned=TRUE,point=TRUE,width=100,breaks=c(0,50,100)))

GW8FI=dfdf(
dsmodel = mcds(key = "hn", formula = '-1'),
mma = glm(~distance + observer + Day),
data = bird.data,
method = "io.fi",
meta.data = list(binned = TRUE, point = TRUE, width = 100, breaks = c(0, 50, 100)))

# GWDS = ddf(
# dsmodel = mcds(key = "hn", formula = '-1'),
data = bird.data,
method = "ds",
meta.data = list(binned = TRUE, point = TRUE, width = 100, breaks = c(0, 50, 100)))

#### GCWA Summary Metrics

# AIC table building code, not exactly elegant, but I did not
# want to add more package dependencies
AIC = c(GW1$criterion, GW2$criterion, GW3$criterion, GW4$criterion,
        GW4FI$criterion, GW5$criterion, GW5FI$criterion,
        GW6$criterion, GW6FI$criterion, GW7$criterion, GW7FI$criterion,
        GW8$criterion, GW8FI$criterion)

# creates a set of row names for me to check my grep() call below
rn <- c("GW1", "GW2", "GW3", "GW4", "GW4FI", "GW5", "GW5FI", "GW6",
        "GW6FI", "GW7", "GW7FI", "GW8", "GW8FI")

# number of parameters for each model
k <- c(length(GW1$par), length(GW2$par), length(GW3$par), length(GW4$par),
        length(GW4FI$par), length(GW5$par), length(GW5FI$par),
        length(GW6$par), length(GW6FI$par), length(GW7$par),
        length(GW7FI$par), length(GW8$par), length(GW8FI$par))

# build AIC table and
AIC.table <- data.frame(AIC = AIC, rn = rn, k = k, dAIC = abs(min(AIC) - AIC),
                         likg = exp(-.5 * (abs(min(AIC) - AIC))))

# row.names(AIC.table) = grep("GW", ls(), value = TRUE)
AIC.table <- AIC.table[with(AIC.table, order(-likg, -dAIC, AIC, k)),]
AIC.table <- data.frame(AIC.table, wi = AIC.table$likg / sum(AIC.table$likg))
AIC.table

# Model average N_hat_covered estimates
# not very clean, but I wanted to show full process, need to use
# collect.models and model.table here

estimate <- c(GW1$Nhat, GW2$Nhat, GW3$Nhat, GW4$Nhat, GW4FI$Nhat,
              GW5$Nhat, GW5FI$Nhat, GW6$Nhat, GW6FI$Nhat, GW7$Nhat,
              GW7FI$Nhat, GW8$Nhat, GW8FI$Nhat)

AIC.values <- AIC

# Nhat.se is calculated in mrds:::summary.io, not in ddf(), so
# it takes a bit to pull out
std.err <- c(summary(GW1)$Nhat.se, summary(GW2)$Nhat.se,
              summary(GW3)$Nhat.se, summary(GW4)$Nhat.se,
summary(GW4FI)$Nhat.se, summary(GW5)$Nhat.se, 
summary(GW5FI)$Nhat.se, summary(GW6)$Nhat.se, 
summary(GW6FI)$Nhat.se, summary(GW7)$Nhat.se, 
summary(GW7FI)$Nhat.se, summary(GW8)$Nhat.se, 
summary(GW8FI)$Nhat.se)

## End(Not run)
## Not run:
#Not Run
#requires RMark
library(RMark)
#uses model.average structure to model average real abundance estimates for 
#covered area of the surveys
mmi.list=list(estimate=estimate, AIC=AIC.values, se=std.err)
model.average(mmi.list, revised=TRUE)

#Not Run
#Best Model FI
#best.modelFI=AIC.table[1,]
#best.model
#Best Model PI
#best.modelPI=AIC.table[2,]
#best.modelPI

#Not Run
#summary(GW7FI, se=TRUE)
#summary(GW7, se=TRUE)

#Not Run
#GOF for models
#ddf.gof(GW7, breaks=c(0,50,100))

#Not Run
#Density estimation across occupancy categories
#out.GW=dht(GW7, region.data, final.samples, obs, se=TRUE, 
#options=list(convert.units=.01))

#Plots--Not Run
#Composite Detection Function examples
#plot(GW7, which=3, showpoints=FALSE, angle=0, density=0, 
# col="black", lwd=3, main="Golden-cheeked Warbler", 
# xlab="Distance (m)", las=1, cex.axis=1.25, cex.lab=1.25)

#Conditional Detection Function
#dd=expand.grid(distance=0:100,Day=(4:82)/82)
#dmat=model.matrix(~distance*Day,dd)
#dd$p=plogis(model.matrix(~distance*Day,dd)%*%coef(GW7$mr)$estimate)
#dd$Day=dd$Day*82
#with(dd[dd$Day==12,],plot(distance,p,ylim=c(0,1), las=1, cex.axis=1.25, cex.lab=1.25))
#with(dd[dd$Day==65,],lines(distance,p,lty=2, lwd=3))
#ch=paste(bird.data$detected[bird.data$observer==1],
# bird.data$detected[bird.data$observer==2],
# sep="")
# tab=table(ch, cut((b^1,bird.data$Day[bird.data$observer==1], c(0,45,83)),
# cut(bird.data$distance[bird.data$observer==1],c(0,50,100))))
# tabmat=cbind(colMeans(rbind(tab[3,1]/colSums(tab[2:3,1]),
# tab[3,1]/colSums(tab[c(1,3),,1]))),
# colMeans(rbind(tab[3,2]/colSums(tab[2:3,2]),
# tab[3,2]/colSums(tab[c(1,3),,2]))))
# colnames(tabmat)=c("0-50","51-100")
# points(c(25,75),tabmat[1,],pch=1, cex=1.5)
# points(c(25,75),tabmat[2,],pch=2, cex=1.5)

# Another alternative plot using barplot instead of points
# (this is one in paper)

# ch=paste(bird.data$detected[bird.data$observer==1],
# bird.data$detected[bird.data$observer==2],
# sep="")
# tab=table(ch, cut((b^1,bird.data$Day[bird.data$observer==1], c(0,45,83)),
# cut(bird.data$distance[bird.data$observer==1],c(0,50,100))))
# tabmat=cbind(colMeans(rbind(tab[3,1]/colSums(tab[2:3,1]),
# tab[3,1]/colSums(tab[c(1,3),,1]))),
# colMeans(rbind(tab[3,2]/colSums(tab[2:3,2]),
# tab[3,2]/colSums(tab[c(1,3),,2]))))
# colnames(tabmat)=c("0-50","51-100")
# par(mfrow=c(2, 1), mai=c(1,1,1,1))
# with(dd[dd$Day==12,],
# plot(distance,p,ylim=c(0,1), las=1,
# ylab="Detection probability", xlab="",
# type="l",lty=1, lwd=4, bty="l", cex.axis=1.5, cex.lab=1.5))
# segments(0, 0, 50, tabmat[1,1], lwd=3)
# segments(50, tabmat[1,1], 100, tabmat[1,1], lwd=4)
# segments(0, tabmat[1,2], 50, tabmat[1,2], lwd=4)
# segments(50, tabmat[1,2], 100, tabmat[1,2], lwd=4)
# segments(0, tabmat[1,1], 50, tabmat[1,1], lwd=4)
# segments(100, tabmat[1,2], 100, 0, lwd=4)
# mtext("a",line=-1, at=90)
# with(dd[dd$Day==65,],
# plot(distance,p,ylim=c(0,1), las=1, ylab="Detection probability",
# xlab="Distance", type="l",lty=1,
# lwd=4, bty="l", cex.axis=1.5, cex.lab=1.5))
# segments(0, 0, 50, tabmat[2,1], lwd=4)
# segments(50, tabmat[2,1], 100, tabmat[2,1], lwd=4)
# segments(0, tabmat[2,2], 50, tabmat[2,2], lwd=4)
# segments(50, tabmat[2,2], 100, tabmat[2,2], lwd=4)
# segments(0, tabmat[2,1], 50, tabmat[2,1], lwd=4)
# segments(100, tabmat[2,2], 100, 0, lwd=4)
# mtext("b",line=-1, at=90)

## End(Not run)
**logisticbyx**  
*Logistic as a function of covariates*

**Description**

Treats logistic as a function of covariates; for a given covariate combination it computes function at with those covariate values at a range of distances.

**Usage**

```r
logisticbyx(distance, x, models, beta, point)
```

**Arguments**

- `distance`: vector of distance values
- `x`: covariate data
- `models`: model list
- `beta`: logistic parameters
- `point`: TRUE if a point transect model

**Value**

vector of probabilities

**Author(s)**

Jeff Laake

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**logisticbyz**  
*Logistic as a function of distance*

**Description**

Treats logistic as a function of distance; for a given distance it computes function at all covariate values in data.

**Usage**

```r
logisticbyz(x, distance, models, beta)
```

**Arguments**

- `x`: covariate data
- `distance`: single distance value
- `models`: model list
- `beta`: logistic parameters
logisticdetfct

Description

Logistic detection function

Usage

logisticdetfct(distance, theta, w, std = FALSE)

Arguments

distance     perpendicular distance vector
theta        scale parameters
w             scale covariate matrix
std          if TRUE uses scale=1

The routine returns a vector of probabilities that the observation were detected given they were at the specified distance and assuming that g(0)=1 (ie a standard line transect detection function).

logisticdupbyx

Description

Treats logistic for duplicates as a function of covariate z; for a given z it computes the function at with those covariate values at a range of distances.

Usage

logisticdupbyx(distance, x1, x2, models, beta, point)
logisticdupbyx_fast

**Arguments**

- **distance**: vector of distance values
- **x1**: covariate data for fct 1
- **x2**: covariate data for fct 2
- **models**: model list
- **beta**: logistic parameters
- **point**: TRUE for point transect data

**Value**

vector of probabilities

**Author(s)**

Jeff Laake

---

**logisticdupbyx_fast**  *Logistic for duplicates as a function of covariates (fast)*

**Description**

As `logisticdupbyx`, but faster when distance is a covariate (but no interactions with distance occur.

**Usage**

`logisticdupbyx_fast(distance, x1, x2, models, beta, point, beta_distance)`

**Arguments**

- **distance**: vector of distance values
- **x1**: linear predictor for 1, without distance
- **x2**: linear predictor for 2, without distance
- **models**: model list
- **beta**: logistic parameters
- **point**: TRUE for point transect data
- **beta_distance**: parameter for distance

**Author(s)**

David L Miller


\begin{verbatim}

logit

Logit function

Description
Computes logit transformation.

Usage
logit(p)

Arguments
p probability

Value
logit(p) returns \( \log(p/(1-p)) \)

Author(s)
Jeff Laake

logLik.ds log-likelihood value for a fitted detection function

Description
Extract the log-likelihood from a fitted detection function.

Usage
## S3 method for class 'ds'
logLik(object, ...)

