

# Package ‘`ea3`’

July 4, 2023

**Title** Wildlife Mortality Estimator for Low Fatality Rates and Imperfect Detection

**Version** 1.0.0.1

**Date** 2023-07-01

**Description** Evidence of Absence software (EoA) is a user-friendly application for estimating bird and bat fatalities at wind farms and designing search protocols. The software is particularly useful in addressing whether the number of fatalities has exceeded a given threshold and what search parameters are needed to give assurance that thresholds were not exceeded. The models are applicable even when zero carcasses have been found in searches, following Huso et al. (2015) <[doi:10.1890/14-0764.1](https://doi.org/10.1890/14-0764.1)>, Dalthorp et al. (2017) <[doi:10.3133/ds1055](https://doi.org/10.3133/ds1055)>, and Dalthorp and Huso (2015) <[doi:10.3133/ofr20151227](https://doi.org/10.3133/ofr20151227)>.

**Depends** R (>= 3.5.0)

**License** GPL-2

**Encoding** UTF-8

**LazyData** true

**Imports** actuar, GenEst, MASS, rjags, survival, VGAM

**RoxygenNote** 7.2.3

**NeedsCompilation** no

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**Repository** CRAN

**Date/Publication** 2023-07-04 13:03:08 UTC

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cpdata0	<i>A template for carcass persistence data with interval-censored carcass persistence times</i>
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### Description

A template for carcass persistence data with interval-censored carcass persistence times

### Usage

cpdata0

### Format

A matrix with 2 columns bracketing persistence times (in days since carcass placement) of each carcass:

**CPmin** the last time the carcass was observed

**CPmax** the first time the carcass was noted missing

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days0	<i>A template for search schedule data</i>
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### Description

A template for search schedule data

### Usage

days0

### Format

A numeric vector with the times when searches were conducted, with days[1] = 0:

**days0** numeric vector of search times

**Description**

This package is designed to analyze searcher efficiency, carcass persistence, search schedule, and carcass observation data for the estimation of bird and bat mortality at wind and solar power facilities. It is specially designed for analyses when few carcasses are observed and detection probability is low. [It works fine for large counts as well, but some estimators (GenEst in particular) are more well-endowed with a wider array of tools for analysis of large-count data.

**Main command-line functions**

`est_pk0`, `est_cp0`, `est_g0` estimate searcher efficiency (pk), carcass persistence (cp), and (g) parameters

`postM`, `postM.ab` estimate posterior distribution of  $M$  given estimated  $g$  and carcass count ( $X$ )

`calcMstar`, `MCI` calculate  $M^*$  and credible interval for  $M$

**Potentially useful calculation functions**

`sim_pk0`, `sim_cp0` simulate estimated SE and CP parameters

`ppersist` calculate probability that a carcass that arrives at an (unknown) uniform random time in an interval persists until a later, specified time. This is the generalized  $r$  statistic for a given persistence distribution, arrival interval width, and search time.

**Behind-the-scenes utility functions**

`fmmax`, `fmmax.ab` functions to calculate a suitable maximum value to truncate improper prior distributions of  $M$

`getab` function to extract MLE of pda and pdb parameters from a persistence distribution

**Description**

Carcass persistence is modeled as survival function

**Usage**

`est_cp0(cpdata, dist)`

**Arguments**

cpdata	matrix of interval censored carcass persistence times
dist	Name of the persistence distribution family: Weibull, lognormal, loglogistic, or exponential

**Details**

uses survival package to fit.

**Value**

answer

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est_g0	<i>estimate g from fitted pk and cp models and search schedule</i>
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**Description**

Given a fitted pk model (from [est\\_pk0](#)), a fitted cp model (from [est\\_cp0](#) or a survreg object), and a search schedule, estimate detection probability.

**Usage**

```
est_g0(pkmodel, cpmodel, days, a = NULL, v = NULL, ...)
```

**Arguments**

pkmodel	fitted pk model (from <a href="#">est_pk0</a> )
cpmodel	fitted cp model (from <a href="#">est_cp0</a> or survreg object)
days	vector of days since searches begin (days[1] == 0)
a	fraction of carcasses arriving in the area searched
v	fraction of carcasses arriving in the period spanned by the monitoring
...	additional arguments (ignored)

**Value**

list of parameters for a beta distributions fit to the vectors of for  $\hat{g}$  for the searched area within the period monitored (`$BabRaw`), for the whole site within the period monitored (`$Bab`), and for the whole site extrapolated to the whole year (`$BabAnn`). In addition, the models and parameters that went into the estimate are included as well (`pkmodel`, `cpmodel`, `a`, `v`).

