Package ‘RSE’

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Author Youhua Chen, Tsung-Jen Shen
Maintainer Youhua Chen <haydi@QR6N.com>
Description A Bayesian-weighted estimator and two unweighted estimators are developed to estimate the number of newly found rare species in additional ecological samples. Among these methods, the Bayesian-weighted estimator and an unweighted (Chao-derived) estimator are of high accuracy and recommended for practical applications. Technical details of the proposed estimators have been well described in the following paper: Shen TJ, Chen YH (2018) A Bayesian weighted approach to predicting the number of newly discovered rare species. Conservation Biology, In press.
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RSE-package Number of Newly Discovered Rare Species Estimation

Description

RSE: an R package for predicting numbers of rare species newly discovered in additional ecological samples

Details

RSE is an R package for estimating the number of new rare species in additional ecological samples. The methods used here include a Bayesian-weighted estimator and two unweighted estimators.

Author(s)

Youhua Chen (Chengdu Institute of Biology, Chinese Academy of Sciences);
Tsung-Jen Shen (Institute of Statistics & Department of Applied Mathematics, National Chung Hsing University)
Maintainer:
Youhua Chen <haydi@126.com>

References


Examples

# for abundance-based data type
data(HerpetologicalData)
## two columns represent two samples of species abundance data
X.merge = HerpetologicalData
## the first column is treated as the original sample
X.col1 = X.merge[,1]
## the second column is treated as the additional sample
X.col2 = X.merge[,2]
boot.abundance.fun

Generate a bootstrap abundance-based sample

Description

Given an abundance-based data, a bootstrap sample is generated from a reconstructed bootstrap assemblage.

Usage

boot.abundance.fun(S.hat, f, b)

Arguments

S.hat
An estimate of species richness.

f
A vector of species frequency counts, i.e., the number of singleton species (only one individual observed in the sample), the number of doubleton species (two individuals observed in the sample), and so forth.

b
A vector of estimates of two parameters for obtaining the estimated relative abundances of observed species in a given sample by Chao et al.’s (2015) method.

Value

The generated bootstrap sample is a vector of species frequency counts, i.e., the number of singleton species (only one individual observed in the bootstrap sample), the number of doubleton species (two individuals observed in the bootstrap sample), and so forth.

Author(s)

Youhua Chen & Tsung-Jen Shen
References


See Also

boot.incidence.fun

Examples

```r
## As an example, Herpetological assemblage data are used here.
data(HerpetologicalData)
## two columns represent two samples of species abundance data
X.merge = HerpetologicalData
## the first column is treated as the original sample
X1 = X.merge[,1]
## Convert species abundance data to species frequency counts data
f = X.to.f(X1)
b = DetAbu(x=X1, zero=FALSE)
## the estimated number of species
S.hat = SpEst.Chao1.abun(f)
boot.abundance.fun(S.hat=S.hat, f=f, b=b)
```

---

**Description**

Given an incidence-based data, a bootstrap sample is generated from a reconstructed bootstrap assemblage.

**Usage**

```r
boot.incidence.fun(S.hat, nT, Q, b)
```

**Arguments**

- `S.hat`: An estimate of species richness.
- `nT`: The number of quadrats of the original sample.
- `Q`: A vector of species frequency counts, i.e., the number of species detected once (in only one quadrat), the number of species detected twice (in exactly two quadrats), and so forth.
- `b`: A vector of estimates of two parameters for obtaining the estimated detection probabilities of observed species in a given sample by Chao et al.'s (2015) method.
**Value**

The generated bootstrap sample is a vector of species frequency counts, i.e., the number of species detected once (in only one quadrat of the bootstrap sample), the number of species detected twice (in exactly two quadrats of the bootstrap sample), and so forth.