Arguments
object a fitted detection function model object
... included for S3 completeness, but ignored

Value
a numeric value giving the log-likelihood with two attributes: "df" the "degrees of freedom" for the model (number of parameters) and "nobs" the number of observations used to fit the model

Author(s)
David L Miller

\end{verbatim}
**mcds**

**MCDS function definition**

**Description**

Creates model formula list for multiple covariate distance sampling using values supplied in call to **ddf**

**Usage**

```r
mcds(
  formula = NULL,
  key = NULL,
  adj.series = NULL,
  adj.order = c(NULL),
  adj.scale = "width",
  adj.exp = FALSE,
  shape.formula = ~1
)
```

**Arguments**

- `formula`: formula for scale function
- `key`: string identifying key function (currently either "hn" (half-normal), "hr" (hazard-rate), "unif" (uniform) or "gamma" (gamma distribution))
- `adj.series`: string identifying adjustment functions cos (Cosine), herm (Hermite polynomials), poly (simple polynomials) or NULL
- `adj.order`: vector of order of adjustment terms to include
- `adj.scale`: whether to scale the adjustment terms by "width" or "scale"
- `adj.exp`: if TRUE uses exp(adj) for adjustment to keep f(x)>0
- `shape.formula`: formula for shape function

**Value**

A formula list used to define the detection function model

```r
fct string "mcds"
key key function string
adj.series adjustment function string
adj.order adjustment function orders
adj.scale adjustment function scale type
formula formula for scale function
shape.formula formula for shape function
```
Description

Occasionally when fitting an 'mrds' model one can run into optimisation issues. In general such problems can be quite complex so these "quick fixes" may not work. If you come up against problems that are not fixed by these tips, or you feel the results are dubious please go ahead and contact the package authors.

Debug mode

One can obtain debug output at each stage of the optimisation using the showit option. This is set via control, so adding control=list(showit=3) gives the highest level of debug output (setting showit to 1 or 2 gives less output).

Re-scaling covariates

Sometimes convergence issues in covariate (MCDS) models are caused by values of the covariate being very large, so a rescaling of that covariate is then necessary. Simply scaling by the standard deviation of the covariate can help (e.g. dat$size.scaled <- dat$scale/sd(dat$scale) for a covariate size, then including size.scaled in the model instead of size).

It is important to note that one needs to use the original covariate (size) when computing Horvitz-Thompson estimates of population size if the group size is used in that estimate. i.e. use the unscaled size in the numerator of the H-T estimator.

Initial values

Initial (or starting) values can be set via the initial element of the control list. initial is a list itself with elements scale, shape and adjustment, corresponding to the associated parameters. If a model has covariates then the scale or shape elements will be vectors with parameter initial values in the same order as they are specific in the model formula (using showit is a good check they are in the correct order). Adjustment starting values are in order of the order of that term (cosine order 2 is before cosine order 3 terms).

One way of obtaining starting values is to fit a simpler model first (say with fewer covariates or adjustments) and then use the starting values from this simpler model for the corresponding parameters.

Another alternative to obtain starting values is to fit the model (or some submodel) using Distance for Windows. Note that Distance reports the scale parameter (or intercept in a covariate model) on the exponential scale, so one must log this before supplying it to ddf.
**Bounds**

One can change the upper and lower bounds for the parameters. These specify the largest and smallest values individual parameters can be. By placing these constraints on the parameters, it is possible to "temper" the optimisation problem, making fitting possible.

Again, one uses the control list, the elements upperbounds and lowerbounds. In this case, each of upperbounds and lowerbounds are vectors, which one can think of as each of the vectors scale, shape and adjustment from the "Initial values" section above, concatenated in that order. If one does not occur (e.g. no shape parameter) then it is simple omitted from the vector.

**Author(s)**

David L. Miller <dave@ninepointeightone.net>

---

**NCovered**

*Compute estimated abundance in covered (sampled) region*

**Description**

Generic function that computes abundance within the covered region. It calls method (class) specific functions for the computation.

**Usage**

```
NCovered(par, model = NULL, group = TRUE)
```

**Arguments**

- `par` : parameter values (used when computing derivatives wrt parameter uncertainty); if NULL parameter values in `model` are used
- `model` : ddf model object
- `group` : if TRUE computes group abundance and if FALSE individual abundance

**Value**

- abundance estimate

**Author(s)**

Jeff Laake
nlminb_wrapper

Wrapper around nlminb

Description
This is a wrapper around nlminb to use scaling, as this is not available (nor will it be) in optimx.

Usage

```r
nlminb_wrapper(par,
    ll,
    ugr = NULL,
    lower = NULL,
    upper = NULL,
    mcontrol,
    hess = NULL,
    ddfobj,
    data,
    ...
)
```

Arguments

- `par` starting parameters
- `ll` log likelihood function
- `ugr` gradient function
- `lower` lower bounds on parameters
- `upper` upper bounds on parameters
- `mcontrol` control options
- `hess` hessian function
- `ddfobj` detection function specification object
- `data` the data
- `...` anything else to pass to `ll`

Value

optimx object

Author(s)

David L Miller, modified from optimx.run by JC Nash, R Varadhan, G Grothendieck.
p.det

Double-platform detection probability

Description
Computes detection probability for detection function computed from mark-recapture data with possibly different link functions.

Usage
p.det(dpformula, dplink, dppars, dpdata)

Arguments
- dpformula: formula for detection function
- dplink: link function ("logit","loglog","cloglog")
- dppars: parameter vector
- dpdata: double platform data

Value
vector of predicted detection probabilities

Author(s)
?????

parse.optimx
Parse optimx results and present a nice object

Description
Take the resulting object from a call to optimx and make it into an object that mrds wants to talk to.

Usage
parse.optimx(lt, lnl.last, par.last)

Arguments
- lt: an optimx object
- lnl.last: last value of the log likelihood
- par.last: last value of the parameters

Value
lt object that can be used later on
pdot.dsr.integrate.logistic

Compute probability that an object was detected by at least one observer

Description

Computes probability that an object was detected by at least one observer (pdot or p_) for a logistic detection function that contains distance.

Usage

pdot.dsr.integrate.logistic(
  right,
  width,
  beta,
  x,
  integral.numeric,
  BT,
  models,
  GAM = FALSE,
  rem = FALSE,
  point = FALSE
)

Arguments

right either an integration range for binned data (vector of 2) or the rightmost value for integration (from 0 to right)
width transect width
beta parameters of logistic detection function
x data matrix
integral.numeric set to TRUE unless data are binned (done in this fct) or the model is such that distance is not linear (eg distance^2), If integral.numeric is FALSE it will compute the integral analytically. It should only be FALSE if is.linear.logistic function is TRUE.
BT FALSE except for the trial configuration; BT stands for Buckland-Turnock who initially proposed a trial configuration for dual observers
models list of models including g0model
GAM Not used at present. The idea was to be able to use a GAM for g(0) portion of detection function; should always be F
rem only TRUE for the removal configuration but not used and could be removed if pulled from the function calls. Originally thought the pdot integral would differ but it is the same as the io formula. The only thing that differs with removal is that p(2|1) = 1. Observer 2 sees everything seen by observer 1,
Author(s)

Jeff Laake

Description

Plot the tables created by `det.tables`. Produces a series of tables for dual observer data that shows the number missed and detected for each observer within defined distance classes.

Usage

```r
## S3 method for class 'det.tables'
plot(
  x,
  which = 1:6,
  angle = NULL,
  density = NULL,
  col1 = "white",
  col2 = "lightgrey",
  new = TRUE,
  ...,
)
```

Arguments

- `x`: object returned by `det.tables`
- `which`: items in x to plot (vector with values in 1:6)
- `angle`: shading angle for hatching
- `density`: shading density for hatching
- `col1`: plotting colour for total histogram bars.
- `col2`: plotting colour for subset histogram bars.
- `new`: if TRUE new plotting window for each plot
- `...`: other graphical parameters, passed to plotting functions

Details

Plots that are produced are as follows (controlled by the `which` argument):

1. Detected by either observer/Detected by observer 1
2. Detected by either observer/Detected by observer 2
plot.ds

Plot fit of detection functions and histograms of data from distance sampling model

Description

Plots the fitted detection function(s) with a histogram of the observed distances to compare visually the fitted model and data.

Usage

```r
## S3 method for class 'ds'
plot(
  x,
  which = 2,
  breaks = NULL,
  nc = NULL,
  jitter.v = rep(0, 3),
  showpoints = TRUE,
  ...)
```

Examples

```r
data(book.tee.data)
region <- book.tee.data$book.tee.region
egdata <- book.tee.data$book.tee.dataframe
samples <- book.tee.data$book.tee.samples
obs <- book.tee.data$book.tee.obs
xx <- ddf(mrmodel=~glm(formula=~distance*observer),
  dsmodel = ~mcds(key = "hn", formula = ~sex),
  data = egdata, method = "io", meta.data = list(width = 4))
tabs <- det.tables(xx,breaks=c(0,.5,1,2,3,4))
par(mfrow=c(2,3))
plot(tabs,which=1:6,new=FALSE)
```
plot.ds

subset = NULL,
pl.col = "lightgrey",
pl.den = NULL,
pl.ang = NULL,
main = NULL,
pages = 0,
pdf = FALSE,
ylim = NULL,
pl.lab = "Distance",
plab = NULL,
...)

Arguments

x fitted model from ddf.
which index to specify which plots should be produced:

1 histogram of observed distances
2 histogram of observed distances with fitted line and points (default)

breaks user defined breakpoints
nc number of equal width bins for histogram
jitter.v apply jitter to points by multiplying the fitted value by a random draw from a
normal distribution with mean 1 and sd jitter.v.
showpoints logical variable; if TRUE plots predicted value for each observation (conditional
on its observed distance).
subset subset of data to plot.
pl.col colour for histogram bars.
pl.den shading density for histogram bars.
pl.ang shading angle for histogram bars.
main plot title.
pages the number of pages over which to spread the plots. For example, if pages=1
then all plots will be displayed on one page. Default is 0, which prompts the
user for the next plot to be displayed.
pdf plot the histogram of distances with the PDF of the probability of detection
overlaid. Ignored (with warning) for line transect models.
ylim vertical axis limits.
xlab horizontal axis label (defaults to "Distance").
ylab vertical axis label (default automatically set depending on plot type).
... other graphical parameters, passed to the plotting functions (plot, hist, lines,
points, etc).
Details

The structure of the histogram can be controlled by the user-defined arguments `nc` or `breaks`. The observation specific detection probabilities along with the line representing the fitted average detection probability.

It is not intended for the user to call `plot.ds` but its arguments are documented here. Instead the generic `plot` command should be used and it will call the appropriate function based on the class of the `ddf` object.

Value

Just plots.