**Examples**

```
pkmodel <- est_pk0(pkdata = pkdata0)
cpmodel <- est_cp0(cpdata = cpdata0, dist = "weibull")
ghat <- est_g0(pkmodel = pkmodel, cpmodel = cpmodel, days = days0, a = 0.4, v = 0.75)
summary(ghat)
```

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est_pk0	<i>Fit pk searcher efficiency models</i>
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**Description**

Searcher efficiency is modeled as a function of the number of times a carcass has been missed in previous searches and any number of covariates.

**Usage**

```
est_pk0(pkdata, kFixed = NULL, n.iter = 1000, ...)
```

**Arguments**

pkdata	Search trial data entered in a list of N-vectors, \$n and \$y, indicating the number of carcasses available and the number discovered in searcher efficiency field trials in which carcasses were available for discovery. [NOTE: In earlier versions of eoa, the vectors were \$M and \$X. The names have been changed to avoid confusion with the M and X for total mortality and carcasses discovered carcass survey.]
kFixed	If trial carcasses are available for discovery for one search and data are insufficient for estimating k, a fixed, assumed value must be entered for k.
n.iter	number of iterations to use in updating the JAGS model for $p$ and $k$
...	Other parameters that may be used in called functions (esp. burn for updating the JAGS function)

**Details**

The probability of finding a carcass that is present at the time of search is  $p$  on the first search after carcass arrival and is assumed to decrease by a factor of  $k$  each time the carcass is missed in searches.

**Value**

A list with an nsim x 2 matrix of simulated  $p$  and  $k$  values the joint posterior for SE.

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fmmx	<i>Find suitable mmax for clipping improper priors for M</i>
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**Description**

Improper priors need to be clipped in order to be usable. fmmx and fmmx.ab find values of  $m$  that are large enough that the probability of exceeding is less than 0.0001 (depends on  $g$  and  $X$ ).

**Usage**

```
fmmax(x, g)
```

```
fmmax.ab(x, pBa, pBb)
```

**Arguments**

x                    carcass count

g                    overall carcass detection probability

pBa, pBb            parameters for beta distribution characterizing estimated  $g$

**Value**

integer  $m$  such that  $Pr(M \geq m) < 0.0001$

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getab	<i>retrieve EoA parameterization from survival parameterization of a fitted cp model (or survreg object with exponential, weibull, lognormal, or loglogistic distribution)</i>
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**Description**

retrieve EoA parameterization from survival parameterization of a fitted cp model (or survreg object with exponential, weibull, lognormal, or loglogistic distribution)

**Usage**

```
getab(cpmodel)
```

**Arguments**

cpmodel            fitted cp model (from est\_cp0 or survreg object)

**Value**

2-vector of pda and pdb

---

MpriorOK                      *Check validity of format of custom prior for M*

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**Description**

Check validity of format of custom prior for M

**Usage**

MpriorOK(prior)

**Arguments**

prior                      a custom prior for M must be a matrix with columns for M and associated probabilities  $P(M = m)$ . The M column must begin at 0 and the probabilities must sum to 1.

**Value**

boolean. Is the prior formatted properly?

---

pkdata0                      *A template for summarized searcher efficiency data with the number of carcasses available and the number discovered for N = 12 search occasions*

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**Description**

A template for summarized searcher efficiency data with the number of carcasses available and the number discovered for N = 12 search occasions

**Usage**

pkdata0

**Format**

A list with 2 numeric N-vectors with numbers of:

**n** searcher efficiency trial carcasses available

**y** carcasses discovered

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postM	<i>Calculate posterior distribution of M and extract statistics (M* and CI)</i>
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### Description

Calculation of the posterior distribution of total mortality ( $M$ ) given the carcass count, overall detection probability ( $g$ ), and prior distribution; calculation of summary statistics from the posterior distribution of  $M$ , including  $M^*$  and credibility intervals.

### Usage

```
postM(x, g, prior = "IbinRef", mmax = NA)
postM.ab(x, Ba, Bb, prior = "IbinRef", mmax = NULL)
calcMstar(pMgX, alpha)
MCI(pMgX, crlev = 0.95)
```

### Arguments

x	carcass count
g	overall carcass detection probability
prior	prior distribution of $M$
mmax	cutoff for prior of $M$ (large max requires large computing resources but does not help in the estimation)
Ba, Bb	parameters for beta distribution characterizing estimated $g$
pMgX	posterior distribution of $M$
crlev, alpha	credibility level ( $1 - \alpha$ ) and its complement ( $\alpha$ )

