# Package 'jointNmix'

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Type Package

Title Joint N-Mixture Models for Site-Associated Species

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**Depends** R (>= 3.0.0), methods, graphics, stats

#### Description

Fits univariate and joint N-mixture models for data on two unmarked site-associated species. Includes functions to estimate latent abundances through empirical Bayes methods.

License GPL (>= 2)

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jointNmix-package

# Description

Fits univariate and joint N-mixture models for data on two unmarked site-associated species. Includes functions to estimate latent abundances through empirical Bayes methods.

### Details

The DESCRIPTION file:

Package:	jointNmix
Type:	Package
Title:	Joint N-Mixture Models for Site-Associated Species
Version:	1.0
Date:	2016-10-04
Authors@R:	c(person("Rafael", "de Andrade Moral", role = c("aut", "cre"), email = "rafael_moral@yahoo.com.br"), person
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Maintainer:	Rafael de Andrade Moral <rafael_moral@yahoo.com.br></rafael_moral@yahoo.com.br>
Depends:	R ( $\geq$ 3.0.0), methods, graphics, stats
Description:	Fits univariate and joint N-mixture models for data on two unmarked site-associated species. Includes function
License:	GPL (>=2)

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jointNmix-package	Joint N-Mixture Models for Site-Associated
	Species
plot.uniNmix	Plot Methods for Nmix objects
	Species

# Author(s)

Rafael de Andrade Moral [aut, cre], Clarice Garcia Borges Demetrio [aut], John Hinde [aut] Maintainer: Rafael de Andrade Moral <rafael\_moral@yahoo.com.br>

#### abundance-prediction

#### References

Moral, R.A., Hinde, J., Demétrio, C.G.B., Reigada, C. and Godoy, W.A.C. (submitted) Models for jointly estimating abundance of two unmarked site-associated species subject to imperfect detection.

abundance-prediction Get empirical Bayes predictions of the latent abundances

#### Description

Computes the conditional abundance distribution given the data and model estimates

# Usage

```
getranef.uniNmix(obj, distr = FALSE)
getranef.jointNmix(obj, distr = FALSE)
```

#### Arguments

obj	fitted model object
distr	logical. If TRUE, returns the matrix of conditional probabilities. If FALSE, returns the expected values of the conditional distributions, i.e. the abundance estimates

#### Details

These functions return the empirical Bayes estimates of the latent abundances from univariate and joint N-mixture models, as described by Royle (2004) and Moral et al. (submitted), respectively.

#### Author(s)

Rafael A. Moral <rafael\_moral@yahoo.com.br>, Clarice G. B. Demétrio and John Hinde

#### References

Royle, J.A. (2004) Models for estimating population size from spatially replicated counts. Biometrics 60:108-105.

Moral, R.A., Hinde, J., Demétrio, C.G.B., Reigada, C. and Godoy, W.A.C. (submitted) Models for jointly estimating abundance of two unmarked site-associated species subject to imperfect detection.

#### See Also

Nmix jointNmix

corr-cov-jointNmix Extract correlations and covariances

#### Description

Extract estimated correlations and covariances between observed abundances for joint N-mixture models

#### Usage

```
getcorr.jointNmix(obj)
getcov.jointNmix(obj)
```

### Arguments

obj fitted model object

# Details

The correlations under each specification of the latent abundance distributions are displayed in Moral et al. (submitted)'s Table 1. The covariances are derived as Supplementary information.

# Author(s)

Rafael A. Moral <rafael\_moral@yahoo.com.br>, Clarice G. B. Demétrio and John Hinde

#### References

Moral, R.A., Hinde, J., Demétrio, C.G.B., Reigada, C. and Godoy, W.A.C. (submitted) Models for jointly estimating abundance of two unmarked site-associated species subject to imperfect detection.