Author(s)

Jeff Laake, Jon Bishop, David Borchers, David L Miller

See Also

`add_df_covar_line`

Examples

```r
# fit a model to the tee data
data(book.tee.data)
egdata <- book.tee.data$book.tee.dataframe
xx <- ddf(dsmodel=mcds(key="hn", formula=~sex),
    data=egdata[egdata$observer==1, ],
    method="ds", meta.data=list(width=4))

# not showing predicted probabilities
plot(xx, breaks=c(0, 0.5, 1, 2, 3, 4), showpoints=FALSE)

# two subsets
plot(xx, breaks=c(0, 0.5, 1, 2, 3, 4), subset=sex==0)
plot(xx, breaks=c(0, 0.5, 1, 2, 3, 4), subset=sex==1)

# put both plots on one page
plot(xx, breaks=c(0, 0.5, 1, 2, 3, 4), pages=1, which=1:2)
```
Description

Plots the fitted detection functions for a distance sampling model and histograms of the distances (for unconditional detection functions) or proportion of observations detected within distance intervals (for conditional detection functions) to compare visually the fitted model and data.

Usage

```r
## S3 method for class 'io'
plot(
    x,
    which = 1:6,
    breaks = NULL,
    nc = NULL,
    maintitle = "",
    showlines = TRUE,
    showpoints = TRUE,
    ylim = c(0, 1),
    angle = NULL,
    density = NULL,
    col = "lightgrey",
    jitter = NULL,
    divisions = 25,
    pages = 0,
    xlab = "Distance",
    ylab = "Detection probability",
    subtitle = TRUE,
    ...
)
```

Arguments

- `x`: fitted model from `ddf`
- `which`: index to specify which plots should be produced.
  1. Plot primary unconditional detection function
  2. Plot secondary unconditional detection function
  3. Plot pooled unconditional detection function
  4. Plot duplicate unconditional detection function
  5. Plot primary conditional detection function
  6. Plot secondary conditional detection function

Note that the order of which is ignored and plots are produced in the above order.

- `breaks`: user define breakpoints
- `nc`: number of equal-width bins for histogram
- `maintitle`: main title line for each plot
showlines logical variable; if TRUE a line representing the average detection probability is plotted
showpoints logical variable; if TRUE plots predicted value for each observation
ylim range of vertical axis; defaults to (0,1)
angle shading angle for histogram bars.
density shading density for histogram bars.
col colour for histogram bars.
jitter scaling option for plotting points. Jitter is applied to points by multiplying the fitted value by a random draw from a normal distribution with mean 1 and sd jitter.
divisions number of divisions for averaging line values; default = 25
pages the number of pages over which to spread the plots. For example, if pages=1 then all plots will be displayed on one page. Default is 0, which prompts the user for the next plot to be displayed.
xlab label for x-axis
ylab label for y-axis
subtitle if TRUE, shows plot type as sub-title
... other graphical parameters, passed to the plotting functions (plot, hist, lines, points, etc)

Details

The structure of the histogram can be controlled by the user-defined arguments nc or breaks. The observation specific detection probabilities along with the line representing the fitted average detection probability.

It is not intended for the user to call plot.io.fi but its arguments are documented here. Instead the generic plot command should be used and it will call the appropriate function based on the class of the ddf object.

Value

Just plots

Author(s)

Jeff Laake, Jon Bishop, David Borchers, David L Miller

Examples

library(mrds)
data(book.tee.data)
egdata <- book.tee.data$book.tee.dataframe
result.io <- ddf(dsmodel=~cds(key = "hn"), mrmodel=~glm(~distance),
data=egdata, method="io", meta.data=list(width=4))
# just plot everything
plot(result.io)

# Plot primary and secondary unconditional detection functions on one page
# and primary and secondary conditional detection functions on another
plot(result.io, which=c(1,2,5,6), pages=2)

---

**plot.io.fi**

*Plot fit of detection functions and histograms of data from distance sampling independent observer model with full independence (io.fi)*

**Description**

Plots the fitted detection functions for a distance sampling model and histograms of the distances (for unconditional detection functions) or proportion of observations detected within distance intervals (for conditional detection functions) to compare visually the fitted model and data.

**Usage**

```r
## S3 method for class 'io.fi'
plot(
  x,
  which = 1:6,
  breaks = NULL,
  nc = NULL,
  maintitle = "",
  showlines = TRUE,
  showpoints = TRUE,
  ylim = c(0, 1),
  angle = NULL,
  density = NULL,
  col = "lightgrey",
  jitter = NULL,
  divisions = 25,
  pages = 0,
  xlab = "Distance",
  ylab = "Detection probability",
  subtitle = TRUE,
  ...
)
```

**Arguments**

- `x` fitted model from `ddf`
which index to specify which plots should be produced.

1 Plot primary unconditional detection function
2 Plot secondary unconditional detection function
3 Plot pooled unconditional detection function
4 Plot duplicate unconditional detection function
5 Plot primary conditional detection function
6 Plot secondary conditional detection function

Note that the order of which is ignored and plots are produced in the above order.

breaks user define breakpoints
nc number of equal-width bins for histogram
maintitle main title line for each plot
showlines logical variable; if TRUE a line representing the average detection probability is plotted
showpoints logical variable; if TRUE plots predicted value for each observation
ylim range of vertical axis; defaults to (0,1)
angle shading angle for histogram bars.
density shading density for histogram bars.
col colour for histogram bars.
jitter scaling option for plotting points. Jitter is applied to points by multiplying the fitted value by a random draw from a normal distribution with mean 1 and sd jitter.
divisions number of divisions for averaging line values; default = 25
pages the number of pages over which to spread the plots. For example, if pages=1 then all plots will be displayed on one page. Default is 0, which prompts the user for the next plot to be displayed.
xlab label for x-axis
ylab label for y-axis
subtitle if TRUE, shows plot type as sub-title
...
other graphical parameters, passed to the plotting functions (plot, hist, lines, points, etc)

Details

The structure of the histogram can be controlled by the user-defined arguments nc or breaks. The observation specific detection probabilities along with the line representing the fitted average detection probability.

It is not intended for the user to call `plot.io.fi` but its arguments are documented here. Instead the generic `plot` command should be used and it will call the appropriate function based on the class of the ddf object.
Value

Just plots.

Author(s)

Jeff Laake, Jon Bishop, David Borchers, David L Miller

Examples

```r
library(mrds)
data(book.tee.data)
egdata <- book.tee.data$book.tee.dataframe
result.io.fi <- ddf(mrmodel=~glm(~distance), data = egdata, method = "io.fi",
meta.data = list(width = 4))

# just plot everything
plot(result.io.fi)

# Plot primary and secondary unconditional detection functions on one page
# and primary and secondary conditional detection functions on another
plot(result.io.fi,which=c(1,2,5,6),pages=2)
```

---

plot.layout

Layout for plot methods in mrds

Description

This function does the paging, using devAskNewPage(). This means we can just call plots and R will make the prompt for us. Warning, this function has side effects! It modifies devAskNewPage!

Usage

```r
## S3 method for class 'layout'
plot(which, pages)
```

Arguments

- `which`: which plots are to be created
- `pages`: number of pages to span the plots across

Details

Code is stolen and modified from `plot.R` in mgcv by Simon Wood

Author(s)

David L. Miller, based on code by Simon N. Wood
plot.rem  

Plot fit of detection functions and histograms of data from removal distance sampling model

Description

Plots the fitted detection functions for a distance sampling model and histograms of the distances (for unconditional detection functions) or proportion of observations detected within distance intervals (for conditional detection functions) to compare visually the fitted model and data.

Usage

```r
## S3 method for class 'rem'
plot(
x, which = 1:3,
breaks = NULL,
nc = NULL,
maintitle = "",
showlines = TRUE,
showpoints = TRUE,
ylim = c(0, 1),
angle = NULL,
density = NULL,
col = "lightgrey",
jitter = NULL,
divisions = 25,
pages = 0,
xlab = "Distance",
ylab = "Detection probability",
subtitle = TRUE,
...
)
```

Arguments

- `x`: fitted model from `ddf`
- `which`: index to specify which plots should be produced.
  - 1: Plot primary unconditional detection function
  - 2: Plot pooled unconditional detection function
  - 3: Plot conditional (1|2) detection function
- `breaks`: user define breakpoints
- `nc`: number of equal-width bins for histogram
### Parameters

- **maintitle**: main title line for each plot
- **showlines**: logical variable; if TRUE a line representing the average detection probability is plotted
- **showpoints**: logical variable; if TRUE plots predicted value for each observation
- **ylim**: range of vertical axis; defaults to (0,1)
- **angle**: shading angle for histogram bars.
- **density**: shading density for histogram bars.
- **col**: colour for histogram bars.
- **jitter**: scaling option for plotting points. Jitter is applied to points by multiplying the fitted value by a random draw from a normal distribution with mean 1 and sd jitter.
- **divisions**: number of divisions for averaging line values; default = 25
- **pages**: the number of pages over which to spread the plots. For example, if pages=1 then all plots will be displayed on one page. Default is 0, which prompts the user for the next plot to be displayed.
- **xlab**: label for x-axis
- **ylab**: label for y-axis
- **subtitle**: if TRUE, shows plot type as sub-title
- **...**: other graphical parameters, passed to the plotting functions (plot, hist, lines, points, etc)

### Details

The structure of the histogram can be controlled by the user-defined arguments `nc` or `breaks`. The observation specific detection probabilities along with the line representing the fitted average detection probability.

It is not intended for the user to call `plot.rem` but its arguments are documented here. Instead the generic `plot` command should be used and it will call the appropriate function based on the class of the `ddf` object.

### Author(s)

Jeff Laake, Jon Bishop, David Borchers, David L Miller

### Description

Plots the fitted detection functions for a distance sampling model and histograms of the distances (for unconditional detection functions) or proportion of observations detected within distance intervals (for conditional detection functions) to compare visually the fitted model and data.
Usage

```r
## S3 method for class 'rem.fi'
plot(
  x,
  which = 1:3,
  breaks = NULL,
  nc = NULL,
  maintitle = "",
  showlines = TRUE,
  showpoints = TRUE,
  ylim = c(0, 1),
  angle = NULL,
  density = NULL,
  col = "lightgrey",
  jitter = NULL,
  divisions = 25,
  pages = 0,
  xlab = "Distance",
  ylab = "Detection probability",
  subtitle = TRUE,
  ...
)
```

Arguments

- `x` fitted model from `ddf`  
- `which` index to specify which plots should be produced.  
  - 1 Plot primary unconditional detection function  
  - 2 Plot pooled unconditional detection function  
  - 3 Plot conditional (1|2) detection function  
- `breaks` user defined breakpoints  
- `nc` number of equal-width bins for histogram  
- `maintitle` main title line for each plot  
- `showlines` logical variable; if `TRUE` a line representing the average detection probability is plotted  
- `showpoints` logical variable; if `TRUE` plots predicted value for each observation  
- `ylim` range of vertical axis; defaults to `(0,1)`  
- `angle` shading angle for histogram bars.  
- `density` shading density for histogram bars.  
- `col` colour for histogram bars.  
- `jitter` scaling option for plotting points. Jitter is applied to points by multiplying the fitted value by a random draw from a normal distribution with mean 1 and sd `jitter`
divisions  number of divisions for averaging line values; default = 25
pages    the number of pages over which to spread the plots. For example, if pages=1 then all plots will be displayed on one page. Default is 0, which prompts the user for the next plot to be displayed.
xlab      label for x-axis
ylab      label for y-axis
subtitle if TRUE, shows plot type as sub-title
...       other graphical parameters, passed to the plotting functions (plot, hist, lines, points, etc)

Details

The structure of the histogram can be controlled by the user-defined arguments nc or breaks. The observation specific detection probabilities along with the line representing the fitted average detection probability.