### Details

The functions `postM` and `postM.ab` return the posterior distributions of  $M|(X, g)$  and  $M|(X, Ba, Bb)$ , respectively, where  $Ba$  and  $Bb$  are beta distribution parameters for the estimated detection probability. `postM` and `postM.ab` include options to specify a prior distribution for  $M$  and a limit for truncating the prior to disregard implausibly large values of  $M$  and make the calculations tractable in certain cases where they otherwise might not be. Use `postM` when  $g$  is fixed and known; otherwise, use `postM.ab` when uncertainty in  $g$  is characterized in a beta distribution with parameters  $Ba$  and  $Bb$ . The non-informative, integrated reference prior for binomial random variables is the default (`prior = "IbinRef"`). Other options include "binRef", "IbetabinRef", and "betabinRef", which are the non-integrated and integrated forms of the binomial and betabinomial reference priors (Berger et al., 2012). For  $X > 2$ , the integrated and non-integrated reference priors give virtually identical posteriors. However, the non-integrated priors assign infinite weight to  $m = 0$  and return a posterior of  $Pr(M = 0|X = 0, \hat{g}) = 1$ , implying absolute certainty that the total number of fatalities



was 0 if no carcasses were observed. In addition, a uniform prior may be specified by `prior = "uniform"`. Alternatively, a custom prior may be given as a 2-dimensional array with columns for  $m$  and  $Pr(M = m)$ , respectively. The first column (`m`) must be sequential integers starting at  $m = 0$ . The second column gives the probabilities associated with  $m$ , which must be non-negative and sum to 1. The named priors ("`IbinRef`", "`binRef`", "`IbetabinRef`", and "`betabinRef`") are functions of  $m$  and defined on  $m = 0, 1, 2, \dots$  without upper bound. However, the posteriors can only be calculated for a finite number of  $m$ 's up to a maximum of `mmax`, which is set by default to the smallest value of  $m$  such that  $Pr(X \leq x|m, \hat{g}) < 0.0001$ , where  $x$  is the observed carcass count, or, alternatively, `mmax` may be specified by the user.

### Value

The functions `postM` and `postM.ab` return the posterior distributions of  $M|(X, g)$  and  $M|(X, Ba, Bb)$ , respectively. The functions `calcMstar` and `MCI` return  $M^*$  value and credibility interval for the given posterior distribution, `pMgX` (which may be the return value of `postM` or `postM.ab`) and  $\alpha$  value or credibility level.

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<code>sim_cp0</code>	<i>generate random cp parameters or persistence times</i>
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### Description

Given a fitted cp model (survreg object), generate random pda and pdb parameters or random persistence times. NOTE: This function is likely to move to call GenEst's `rcp` in future. This will not change the results, but the GenEst version is more nicely coded and keeping some coherence among the models is helpful.

### Usage

```
sim_cp0(cpmode1, nsim, option = "parms")
```

### Arguments

<code>cpmodel</code>	Fitted cp model ((survreg object))
<code>nsim</code>	Number of simulation draws
<code>option</code>	<code>option = "parms"</code> returns random draws of parameters from the fitted model; <code>option != "parms"</code> returns random draws of carcass persistence times

### Value

answer

---

 sim\_pk0

*Simulate pk parameters from model*


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**Description**

Simple simulation

**Usage**

```
sim_pk0(pkmodel, nsim = 1000)
```

**Arguments**

pkmodel	A model returned from est_pk0
nsim	Number of simulation reps for estimating the joint posterior distribution of p and k.

**Value**

An nsim x 2 matrix of simulated p and k values the joint posterior for SE.

---

summary.estg

*Summary statistics for estimated g*


---

**Description**

Summary statistics for estimated g

**Usage**

```
## S3 method for class 'estg'
summary(object, crlev = 0.95, ...)
```

**Arguments**

object	An <a href="#">estg</a> object
crlev	Credibility level of estimated CI to be returned
...	additional (optional) arguments passed to <code>rjags::coda.samples</code>

**Value**

summary statistics for estimated g. searched is for the fraction of carcasses falling in the searched area during the monitored period, site is area-adjusted to account for carcasses falling outside the searched area, and full is further extrapolated to the full year.

---

summary.estpk	<i>Summary statistics for estimated p and p parameters</i>
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---

**Description**

Summary statistics for estimated p and p parameters

**Usage**

```
## S3 method for class 'estpk'  
summary(object, crlev = 0.95, n.iter = 10000, ...)
```

**Arguments**

object	An <a href="#">estpk</a> object
crlev	Credibility level of estimated CI to be returned
n.iter	Number of iterations of the JAGS model for estimating the joint posterior distribution of p and k (relevant only if <code>object\$type == "pk"</code> ).
...	additional (optional) arguments passed to <code>rjags::coda.samples</code>

**Value**

array of summary statistics for p and k

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