### See Also

jointNmix

dneymanA

The Neyman-A probability function

#### Description

Computes the probability function of the Neyman-A distribution

#### Usage

```
dneymanA(x, lambda1, lambda2, K, log = FALSE)
```

# jointNmix

#### Arguments

х	vector of values
lambda1, lambda	a2
	parameters of the distribution
К	truncation value for the infinite summation
log	logical. If TRUE, the logarithm of the probabilities is returned

#### Details

The Neyman-A distribution has probability function

$$\frac{e^{-\lambda_1}\lambda_2^x}{x!}\sum_{k=0}^{\infty}\frac{(\lambda_1e^{-\lambda_2})^kk^x}{k!}$$

and is an overdispersion model. The summation is truncated to K.

# Author(s)

Rafael A. Moral <rafael\_moral@yahoo.com.br>, Clarice G. B. Demétrio and John Hinde

# Examples

```
x <- 0:10
dneymanA(x, lambda1 = 2, lambda2 = 1.5, K = 50)
```

jointNmix Joint N-mixture models

# Description

Fits joint N-mixture models for site-associated species

#### Usage

```
jointNmix(sp1, sp2, start, method = "BFGS", K, mixture = c("P", "P"),
Xp1, Xp2, Xl1, Xl2, Xpsi, includepsi = TRUE)
```

#### Arguments

sp1	observation matrix for species 1
sp2	observation matrix for species 2
start	initial values for the optimization process
method	optimization method passed to optim. Defaults to "BFGS"
К	truncation number of the infinite summations in the log-likelihood. Defaults to $max(sp1, sp2) + 100$

mixture	two-character vector for latent abundance distributions. "P" for Poisson and "NB" for negative binomial. Defaults to $c("P", "P")$
Xp1	model matrix for detection probabilities of species 1
Xp2	model matrix for detection probabilities of species 2
X11	model matrix for abundance of species 1
X12	model matrix for linking parameter of species 2
Xpsi	model matrix for abundance of species 2
includepsi	logical. If FALSE, psi is not estimated and set to zero

#### Details

The function fits a bivariate extension to Royle's (2004) N-mixture model to data on the abundance of two species collected at R sites over T time occasions. The model for observation on site i at time t for species 1 can be specified as

$$Y_{1it}|N_{1i} Bin(N_{1i}, p_{1it})$$

 $N_{1i}$  acount distribution with mean  $\lambda_{1i}$ .

The model for species 2 is

 $Y_{2it}|N_{1i}, N_{2i} Bin(N_{2i}, p_{2it})$ 

 $N_{2i}|N_{1i}$  acount distribution with mean  $\psi + \lambda_{2i}N_{1i}$ .

Here, users may define a Poisson or negative binomial distribution for the latent abundances N\_1i and N\_2i.

#### Value

An object of class jointNmix and Nmix, for which many methods are available (see methods(class = "jointNmix") and methods(class = "Nmix"))

## Author(s)

Rafael A. Moral <rafael\_moral@yahoo.com.br>, Clarice G. B. Demétrio and John Hinde

# References

Moral, R.A., Hinde, J., Demétrio, C.G.B., Reigada, C. and Godoy, W.A.C. (submitted) Models for jointly estimating abundance of two unmarked site-associated species subject to imperfect detection.

#### See Also

Nmix

#### jointNmix-methods

#### Examples

```
## simulating data with poisson latent abundances
R <- 10 # sites
T <- 10 # time occasions
lambda1 <- 5
psi <- 3
p1 <- .3
p2 <- .6
lambda2 <- .5
set.seed(1234); N1 <- rpois(R, lambda1)</pre>
set.seed(1234); N2 <- rpois(R, psi + lambda2*N1)</pre>
y1 <- y2 <- matrix(0, ncol=T, nrow=R)</pre>
set.seed(1234); for(i in 1:R) y1[,i] <- rbinom(T, N1, p1)</pre>
set.seed(1234); for(i in 1:R) y2[,i] <- rbinom(T, N2, p2)</pre>
Xp <- cbind(rep(1, R*T))</pre>
Xl <- cbind(rep(1, R))</pre>
## Not run:
## fitting the Poisson-Poisson joint N-mixture model
fitpp <- jointNmix(y1, y2, Xp1=Xp, Xp2=Xp, Xl1=X1, Xl2=Xl, mixture=c("P","P"), K=30)
## fitting the negbin-Poisson joint N-mixture model
fitnbp <- jointNmix(y1, y2, Xp1=Xp, Xp2=Xp, Xl1=Xl, Xl2=Xl, mixture=c("NB","P"), K=30)</pre>
## likelihood-ratio test between P-P and NB-P models
anova(fitpp, fitnbp)
## comparing using AIC
lapply(list(fitpp, fitnbp), AIC)
## conditional posterior probability functions for abundances
plot(fitpp, posterior = TRUE)
## estimated abundances vs. true abundances
data.frame(getranef.jointNmix(fitpp), N1, N2)
## End(Not run)
```