It is not intended for the user to call plot.rem.fi but its arguments are documented here. Instead the generic plot command should be used and it will call the appropriate function based on the class of the ddf object.

Author(s)

Jeff Laake, Jon Bishop, David Borchers, David L Miller

plot.trial  Plot fit of detection functions and histograms of data from distance sampling trial observer model

Description

Plots the fitted detection functions for a distance sampling model and histograms of the distances (for unconditional detection functions) or proportion of observations detected within distance intervals (for conditional detection functions) to compare visually the fitted model and data.

Usage

```r
## S3 method for class 'trial'
plot(
x, 
which = 1:2,
breaks = NULL,
nc = NULL,
maintitle = "",
showlines = TRUE,
showpoints = TRUE,
ylim = c(0, 1),
```

angle = NULL,  
density = NULL,  
col = "lightgrey",  
jitter = NULL,  
divisions = 25,  
pages = 0,  
xlab = "Distance",  
ylab = "Detection probability",  
subtitle = TRUE,  
...  
)

Arguments

x                 fitted model from ddf
which             index to specify which plots should be produced.

1  Unconditional detection function for observer 1
2  Conditional detection function plot (1|2)

breaks           user defined breakpoints
nc                number of equal-width bins for histogram
maintitle        main title line for each plot
showlines        logical variable; if TRUE a line representing the average detection probability is plotted
showpoints       logical variable; if TRUE plots predicted value for each observation
ylim             range of vertical axis; defaults to (0,1)
angle            shading angle for histogram bars.
density          shading density for histogram bars.
col              colour for histogram bars.
jitter           scaling option for plotting points. Jitter is applied to points by multiplying the fitted value by a random draw from a normal distribution with mean 1 and sd jitter.
divisions        number of divisions for averaging line values; default = 25
pages            the number of pages over which to spread the plots. For example, if pages=1 then all plots will be displayed on one page. Default is 0, which prompts the user for the next plot to be displayed.
xlab             label for x-axis
ylab             label for y-axis
subtitle         if TRUE, shows plot type as sub-title
...              other graphical parameters, passed to the plotting functions (plot, hist, lines, points, etc)
Details

The structure of the histogram can be controlled by the user-defined arguments nc or breaks. The observation specific detection probabilities along with the line representing the fitted average detection probability.

It is not intended for the user to call plot.io.fi but its arguments are documented here. Instead the generic plot command should be used and it will call the appropriate function based on the class of the ddf object.

Author(s)

Jeff Laake, Jon Bishop, David Borchers

Description

Plots the fitted detection functions for a distance sampling model and histograms of the distances (for unconditional detection functions) or proportion of observations detected within distance intervals (for conditional detection functions) to compare visually the fitted model and data.

Usage

```r
## S3 method for class 'trial.fi'
plot(
x,
  which = 1:2,
  breaks = NULL,
  nc = NULL,
  maintitle = "",
  showlines = TRUE,
  showpoints = TRUE,
  ylim = c(0, 1),
  angle = NULL,
  density = NULL,
  col = "lightgrey",
  jitter = NULL,
  divisions = 25,
  pages = 0,
  xlab = "Distance",
  ylab = "Detection probability",
  subtitle = TRUE,
  ...
)
```
Arguments

- **x**: fitted model from `ddf`
- **which**: index to specify which plots should be produced.
  1. Unconditional detection function for observer 1
  2. Conditional detection function plot (1|2)

- **breaks**: user define breakpoints
- **nc**: number of equal-width bins for histogram
- **maintitle**: main title line for each plot
- **showlines**: logical variable; if `TRUE` a line representing the average detection probability is plotted
- **showpoints**: logical variable; if `TRUE` plots predicted value for each observation
- **ylim**: range of vertical axis; defaults to (0,1)
- **angle**: shading angle for histogram bars.
- **density**: shading density for histogram bars.
- **col**: colour for histogram bars.
- **jitter**: scaling option for plotting points. Jitter is applied to points by multiplying the fitted value by a random draw from a normal distribution with mean 1 and sd jitter.
- **divisions**: number of divisions for averaging line values; default = 25
- **pages**: the number of pages over which to spread the plots. For example, if `pages=1` then all plots will be displayed on one page. Default is 0, which prompts the user for the next plot to be displayed.
- **xlab**: label for x-axis
- **ylab**: label for y-axis
- **subtitle**: if `TRUE`, shows plot type as sub-title
- **...**: other graphical parameters, passed to the plotting functions (`plot`, `hist`, `lines`, `points`, etc)

Details

The structure of the histogram can be controlled by the user-defined arguments `nc` or `breaks`. The observation specific detection probabilities along with the line representing the fitted average detection probability.

It is not intended for the user to call `plot.io.fi` but its arguments are documented here. Instead the generic `plot` command should be used and it will call the appropriate function based on the class of the `ddf` object.

Author(s)

Jeff Laake, Jon Bishop, David Borchers
plot_cond

Plot conditional detection function from distance sampling model

Description

Plot proportion of observations detected within distance intervals (for conditional detection functions) to compare visually the fitted model and data. Internal function called by plot methods.

Usage

plot_cond(
  obs, xmat, gxvalues, model, nc, breaks, finebr, showpoints, showlines, maintitle, ylim, angle = -45, density = 20, col = "black", jitter = NULL, xlab = "Distance", ylab = "Detection probability", subtitle = TRUE,
  ...
)

Arguments

obs observer code
xmat processed data
gxvalues detection function values for each observation
model fitted model from ddf
nc number of equal-width bins for histogram
breaks user define breakpoints
finebr fine break values over which line is averaged
showpoints logical variable; if TRUE plots predicted value for each observation
showlines logical variable; if TRUE plots average predicted value line
maintitle main title line for each plot
 ylim range of y axis (default c(0,1))
angle shading angle for hatching
density shading density for hatching
col plotting colour
jitter scaling option for plotting points. Jitter is applied to points by multiplying the fitted value by a random draw from a normal distribution with mean 1 and sd jitter.
xlab label for x-axis
ylab label for y-axis
subtitle if TRUE, shows plot type as sub-title
... other graphical parameters, passed to the plotting functions (plot, hist, lines, points, etc)

Author(s)
Jeff Laake, Jon Bishop, David Borchers

plot_uncond  Plot unconditional detection function from distance sampling model

Description
Plots unconditional detection function for observer=obs observations overlays histogram, average detection function and values for individual observations data. Internal function called by plot methods.

Usage
plot_uncond(
  model,
  obs,
  xmat,
  gxvalues,
  nc,
  finebr,
  breaks,
  showpoints,
  showlines,
  maintitle,
  ylim,
  return.lines = FALSE,
  angle = -45,
  density = 20,
  col = "black",
  jitter = NULL,
xlab = "Distance",
ylab = "Detection probability",
subtitle = TRUE,
...
)

Arguments

model fitted model from ddf
obs value of observer for plot
xmat processed data
gxvalues detection function values for each observation
nc number of equal-width bins for histogram
finebr fine break values over which line is averaged
breaks user define breakpoints
showpoints logical variable; if TRUE plots predicted value for each observation
showlines logical variable; if TRUE plots average predicted value line
maintitle main title line for each plot
ylim range of y axis; defaults to (0,1)
return.lines if TRUE, returns values for line
angle shading angle for hatching
density shading density for hatching
col plotting colour
jitter scaling option for plotting points. Jitter is applied to points by multiplying the fitted value by a random draw from a normal distribution with mean 1 and sd jitter.

xlab label for x-axis
ylab label for y-axis
subtitle if TRUE, shows plot type as sub-title
...
other graphical parameters, passed to the plotting functions (plot, hist, lines, points, etc)

Value

if return.lines==TRUE returns dataframe average.line otherwise just plots

Author(s)

Jeff Laake, Jon Bishop, David Borchers
Predictions from mrds models

Description

Predict detection probabilities (or effective strip widths/effective areas of detection) from a fitted distance sampling model using either the original data (i.e. "fitted" values) or using new data.

Usage

## S3 method for class 'ds'
predict(object, newdata=NULL, compute=FALSE, int.range=NULL, esw=FALSE, se.fit=FALSE, ...)
## S3 method for class 'io.fi'
predict(object, newdata=NULL, compute=FALSE, int.range=NULL, integrate=FALSE, ...)
## S3 method for class 'io'
predict(object, newdata=NULL, compute=FALSE, int.range=NULL, ...)
## S3 method for class 'trial'
predict(object, newdata=NULL, compute=FALSE, int.range=NULL, ...)
## S3 method for class 'trial.fi'
predict(object, newdata=NULL, compute=FALSE, int.range=NULL, integrate=FALSE, ...)
## S3 method for class 'rem'
predict(object, newdata=NULL, compute=FALSE, int.range=NULL, ...)
## S3 method for class 'rem.fi'
predict(object, newdata=NULL, compute=FALSE, int.range=NULL, integrate=FALSE, ...)

Arguments

- **object**: ddf model object.
- **newdata**: new data.frame for prediction, this must include a column called "distance".
- **compute**: if TRUE compute values and don’t use the fitted values stored in the model object.
- **int.range**: integration range for variable range analysis; either vector or 2 column matrix.
- **esw**: if TRUE, returns effective strip half-width (or effective area of detection for point transect models) integral from 0 to the truncation distance (width) of \( p(y)dy \); otherwise it returns the integral from 0 to truncation width of \( p(y)\pi(y) \) where \( \pi(y) = 1/w \) for lines and \( \pi(y) = 2r/w^2 \) for points.
- **se.fit**: for *.ds models only, generate standard errors on the predicted probabilities of detection (or ESW if esw=TRUE), stored in the se.fit element
- **integrate**: for *.fi methods, see Details below.

Details

The first 4 arguments are the same in each predict function. The latter 2 are specific to certain functions. For line transects, the effective strip half-width (esw=TRUE) is the integral of the fitted
detection function over either 0 to W or the specified int.range. The predicted detection probability is the average probability which is simply the integral divided by the distance range. For point transect models, esw=TRUE calculates the effective area of detection (commonly referred to as "nu", this is the integral of $2/width^2 \times rg(r)$.

Fitted detection probabilities are stored in the model object and these are returned unless compute=TRUE or newdata is specified. compute=TRUE is used to estimate numerical derivatives for use in delta method approximations to the variance.

For method="io.fi" or method="trial.fi" if integrate=FALSE, predict returns the value of the conditional detection probability and if integrate=TRUE, it returns the average conditional detection probability by integrating over x (distance) with respect to a uniform distribution.

Note that the ordering of the returned results when no new data is supplied (the "fitted" values) will not necessarily be the same as the data supplied to ddf, the data (and hence results from predict) will be sorted by object ID (object) then observer ID (observer).

Value

For all but the exceptions below, the value is a list with a single element: fitted, a vector of average detection probabilities or esw values for each observation in the original data or newdata.

For predict.ds, if se.fit=TRUE there is an additional element $se.fit$, which contains the standard errors of the probabilities of detection or ESW.

For predict.io.fi,predict.trial.fi,predict.rem.fi with integrate=TRUE, the value is a list with one element: fitted, which is a vector of integrated (average) detection probabilities for each observation in the original data or newdata.

For predict.io.fi,predict.trial.fi, or predict.rem.fi with integrate=FALSE, the value is a list with the following elements:

fitted $p(y)$ values

p1 $p_{1|2}(y)$, conditional detection probability for observer 1
p2 $p_{2|1}(y)$, conditional detection probability for observer 2

fitted $p(y) = p_{1|2}(y) + p_{2|1}(y) - p_{1|2}(y) \times p_{2|1}(y)$, conditional detection probability of being seen by either observer

Note

Each function is called by the generic function predict for the appropriate ddf model object. They can be called directly by the user, but it is typically safest to use predict which calls the appropriate function based on the type of model.

Author(s)

Jeff Laake, David L Miller

See Also

ddf, summary.ds, plot.ds
print.ddf

**Simple pretty printer for distance sampling analyses**

**Description**

Simply prints out summary of the model which was fitted. For more detailed information see `summary`.

**Usage**

```r
## S3 method for class 'ddf'
print(x, ...)
```

**Arguments**

- `x`: a ddf object
- `...`: not passed through, just for S3 compatibility.

**Author(s)**

David L. Miller

---

print.ddf.gof

**Prints results of goodness of fit tests for detection functions**

**Description**

Provides formatted output for results of goodness of fit tests: chi-square, Kolmogorv-Smirnov and Cramer-von Mises test as appropriate.

**Usage**

```r
## S3 method for class 'ddf.gof'
print(x, ...)
```

**Arguments**

- `x`: result of call to `ddf.gof`
- `...`: unused unspecified arguments for generic print

**Value**

None
**print.det.tables**

**Author(s)**

Jeff Laake

**See Also**

ddf.gof

---

**print.det.tables**  
*Print results of observer detection tables*

**Description**

Provides formatted output for detection tables

**Usage**

```r
## S3 method for class 'det.tables'
print(x, ...)
```

**Arguments**

- `x`       result of call to ddf
- `...`     unused unspecified arguments for generic print

**Value**

None

**Author(s)**

Jeff Laake

**See Also**

plot.det.tables
print.dht  
*Prints density and abundance estimates*

**Description**

Outputs summary statistics, abundance and density by region (if any) and optionally a correlation matrix if more than one region.

**Usage**

```r
## S3 method for class 'dht'
print(x, cor = FALSE, bysample = FALSE, vcmatrices = FALSE, ...)
```

**Arguments**

- `x`  
  dht object that results from call to dht for a specific ddf object
- `cor`  
  if TRUE outputs correlation matrix of estimates
- `bysample`  
  if TRUE, prints results for each sample
- `vcmatrices`  
  if TRUE, prints variance-covariance matrices
- `...`  
  unspecified and unused arguments for S3 consistency

**Value**

None

**Author(s)**

Jeff Laake

**See Also**

dht

---

print.p_dist_table  
*Print distribution of probabilities of detection*

**Description**

Just a pretty printer for the table of probabilities of detection.

**Usage**

```r
## S3 method for class 'p_dist_table'
print(x, digits = 2, ...)
```
Arguments

- `x` output from `p_dist_table`
- `digits` number of significant digits to print
- `...` other arguments to be passed to `print.data.frame`

Value

Just prints the table and the range of ps

Author(s)

David L Miller

Description

Provides a brief summary of data and fitted detection probability model parameters, model selection criterion, and optionally abundance in the covered (sampled) region and its standard error. What is printed depends on the corresponding call to `summary`.

Usage

```r
## S3 method for class 'summary.ds'
print(x, ...)
```

Arguments

- `x` a summary of ddf model object
- `...` unspecified and unused arguments for S3 consistency

Author(s)

Jeff Laake

See Also

`summary.ds`
print.summary.io

Print summary of distance detection function model object

Description

Provides a brief summary of data and fitted detection probability model parameters, model selection criterion, and optionally abundance in the covered (sampled) region and its standard error. What is printed depends on the corresponding call to summary.

Usage

## S3 method for class 'summary.io'
print(x, ...)

Arguments

x  
a summary of ddf model object

...  
unspecified and unused arguments for S3 consistency

Author(s)

Jeff Laake

See Also

summary.io

print.summary.io.fi

Print summary of distance detection function model object

Description

Provides a brief summary of data and fitted detection probability model parameters, model selection criterion, and optionally abundance in the covered (sampled) region and its standard error. What is printed depends on the corresponding call to summary.

Usage

## S3 method for class 'summary.io.fi'
print(x, ...)

Arguments

x  
a summary of ddf model object

...  
unspecified and unused arguments for S3 consistency
Author(s)

Jeff Laake

See Also

summary.io.fi

Description

Provides a brief summary of data and fitted detection probability model parameters, model selection
criterion, and optionally abundance in the covered (sampled) region and its standard error. What is
printed depends on the corresponding call to summary.

Usage

## S3 method for class 'summary.rem'
print(x, ...)

Arguments

x a summary of ddf model object

... unspecified and unused arguments for S3 consistency

Author(s)

Jeff Laake

See Also

summary.rem
print.summary.trial

---

**print.summary.rem.fi**  
*Print summary of distance detection function model object*

---

**Description**

Provides a brief summary of data and fitted detection probability model parameters, model selection criterion, and optionally abundance in the covered (sampled) region and its standard error. What is printed depends on the corresponding call to summary.

**Usage**

```r
## S3 method for class 'summary.rem.fi'
print(x, ...)
```

**Arguments**

- `x`: a summary of ddf model object
- `...`: unspecified and unused arguments for S3 consistency

**Author(s)**

Jeff Laake

**See Also**

- `summary.rem.fi`

---

**print.summary.trial**  
*Print summary of distance detection function model object*

---

**Description**

Provides a brief summary of data and fitted detection probability model parameters, model selection criterion, and optionally abundance in the covered (sampled) region and its standard error. What is printed depends on the corresponding call to summary.

**Usage**

```r
## S3 method for class 'summary.trial'
print(x, ...)
```

**Arguments**

- `x`: a summary of ddf model object
- `...`: unspecified and unused arguments for S3 consistency
Author(s)
Jeff Laake

See Also
summary.trial

Description
Provides a brief summary of data and fitted detection probability model parameters, model selection
criterion, and optionally abundance in the covered (sampled) region and its standard error. What is
printed depends on the corresponding call to summary.

Usage

```r
## S3 method for class 'summary.trial.fi'
print(x, ...)
```

Arguments

- `x` a summary of ddf model object
- `...` unspecified and unused arguments for S3 consistency

Author(s)
Jeff Laake

See Also
summary.trial.fi
prob.deriv

Derivatives for variance of average p and average p(0) variance

Description

Used in call to DeltaMethod from prob.se to get first derivatives

Usage

prob.deriv(par, model, parfct, observer = NULL, fittedmodel = NULL)

Arguments

- **par**: detection function parameter values
- **model**: ddf model object
- **parfct**: function of detection probabilities; currently only average (over covariates) detection probability p integrated over distance or average (over covariates) detection probability at distance 0; p(0)
- **observer**: 1,2,3 for primary, secondary, or duplicates for average p(0); passed to fct
- **fittedmodel**: full fitted ddf model when trial.fi or io.fi is called from trial or io respectively

Details

Need to add equations here as I do not think they exist in any of the texts. These should probably be checked with simulation.

Value

Vector of values from fct at specified parameter values

Author(s)

Jeff Laake

See Also

prob.se
Description
Computes components of variance for average $p=n/N$ and average $p(0)$ with weights based on empirical covariate distribution, if it contains covariates.

Usage
`prob.se(model, fct, vcov, observer = NULL, fittedmodel = NULL)`

Arguments
- `model`: ddf model object
- `fct`: function of detection probabilities; currently only average (over covariates) detection probability $p$ integrated over distance or average (over covariates) detection probability at distance 0; $p(0)$
- `vcov`: variance-covariance matrix of parameter estimates
- `observer`: 1,2,3 for primary, secondary, or duplicates for average $p(0)$; passed to fct
- `fittedmodel`: full fitted ddf model when `trial.fi` or `io.fi` is called from `trial` or `io` respectively

Details
Need to add equations here as I do not think they exist in any of the texts. These should probably be checked with simulation.

Value
- `var`: variance
- `partial`: partial derivatives of parameters with respect to fct
- `covar`: covariance of $n$ and average $p$ or $p(0)$

Author(s)
Jeff Laake

See Also
`prob.deriv`
process.data

Process data for fitting distance sampling detection function

Description

Sets up dataframe and does some basic error checking. Adds needed fields to dataframe and to meta.data.

Usage

process.data(data, meta.data = list(), check = TRUE)

Arguments

data: dataframe object
meta.data: meta.data options; see ddf for a description
check: if TRUE check data for errors in the mrds structure; for method="ds" check=FALSE

Details

The function does a number of error checking tasks, creating fields and adding to meta.data including:

1) If check=TRUE, check to make sure the record structure is okay for mrds data. The number of primary records (observer=1) must equal the number of secondary records (observer=2). Also, a field in the dataframe is created timesseen which counts the number of times an object was detected 0,1,2; if timesseen=0 then the record is tossed from the analysis. Also if there are differences in the data (distance, size, covariates) for observer 1 and 2 a warning is issued that the analysis may fail. The code assumes these values are the same for both observers.

2) Based on the presence of fields distbegin and distend, a determination is made of whether the data analysis should be based on binned distances and a field binned is created, which is TRUE if the distance for the observation is binned. By assigning for each observation this allows an analysis of a mixture of binned and unbinned distances.

4) Data are restricted such that distances are not greater than width and not less than left if those values are specified in meta.data. If they are not specified then left defaults to 0 and width defaults to the largest distance measurement.

5) Determine if an integration range (int.begin and int.end has been specified for the observations. If it has, add the structure to meta.data. The integration range is typically used for aerial surveys in which the altitude varies such that the strip width (left to width) changes with a change in altitude.