jointNmix-methods Methods for joint Nmix objects

#### Description

Fitted values, residual extraction, simulation and print method for joint N-mixture models

# Usage

## S3 method for class 'jointNmix'

```
fitted(object, ...)
## S3 method for class 'jointNmix'
print(x, round = TRUE, ...)
## S3 method for class 'jointNmix'
residuals(object, type = c("ordinary", "standardized"), ...)
## S3 method for class 'jointNmix'
simulate(object, ...)
```

#### Arguments

object, x	fitted model object
round	logical. Round the estimates?
type	type of residuals to be returned. May be ordinary or standardized
	not used

# Details

fitted.uniNmix prints the fitted values for the model fit; residuals.uniNmix returns the ordinary (

$$r_i^o = y_i - \hat{\mu}_i$$

) or standardized (

$$r_i^s = \frac{y_i - \hat{\mu}_i}{\operatorname{Var}(Y_i)}$$

) residuals; simulate.uniNmix simulates one sample from the fitted model.

# Author(s)

Rafael A. Moral <rafael\_moral@yahoo.com.br>, Clarice G. B. Demétrio and John Hinde

#### References

Moral, R.A., Hinde, J., Demétrio, C.G.B., Reigada, C. and Godoy, W.A.C. (submitted) Models for jointly estimating abundance of two unmarked site-associated species subject to imperfect detection.

# See Also

Nmix jointNmix

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# Description

Fits univariate N-mixture models

#### Usage

Nmix(sp1, start, method = "BFGS", K, mixture, Xp, Xl)

# Arguments

sp1	observation matrix for the species
start	initial values for the optimization process
method	optimization method passed to optim. Defaults to "BFGS"
К	truncation number of the infinite summations in the log-likelihood. Defaults to $max(sp1, sp2) + 100$
mixture	latent abundance distribution specification. "P" for Poisson, "NB" for negative binomial and "NeymanA" for Neyman-A distributions are available. Defaults to "P"
Хр	model matrix for detection probabilities
X1	model matrix for abundance parameter

# Details

The function fits Royle's (2004) N-mixture model to data on species abundance collected at R sites over T time occasions. The model for observation on site i at time t can be specified as

 $Y_{it}|N_i Bin(N_i, p_{it})$ 

# $N_i$ acount distribution with mean $\lambda_i$ .

Here, users may define a Poisson, negative binomial or Neyman-A distributions for the latent abundances N\_i.

# Value

An object of class uniNmix and Nmix, for which many methods are available (see methods(class = "uniNmix") and methods(class = "Nmix"))

#### Author(s)

Rafael A. Moral <rafael\_moral@yahoo.com.br>, Clarice G. B. Demétrio and John Hinde

#### References

Royle, J.A. (2004) Models for estimating population size from spatially replicated counts. Biometrics 60:108-105.