6) Fields defined as factors are cleaned up such that any unused levels are eliminated.

7) If the restrictions placed on the data, eliminated all of the data, the function stops with an error message.
Value

- `xmat`  processed data frame with added fields
- `meta.data`  meta.data list

Author(s)

Jeff Laake

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**pronghorn**  

*Pronghorn aerial survey data from Wyoming*

**Description**

Detections of pronghorn from fixed-wing aerial surveys in Southeastern Wyoming using four angular bins defined by strut marks. Illustrates data where altitude above ground level (AGL) varies during the survey.

**Format**

A data frame with 660 observations on the following 5 variables.

- **STRATUM**  a numeric vector
- **direction**  a factor with levels `N S` representing the survey direction
- **AGL**  height above ground level
- **Band**  a factor with levels `A B C D` which represent angular bands between breaks at 35.42, 44.56, 51.52, 61.02, 70.97 degrees. These angles were set based on selected distance bins based on the target AGL.
- **cluster**  number of pronghorn in the observed cluster

**Details**

Each record is an observed cluster of pronghorn. The data provide the stratum for the observation, the direction of travel, the AGL at the time of the observation, the angular bin which contained the center of the pronghorn cluster(group), and the number of pronghorn in the group. The angular bins were defined by a combination of two window and five wing strut marks to define bin cutpoints for perpendicular ground distances of 0-65, 65-90, 90-115, 115-165 and 165-265 meters when the plane is 300' (91.4 meters) above ground level. The inner band is considered a blind region due to obstruction of view beneath the plane; thus the line is offset 65 meters from underneath the plane.

**Source**

Data provided courtesy of Rich Guenzel of Wyoming Game and Fish.

**References**

ptdata.dual  

**Description**

Simulated dual observer point count data with detection \( p(0)=0.8 \); \( \text{hn} \) \( \sigma=30 \); \( w=100 \) for both observers with dependency \( \gamma>0 \), \( \gamma=0.1 \)

**Format**

The format is 420 obs of 6 variables: distance: numeric distance from center observer: Factor w/ 2 levels "1","2": 1 2 1 2 1 2 1 2 ... detected: numeric 0/1 person: Factor with 2 levels A,B pair: Factor with 2 levels "AB" BA $ object : sequential object number
Examples

```r
data(ptdata.dual)
xx <- ddf(mrmodel=~glm(formula=~distance),
          dsmodel = ~cds(key="hn", formula = ~1),
          data = ptdata.dual, method = "io", meta.data = list(point=TRUE))
summary(xx)
plot(xx, main="Simulated point count data")
```

Description

Simulated removal observer point count data with detection p(0)=0.8; hn sigma=30; w=100 for both observers with dependency y>0, gamma=0.1

Format

The format is 408 obs of 6 variables: distance: numeric distance from center observer: Factor w/ 2 levels "1","2": 1 2 1 2 1 2 1 2 ... detected: numeric 0/1 person: Factor with 2 levels A,B pair: Factor with 2 levels "AB" BA" object: sequential object number

Examples

```r
data(ptdata.removal)
xx <- ddf(mrmodel=~glm(formula=~distance),
          dsmodel = ~cds(key="hn", formula = ~1),
          data = ptdata.removal, method = "rem",
          meta.data = list(point=TRUE))
summary(xx)
plot(xx, main="Simulated point count data")
```

ptdata.removal  Simulated removal observer point count data

Description

Simulated single observer point count data with detection p(0)=1; hn sigma=30; w=100

Format

The format is 341 obs of 4 variables: ..$ distance: numeric distance from center $ observer: Factor w/ 2 levels "1","2": 1 2 1 2 1 2 1 2 ... ..$ detected: numeric 0/1 $ object : sequential object number
Examples

data(ptdata.single)
xx=ddf(dsmodel = ~cds(key="hn", formula = ~1), data = ptdata.single,
       method = "ds", meta.data = list(point=TRUE))
summary(xx)
plot(xx,main="Simulated point count data")

p_dist_table

Distribution of probabilities of detection

Description

Generate a table of frequencies of probability of detection from a detection function model. This is particularly useful when employing covariates, as it can indicate if there are detections with very small detection probabilities that can be unduly influential when calculating abundance estimates.

Usage

p_dist_table(object, bins = seq(0, 1, by = 0.1), proportion = FALSE)

Arguments

object         fitted detection function
bins           how the results should be binned
proportion     should proportions be returned as well as counts?

Details

Because dht uses a Horvitz-Thompson-like estimator, abundance estimates can be sensitive to errors in the estimated probabilities. The estimator is based on \( \sum 1/\hat{P}_a(z_i) \), which means that the sensitivity is greater for smaller detection probabilities. As a rough guide, we recommend that the method be not used if more than say 5% of the \( \hat{P}_a(z_i) \) are less than 0.2, or if any are less than 0.1. If these conditions are violated, the truncation distance \( w \) can be reduced. This causes some loss of precision relative to standard distance sampling without covariates.

Value

a data.frame with probability bins, counts and (optionally) proportions. The object has an attribute \( p\_range \) which contains the range of estimated detection probabilities

Author(s)

David L Miller
References


Examples

```r
## Not run:
# try out the tee data
data(book.tee.data)
egdata <- book.tee.data$book.tee.dataframe
# fit model with covariates
result <- ddf(dsmodel = ~mcds(key = "hn", formula = ~sex+size),
               data = egdata[egdata$observer==1, ], method = "ds",
               meta.data = list(width = 4))
# print table
p_dist_table(result)
# with proportions
p_dist_table(result, proportion=TRUE)
## End(Not run)
```

Description

Constructs a quantile-quantile (Q-Q) plot for fitted model as a graphical check of goodness of fit. Formal goodness of fit testing for detection function models using Kolmogorov-Smirnov and Cramer-von Mises tests. Both tests are based on looking at the quantile-quantile plot produced by `qqplot.ddf` and deviations from the line x=y.

Usage

`qqplot.ddf(model, plot = TRUE, nboot = 100, ks = FALSE, ...)`

Arguments

- `model`: fitted distance detection function model object
- `plot`: the Q-Q plot be plotted or just report statistics?
- `nboot`: number of replicates to use to calculate p-values for the goodness of fit test statistics
- `ks`: perform the Kolmogorov-Smirnov test (this involves many bootstraps so can take a while)
- `...`: additional arguments passed to `plot`
Details

The Kolmogorov-Smirnov test asks the question "what's the largest vertical distance between a point and the y=x line?" It uses this distance as a statistic to test the null hypothesis that the samples (EDF and CDF in our case) are from the same distribution (and hence our model fits well). If the deviation between the y=x line and the points is too large we reject the null hypothesis and say the model doesn’t have a good fit.

Rather than looking at the single biggest difference between the y=x line and the points in the Q-Q plot, we might prefer to think about all the differences between line and points, since there may be many smaller differences that we want to take into account rather than looking for one large deviation. Its null hypothesis is the same, but the statistic it uses is the sum of the deviations from each of the point to the line.

Value

A list of goodness of fit related values:

- `edf` matrix of lower and upper empirical distribution function values
- `cdf` fitted cumulative distribution function values
- `ks` list with K-S statistic ($D_n$) and p-value ($p$)
- `CvM` list with CvM statistic ($W$) and p-value ($p$)

Details

Note that a bootstrap procedure is required to ensure that the p-values from the procedure are correct as we are comparing the cumulative distribution function (CDF) and empirical distribution function (EDF) and we have estimated the parameters of the detection function.

Author(s)

Jeff Laake, David L Miller

References


See Also

`ddf.gof`, `cdf.ds`
Iterative offset model fitting of mark-recapture with removal model

**Description**
Detection function fitting from mark-recapture data with a removal configuration in which a secondary observer knows what the primary observer detects and detects objects missed by the primary observer. The iterative offset glm/gam uses an offset to compensate for the conditioning on the set of objects seen by either observer (eg 00 those missed by both observers are not included in the analysis. This function is similar to `io.glm`.

**Usage**

```r
rem.glm(
  datavec, fitformula, eps = 1e-05, iterlimit = 500, GAM = FALSE, gamplot = TRUE, datavec2
)
```

**Arguments**

- `datavec`: dataframe containing records seen by either observer 1 or 2
- `fitformula`: logit link formula
- `eps`: convergence criterion
- `iterlimit`: maximum number of iterations allowed
- `GAM`: uses GAM instead of GLM for fitting
- `gamplot`: set to TRUE to get a gam plot object if GAM=TRUE
- `datavec2`: dataframe containing all records for observer 1 and observer 2 as in io.glm form; this is used in case there is an observer(not platform effect)

**Details**
The only difference between this function and `io.glm` is the offset and the data construction because there is only one detection function being estimated for the primary observer. The two functions could be merged.

**Value**

- `glmobj`: GLM or GAM object
- `offsetvalue`: offsetvalues from iterative fit
- `plotobj`: gam plot object (if GAM & gamplot==TRUE, else NULL)
Note
currently the code in this function for GAMs has been commented out until the remainder of the
mrd package will work with GAMs.

Author(s)
Jeff Laake

References
Oxford University Press.

rescale_pars Calculate the parameter rescaling for parameters associated with co-
variates

Description
This will calculate the rescaling needed when covariates to be included in the scale of the detection
function are “too big”. Based on code from optimx.

Usage
rescale_pars(initialvalues, ddfobj)

Arguments
initialvalues starting values for the optimisation
ddfobj detection function object

Details
Derivative-free methods like nlminb are sensitive to the parameters being poorly scaled. This can
also cause problems for quasi-Newton methods too (at least, bad scaling won’t _help_ the optimi-
sation). So here we rescale the parameters if necessary (unless we already got scaling from control)

Author(s)
David L Miller
sample_ddf

Generate data from a fitted detection function and refit the model

Description

Generate data from a fitted detection function and refit the model

Usage

sample_ddf(ds.object)

Arguments

ds.object a fitted detection function object

Note

This function changes the random number generator seed. To avoid any potential side-effects, use something like: seed <- get(".Random.seed", envir=.GlobalEnv) before running code and assign(".Random.seed", seed, envir=.GlobalEnv) after.

Author(s)

David L. Miller

setbounds

Set parameter bounds

Description

Set values of lower and upper bounds and check lengths of any user-specified values

Usage

setbounds(lowerbounds, upperbounds, initialvalues, ddfobj)

Arguments

lowerbounds vector of lower bounds
upperbounds vector of upper bounds
initialvalues vector of initial parameter estimates
ddfobj distance detection function object
Value

<table>
<thead>
<tr>
<th>lower</th>
<th>vector of lower bounds</th>
</tr>
</thead>
<tbody>
<tr>
<td>upper</td>
<td>vector of upper bounds</td>
</tr>
<tr>
<td>setlower</td>
<td>logical indicating whether user set lower bounds</td>
</tr>
<tr>
<td>setupper</td>
<td>logical indicating whether user set upper bounds</td>
</tr>
</tbody>
</table>

Author(s)

Jeff Laake

---

setcov

*Creates design matrix for covariates in detection function*

Description

This function creates a design matrix for the g(0) or scale covariates using the input model formula. It returns a list which contains 2 elements: 1) dim: the dimension (number of columns) of the design matrix, and 2) cov: the constructed design matrix. This function is relatively simple because it uses the built-in function `model.matrix` which does the majority of the work. This function handles 2 exceptions "~.", the null model with 0 columns and "~1" the intercept only model - a column of 1s. If a model other than the 2 exceptions is provided, it calls `model.matrix` to construct the columns. If any of the columns of the design matrix are all 0's the column is removed. This occurs when there is no data for a particular factor.

Usage

`setcov(dmat, model)`

Arguments

<table>
<thead>
<tr>
<th>dmat</th>
<th>data matrix</th>
</tr>
</thead>
<tbody>
<tr>
<td>model</td>
<td>model formula</td>
</tr>
</tbody>
</table>

Value

a design matrix for the specified data and model

Author(s)

Jeff Laake
**setinitial.ds**

*Set initial values for detection function based on distance sampling*

**Description**

For a given detection function, it computes the initial values for the parameters including scale and shape parameters and adjustment function parameters if any. If there are user-defined initial values only the parameters not specified by the user are computed.

**Usage**

```
setinitial.ds(ddfobj, width, initial, point, left)
sethazard(ddfobj, dmat, width, left)
```

**Arguments**

- **ddfobj**: distance detection function object
- **width**: half-width of transect or radius of point count
- **initial**: list of user-defined initial values with possible elements scale, shape, adjustment
- **point**: if TRUE, point count data; otherwise, line transect data
- **left**: left truncation
- **dmat**: xmat from ddfobj

**Value**

- **scale**: vector of initial scale parameter values
- **shape**: vector of initial shape parameter values
- **adjustment**: vector of initial adjustment function parameter values

**Author(s)**

Jeff Laake, David L Miller

**sim.mix**

*Simulation of distance sampling data via mixture models Allows one to simulate line transect distance sampling data using a mixture of half-normal detection functions.*

**Description**

Simulation of distance sampling data via mixture models Allows one to simulate line transect distance sampling data using a mixture of half-normal detection functions.
Usage

\texttt{sim.mix(n, sigma, mix.prop, width, means = 0)}

Arguments

- \textit{n} \hspace{1cm} \text{number of samples to generate}
- \textit{sigma} \hspace{1cm} \text{vector of scale parameters}
- \textit{mix.prop} \hspace{1cm} \text{vector of mixture proportions (same length as sigma)}
- \textit{width} \hspace{1cm} \text{truncation}
- \textit{means} \hspace{1cm} \text{vector of means (used to generate wacky, non-monotonic data)}

Value

- \textit{distances} \hspace{1cm} \text{a vector of distances}

Note

At the moment this is TOTALLY UNSUPPORTED! Please don’t use it for anything important!

Author(s)

David Lawrence Miller

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\textbf{solvecov} \hspace{1cm} \textit{Invert of covariance matrices}

Description

Tries to invert a matrix by \texttt{solve}. If this fails because of singularity, an eigenvector decomposition is computed, and eigenvalues below $1/c_{\text{max}}$ are replaced by $1/c_{\text{max}}$, i.e., $c_{\text{max}}$ will be the corresponding eigenvalue of the inverted matrix.

Usage

\texttt{solvecov(m, cmax = 1e+10)}

Arguments

- \textit{m} \hspace{1cm} \text{a numeric symmetric matrix.}
- \textit{cmax} \hspace{1cm} \text{a positive value, see above.}

Value

A list with the following components: \texttt{inv} the inverted matrix, \texttt{coll} \texttt{TRUE} if \texttt{solve} failed because of singularity.
stake77

Source

solvecov code was taken from package fpc: Christian Hennig [http://www.homepages.ucl.ac.uk/~ucakche/](http://www.homepages.ucl.ac.uk/~ucakche/)

Author(s)

Christian Hennig [http://www.homepages.ucl.ac.uk/~ucakche/](http://www.homepages.ucl.ac.uk/~ucakche/)

See Also

solve, eigen

stake77

Wooden stake data from 1977 survey

Description

Multiple surveys by different observers of a single 1km transect containing 150 wooden stakes placed randomly throughout a 40 m strip (20m on either side).

Format

A data frame with 150 observations on the following 10 variables.

StakeNo  unique number for each stake 1-150
PD  perpendicular distance at which the stake was placed from the line
Obs1 0/1 whether missed/seen by observer 1
Obs2 0/1 whether missed/seen by observer 2
Obs3 0/1 whether missed/seen by observer 3
Obs4 0/1 whether missed/seen by observer 4
Obs5 0/1 whether missed/seen by observer 5
Obs6 0/1 whether missed/seen by observer 6
Obs7 0/1 whether missed/seen by observer 7
Obs8 0/1 whether missed/seen by observer 8

Source


References

Examples

data(stake77)
# Extract functions for stake data and put in the mrds format
extract.stake <- function(stake,obs){
  extract.obs <- function(obs){
    example <- subset(stake,eval(parse(text=paste("Obs",obs,"==1",sep=""))),
                     select="PD")
    example$distance <- example$PD
    example$object <- 1:nrow(example)
    example$PD <- NULL
    return(example)
  }
  if(obs!="all"){
    return(extract.obs(obs=obs))
  }else{
    example <- NULL
    for(i in 1:(ncol(stake)-2)){
      df <- extract.obs(obs=i)
      df$person <- i
      example <- rbind(example,df)
    }
    example$person <- factor(example$person)
    example$object <- 1:nrow(example)
    return(example)
  }
}

extract.stake.pairs <- function(stake,obs1,obs2,removal=FALSE){
  obs1 <- paste("Obs",obs1,sep="")
  obs2 <- paste("Obs",obs2,sep="")
  example <- subset(stake,eval(parse(text=paste(obs1,"==1 |",obs2,"==1 ", sep=""))),select=c("PD",obs1,obs2))
  names(example) <- c("distance","obs1","obs2")
  detected <- c(example$obs1,example$obs2)
  example <- data.frame(object = rep(1:nrow(example),2),
                       distance = rep(example$distance,2),
                       detected = detected,
                       observer = c(rep(1,nrow(example)),rep(2,nrow(example))))
  if(removal) example$detected[example$observer==2] <- 1
  return(example)
}

# extract data for observer 1 and fit a single observer model
stakes <- extract.stake(stake77,1)
ds.model <- ddf(dsmodel = ~mcds(key = "hn", formula = ~1), data = stakes,
               method = "ds", meta.data = list(width = 20))
plot(ds.model,breaks=seq(0,20,2),showpoints=TRUE)
ddf.gof(ds.model)

# extract data from observers 1 and 3 and fit an io model
stkpairs <- extract.stake.pairs(stake77,1,3,removal=FALSE)
io.model <- ddf(dsmodel = ~mcds(key = "hn", formula=~1),
                mrmodel=~glm(formula=-distance),
                method="ds")
stake78

Wooden stake data from 1978 survey

Description

Multiple surveys by different observers of a single 1km transect containing 150 wooden stakes placed based on expected uniform distribution throughout a 40 m strip (20m on either side).

Format

A data frame with 150 observations on the following 13 variables.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>StakeNo</td>
<td>unique number for each stake 1-150</td>
</tr>
<tr>
<td>PD</td>
<td>perpendicular distance at which the stake was placed from the line</td>
</tr>
<tr>
<td>Obs1</td>
<td>0/1 whether missed/seen by observer 1</td>
</tr>
<tr>
<td>Obs2</td>
<td>0/1 whether missed/seen by observer 2</td>
</tr>
<tr>
<td>Obs3</td>
<td>0/1 whether missed/seen by observer 3</td>
</tr>
<tr>
<td>Obs4</td>
<td>0/1 whether missed/seen by observer 4</td>
</tr>
<tr>
<td>Obs5</td>
<td>0/1 whether missed/seen by observer 5</td>
</tr>
<tr>
<td>Obs6</td>
<td>0/1 whether missed/seen by observer 6</td>
</tr>
<tr>
<td>Obs7</td>
<td>0/1 whether missed/seen by observer 7</td>
</tr>
<tr>
<td>Obs8</td>
<td>0/1 whether missed/seen by observer 8</td>
</tr>
<tr>
<td>Obs9</td>
<td>0/1 whether missed/seen by observer 9</td>
</tr>
<tr>
<td>Obs10</td>
<td>0/1 whether missed/seen by observer 10</td>
</tr>
<tr>
<td>Obs11</td>
<td>0/1 whether missed/seen by observer 11</td>
</tr>
</tbody>
</table>

Details

The 1997 survey was based on a single realization of a uniform distribution. Because it was a single transect and there was no randomization of the distances for each survey, we repeated the experiment and used distances that provided a uniform distribution but randomly sorted the positions along the line so there was no pattern obvious to the observer.

Source

References


Examples

data(stake78)
data(stake77)
# compare distribution of distances for all stakes
hist(stake77$PD)
hist(stake78$PD)
# Extract stake data and put in the mrds format for model fitting.
extact.stake <- function(stake,obs){
exact.obs <- function(obs){
exact <- subset(stake,eval(parse(text=paste("Obs",obs,"==1",sep=""))),
     select="PD")
exact$distance <- exact$PD
exact$object <- 1:nrow(exact)
exact$PD <- NULL
return(exact)
}
if(obs!="all"){
    return(exact.obs(obs=obs))
}else{
    example <- NULL
    for(i in 1:(ncol(stake)-2)){
        df <- exact.obs(obs=i)
        df$person <- i
        example <- rbind(example,df)
    }
    example$person <- factor(example$person)
    example$object <- 1:nrow(example)
    return(example)
}
estact.stake.pairs <- function(stake,obs1,obs2,removal=FALSE){
obs1 <- paste("Obs",obs1,sep="")
obs2 <- paste("Obs",obs2,sep="")
example <- subset(stake,eval(parse(text=paste(obs1,"==1 | ",obs2,"==1 ",sep=""))), select=c("PD",obs1,obs2))
names(example) <- c("distance","obs1","obs2")
detected <- c(example$obs1,example$obs2)
example <- data.frame(object=rep(1:nrow(example),2),
distance=rep(example$distance,2),
detected = detected,
observer=c(rep(1,nrow(example)),
rep(2,nrow(example))))
if(removal) example$detected[example$observer==2] <- 1
return(example)"
# extract data for observer 10 and fit a single observer model
stakes <- extract.stake(stake78,10)
ds.model <- ddf(dsmodel = ~mcds(key = "hn", formula = ~1), data = stakes,
    method = "ds", meta.data = list(width = 20))
plot(ds.model,breaks=seq(0,20,2),showpoints=TRUE)
ddf.gof(ds.model)

# extract data from observers 5 and 7 and fit an io model
stkpairs <- extract.stake.pairs(stake78,5,7,removal=FALSE)
io.model <- ddf(dsmodel = ~mcds(key = "hn", formula=~1),
    mrmodel=~glm(formula=~distance),
    data = stkpairs, method = "io")
summary(io.model)
par(mfrow=c(3,2))
plot(io.model,breaks=seq(0,20,2),showpoints=TRUE,new=FALSE)
ddf.gof(io.model)

summary.ds

**Summary of distance detection function model object**

**Description**

Provides a brief summary of data and fitted detection probability model parameters, model selection criterion, and optionally abundance in the covered (sampled) region and its standard error.