#### See Also

jointNmix

#### Examples

```
## simulating data with negative binomial latent abundances
R <- 10 # sites
T <- 10 # time occasions
lambda <- 5 # abundance parameter</pre>
p <- .3 # probability of detection</pre>
phi <- 1 # dispersion parameter
set.seed(1234); Ni <- rnbinom(R, mu=lambda, size=phi) # latent abundances</pre>
y <- matrix(0, ncol=T, nrow=R)</pre>
set.seed(1234); for(i in 1:R) y[,i] <- rbinom(T, Ni, p) # observed abundances</pre>
## fitting the Poisson N-mixture model
fitp <- Nmix(y, Xp=cbind(rep(1, R*T)), Xl=cbind(rep(1, R)), mixture="P", K=25)</pre>
## fitting the negative binomial N-mixture model
fitnb <- Nmix(y, Xp=cbind(rep(1, R*T)), Xl=cbind(rep(1, R)), mixture="NB", K=25)</pre>
## fitting the Neyman-A N-mixture model
fitna <- Nmix(y, Xp=cbind(rep(1, R*T)), X1=cbind(rep(1, R)), mixture="NeymanA", K=25)</pre>
## likelihood-ratio test between Poisson and negbin models
anova(fitp, fitnb)
## comparing using AIC
lapply(list(fitp, fitnb, fitna), AIC)
## conditional posterior probability functions for abundances
plot(fitnb, posterior = TRUE)
## estimated abundances vs. true abundances
data.frame(getranef.uniNmix(fitnb), Ni)
```

Nmix-methods

Methods for Nmix objects

#### Description

AIC, anova, coef, and logLik methods for univariate and joint N-mixture models

#### plot-methods-Nmix

# Usage

```
## S3 method for class 'Nmix'
AIC(object, ...)
## S3 method for class 'Nmix'
anova(object, object2, ...)
## S3 method for class 'Nmix'
coef(object, ...)
## S3 method for class 'Nmix'
logLik(object, ...)
```

#### Arguments

object	fitted model object
object2	second model object for the anova method
	not used

# Details

AIC.Nmix provides the Akaike Information Criterion for the model fit; anova.Nmix performs a likelihood-ratio test between two nested model fits; coef.Nmix prints the estimates of the model fit; logLik.Nmix prints the log-likelihood of the model fit.

#### Author(s)

Rafael A. Moral <rafael\_moral@yahoo.com.br>, Clarice G. B. Demétrio and John Hinde

# See Also

Nmix jointNmix

plot-methods-Nmix Plot Methods for Nmix objects

# Description

Plots residuals vs. fitted values or the conditional distributions of the abundances for each site

# Usage

```
## S3 method for class 'uniNmix'
plot(x, posterior = FALSE, layout, sites, restype, ...)
## S3 method for class 'jointNmix'
plot(x, posterior = FALSE, layout, sites, restype, ...)
```

#### Arguments

х	fitted model object
posterior	logical. If TRUE, plots the conditional distributions of the abundances. If FALSE, plots residuals vs. fitted values
layout	two-number vector indicating the partition of the graphical window in rows and columns, respectively
sites	index of sites to plot the conditional distribution of the abundances. If left un- specified, all sites are plotted
restype	type of residuals to be plotted. Defaults to "ordinary"
	not used

# Author(s)

Rafael A. Moral <rafael\_moral@yahoo.com.br>, Clarice G. B. Demétrio and John Hinde

# References

Moral, R.A., Hinde, J., Demétrio, C.G.B., Reigada, C. and Godoy, W.A.C. (submitted) Models for jointly estimating abundance of two unmarked site-associated species subject to imperfect detection.

#### See Also

Nmix jointNmix

uniNmix-methods Methods for univariate Nmix objects

# Description

Fitted values, residual extraction, simulation and print method for univariate N-mixture models

#### Usage

```
## S3 method for class 'uniNmix'
fitted(object, ...)
## S3 method for class 'uniNmix'
print(x, round = TRUE, ...)
## S3 method for class 'uniNmix'
residuals(object, type = c("ordinary", "standardized"), ...)
## S3 method for class 'uniNmix'
simulate(object, ...)
```

# Arguments

object, x	fitted model object
round	logical. Round the estimates?
type	type of residuals to be returned. May be ordinary or standardized
	not used

# Details

fitted.uniNmix prints the fitted values for the model fit; residuals.uniNmix returns the ordinary (  $r_i^o=y_i-\hat{\mu}_i$ 

) or standardized (

$$r_i^s = \frac{y_i - \hat{\mu}_i}{\operatorname{Var}(Y_i)}$$

) residuals; simulate.uniNmix simulates one sample from the fitted model.

# Author(s)

Rafael A. Moral <rafael\_moral@yahoo.com.br>, Clarice G. B. Demétrio and John Hinde

# See Also

Nmix jointNmix

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