**Usage**

```r
# S3 method for class 'ds'
summary(object, se = TRUE, N = TRUE, ...)
```

**Arguments**

- `object` a ddf model object
- `se` if TRUE, computes standard errors
- `N` if TRUE, computes abundance in covered (sampled) region
- `...` unspecified and unused arguments for S3 consistency

**Details**

The argument `N` is used to suppress computation of abundance and average detection probability in calls to summarize the ds and either io.fi or trial.fi for summaries of io and trial objects respectively which are composed of a ds model object and a mark-recapture model object. The corresponding print function is called to print the summary results.

**Value**

list of extracted and summarized objects
Note

This function is called by the generic function summary for any ddf model object. Each function can be called directly by the user, but it is typically safest to use the generic function summary which calls the appropriate function based on the type of ddf model.

Author(s)

Jeff Laake

summary.io Summary of distance detection function model object

Description

Provides a brief summary of data and fitted detection probability model parameters, model selection criterion, and optionally abundance in the covered (sampled) region and its standard error.

Usage

```r
## S3 method for class 'io'
summary(object, se = TRUE, ...)
```

Arguments

- `object` a ddf model object
- `se` if TRUE, computes standard errors
- `...` unspecified and unused arguments for S3 consistency

Details

The argument N is used to suppress computation of abundance and average detection probability in calls to summarize the ds and either io.fi or trial.fi for summaries of io and trial objects respectively which are composed of a ds model object and a mark-recapture model object. The corresponding print function is called to print the summary results.

Value

list of extracted and summarized objects

Note

This function is called by the generic function summary for any ddf model object. Each function can be called directly by the user, but it is typically safest to use the generic function summary which calls the appropriate function based on the type of ddf model.

Author(s)

Jeff Laake
Description

Provides a brief summary of data and fitted detection probability model parameters, model selection criterion, and optionally abundance in the covered (sampled) region and its standard error.

Usage

```r
## S3 method for class 'io.fi'
summary(object, se = TRUE, N = TRUE, fittedmodel = NULL, ddfobj = NULL, ...)
```

Arguments

- `object`: a `ddf` model object
- `se`: if `TRUE`, computes standard errors
- `N`: if `TRUE`, computes abundance in covered (sampled) region
- `fittedmodel`: full fitted model when called from `trial` or `io`
- `ddfobj`: distance sampling object description
- `...`: unspecified and unused arguments for S3 consistency

Details

The argument `N` is used to suppress computation of abundance and average detection probability in calls to summarize the `ds` and either `io.fi` or `trial.fi` for summaries of `io` and `trial` objects respectively which are composed of a `ds` model object and a mark-recapture model object. The corresponding print function is called to print the summary results.

Value

list of extracted and summarized objects

Note

This function is called by the generic function `summary` for any `ddf` model object. Each function can be called directly by the user, but it is typically safest to use the generic function `summary` which calls the appropriate function based on the type of `ddf` model.

Author(s)

Jeff Laake
Summary of distance detection function model object

Description

Provides a brief summary of data and fitted detection probability model parameters, model selection criterion, and optionally abundance in the covered (sampled) region and its standard error.

Usage

```r
## S3 method for class 'rem'
summary(object, se = TRUE, ...)
```

Arguments

- `object`: a `ddf` model object
- `se`: if TRUE, computes standard errors
- `...`: unspecified and unused arguments for S3 consistency

Details

The argument `N` is used to suppress computation of abundance and average detection probability in calls to summarize the `ds` and either `io.fi` or `trial.fi` for summaries of `io` and `trial` objects respectively which are composed of a `ds` model object and a mark-recapture model object. The corresponding print function is called to print the summary results.

Value

list of extracted and summarized objects

Note

This function is called by the generic function `summary` for any `ddf` model object. Each function can be called directly by the user, but it is typically safest to use the generic function `summary` which calls the appropriate function based on the type of `ddf` model.

Author(s)

Jeff Laake
**Summary of distance detection function model object**

**Description**

Provides a brief summary of data and fitted detection probability model parameters, model selection criterion, and optionally abundance in the covered (sampled) region and its standard error.

**Usage**

```r
## S3 method for class 'rem.fi'
summary(object, se = TRUE, N = TRUE, fittedmodel = NULL, ...)
```

**Arguments**

- `object`: a `ddf` model object
- `se`: if TRUE, computes standard errors
- `N`: if TRUE, computes abundance in covered (sampled) region
- `fittedmodel`: full fitted model when called from `trial` or `io`
- `...`: unspecified and unused arguments for S3 consistency

**Details**

The argument `N` is used to suppress computation of abundance and average detection probability in calls to summarize the `ds` and either `io.fi` or `trial.fi` for summaries of `io` and `trial` objects respectively which are composed of a `ds` model object and a mark-recapture model object. The corresponding print function is called to print the summary results.

**Value**

- list of extracted and summarized objects

**Note**

This function is called by the generic function `summary` for any `ddf` model object. Each function can be called directly by the user, but it is typically safest to use the generic function `summary` which calls the appropriate function based on the type of `ddf` model.

**Author(s)**

Jeff Laake
Summary of distance detection function model object

Description

Provides a brief summary of data and fitted detection probability model parameters, model selection criterion, and optionally abundance in the covered (sampled) region and its standard error.

Usage

```r
## S3 method for class 'trial'
summary(object, se = TRUE, ...)
```

Arguments

- `object`: a ddf model object
- `se`: if TRUE, computes standard errors
- `...`: unspecified and unused arguments for S3 consistency

Details

The argument `N` is used to suppress computation of abundance and average detection probability in calls to summarize the `ds` and either `io.fi` or `trial.fi` for summaries of `io` and `trial` objects respectively which are composed of a `ds` model object and a mark-recapture model object. The corresponding print function is called to print the summary results.

Value

list of extracted and summarized objects

Note

This function is called by the generic function `summary` for any ddf model object. Each function can be called directly by the user, but it is typically safest to use the generic function `summary` which calls the appropriate function based on the type of ddf model.

Author(s)

Jeff Laake
Summary of distance detection function model object

**Description**

Provides a brief summary of data and fitted detection probability model parameters, model selection criterion, and optionally abundance in the covered (sampled) region and its standard error.

**Usage**

```r
## S3 method for class 'trial.fi'
summary(object, se = TRUE, N = TRUE, fittedmodel = NULL, ...)
```

**Arguments**

- `object`: a ddf model object
- `se`: if TRUE, computes standard errors
- `N`: if TRUE, computes abundance in covered (sampled) region
- `fittedmodel`: full fitted model when called from `trial` or `io`
- `...`: unspecified and unused arguments for S3 consistency

**Details**

The argument `N` is used to suppress computation of abundance and average detection probability in calls to summarize the `ds` and either `io.fi` or `trial.fi` for summaries of `io` and `trial` objects respectively which are composed of a `ds` model object and a mark-recapture model object. The corresponding print function is called to print the summary results.

**Value**

list of extracted and summarized objects

**Note**

This function is called by the generic function `summary` for any ddf model object. Each function can be called directly by the user, but it is typically safest to use the generic function `summary` which calls the appropriate function based on the type of ddf model.

**Author(s)**

Jeff Laake
```r
survey.region.dht

Extrapolate Horvitz-Thompson abundance estimates to entire surveyed region

Description
Extrapolate Horvitz-Thompson abundance estimates to entire surveyed region

Usage
survey.region.dht(Nhat.by.sample, samples, width, point)

Arguments
Nhat.by.sample dataframe of abundance by sample
samples samples table
width transect width
point if TRUE point count otherwise line transect

Value
Revised Nhat.by.sample dataframe containing estimates extrapolated to survey region

Note
Internal function called by dht and related functions.

Author(s)
Jeff Laake
```

```r
test.breaks

Test validity for histogram breaks(cutpoints)

Description
Determines whether user specified breaks for histograms are properly ordered and match the left and right truncation.

Usage
test.breaks(breaks, left, width)
```
varn

Arguments

breaks          vector of cutpoints (breaks) for distance histogram
left            left truncation value
width           right truncation value; either radius of point count or half-width of transect

Value

vector of breaks modified to be valid if necessary

Author(s)

Jeff Laake

---

varn    Compute empirical variance of encounter rate

Description

Computes one of a series of possible variance estimates for the observed encounter rate for a set of sample measurements (e.g., line lengths) and number of observations per sample.

Usage

varn(lvec,nvec,type)

covn(lvec, groups1, groups2, type)

Arguments

lvec            vector of sample measurements (e.g., line lengths)
nvec            vector of number observed
type            choice of variance estimator to use for encounter rate
groups1         vector of number of groups observed
groups2         vector of number of individuals observed

Details

The choice of type follows the notation of Fewster et al. (2009) in that there are 8 choices of encounter rate variance that can be computed for lines and one for points:

R2  random line placement with unequal line lengths (design-assisted estimator)
R3  random line placement, model-assisted estimator, based on true contagion process
R4  random line placement, model-assisted estimator, based on apparent contagion process
S1  systematic line placement, post-stratification with no strata overlap
S2  systematic line placement, post-stratification with no strata overlap, variances weighted by line length per stratum
01  systematic line placement, post-stratification with overlapping strata (akin to S1)
02  systematic line placement, post-stratification with overlapping strata (weighted by line length per stratum, akin to S2)
03  systematic line placement, post-stratification with overlapping strata, model-assisted estimator with trend in encounter rate with line length
P2  random point placement, potentially unequal number of visits per point, design-based estimator
P3  random point placement, potentially unequal number of visits per point, model-based estimator

Default value is "R2", shown in Fewster et al. (2009) to have good performance for completely random designs for lines. For systematic parallel line transect designs, Fewster et al. recommend "O2". For point transects the default (and currently only implemented option) is "P3".

For the systematic estimators, pairs are assigned in the order they are given in the lengths and groups vectors.

Value

Variance of encounter rate as defined by arguments

Note

This function is also used with different calling arguments to compute Innes et al variance of the estimated abundances/length rather than observation encounter rate. The function covn is probably only valid for R3 and R2. Currently, the R2 form is used for all types other than R3.

Author(s)

Jeff Laake, David L Miller

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