**Author(s)**

Youhua Chen & Tsung-Jen Shen

**References**


**See Also**

`boot.abundance.fun`

**Examples**

```r
## As an example, Canadian-mite data are used here.
data(CanadaMite)
## two columns represent two samples of incidence counts
X.merge = CanadaMite
## the first column is treated as the sample
X.col1 = X.merge[,1]
X1 = X.col1
## Convert species incidence count data to frequency counts data
data = X.to.f(X1)
## the number of quadrats in the first sample
nT = 16
b = DetInc(X1, nT)
boot.incidence.fun(S.hat=sum(Q)+b[3], nT=nT, Q=Q, b=b[1:2])
```

**Description**

The mite data were collected by Youhua Chen (Chen et al. 2015) in western coast of Canada. In Chen et al. (2015), 16 moss sampling locations were surveyed from the early days to the midst of June 2011, while another 16 moss sampling units were surveyed from the midst of June to the early days of July 2011.

Therefore, for the dataset, it has two columns, the first column contained the incidence or occurrence information of mites collected by first days of sampling (the early days to the midst of June 2011), while the second column contained the incidence of mites that were collected by the last days of the sampling
DetAbu

Usage

data(CanadaMite)

Author(s)

Youhua Chen & Tsung-Jen Shen

References


See Also

HerpetologicalData

Examples

data(CanadaMite)
## two columns represent two samples of species incidence data
X.merge = CanadaMite
head(X.merge)

```
DetAbu(x, zero = FALSE)
```

Arguments

- **x**
  
  A vector of species abundance data, i.e., the number of individuals of species 1, the number of individuals of species 2, and so forth.

- **zero**
  
  A logical value for whether reserving zero frequency or not.
**Value**

A vector of 3 elements: the first two values are the estimates of two parameters in Chao et al. (2015) for jointly estimating relative abundances of observed species and the third one is the estimated number of unseen species in the sample by Chao 1 estimator (Chao, 1984).

**Note**

This function is a part of the original R code JADE by Chao et al. (2015) and is slightly modified for the output format.

**Author(s)**

Youhua Chen & Tsung-Jen Shen

**References**


R code for JADE: http://chao.stat.nthu.edu.tw/wordpress/paper/107_Rcode.txt

**See Also**

DetInc

**Examples**

```r
## As an example, Herpetological assemblage data are used here.
data(HerpetologicalData)
## two columns represent two samples of species abundance data
X.merge = HerpetologicalData
## the first column is treated as the original sample
X1 = X.merge[,1]
DetAbu(x=X1)
```

```r
detinc

Incidence-based data: the estimation of parameters for obtaining the estimation of detection probabilites of observed species

**Description**

The estimation of parameters for obtaining the estimation of detection probabilites of observed species

**Usage**

```r
DetInc(y, nT, zero = FALSE)
```
Arguments

- **y**: A vector of species incidence counts, i.e., the number of quadrats with species 1, the number of quadrats with species 2, and so forth.
- **nt**: The number of quadrats of the original sample.
- **zero**: A logical value for whether reserving zero frequency or not.

Value

A vector of 3 elements: the first two values are the estimates of two parameters in Chao et al. (2015) for jointly estimating detection probabilities of observed species and the third one is the estimated number of unseen species in the sample by Chao 2 estimator (Chao, 1987).

Note

This function is a part of the original R code JADE by Chao et al. (2015) and is slightly modified for the output format.

Author(s)

Youhua Chen & Tsung-Jen Shen

References


See Also

- **DetAbu**

Examples

```r
## As an example, Canadian-mite data are used here.
data(CanadaMite)
## two columns represent two samples of incidence counts
X.merge = CanadaMite
## the first column is treated as the original sample
X.col1 = X.merge[,1]
Xi = X.col1
## the number of quadrats in the first sample
nt = 16
DetInc(y = Xi, nT = nt)
```
**Description**

This function is to convert species frequency counts data to species abundance data.

**Usage**

\[ f \to X(f) \]

**Arguments**

- \( f \) Species frequency counts data.

**Value**

Species abundance data is returned.

**Author(s)**

Youhua Chen & Tsung-Jen Shen

**See Also**

\( X \to f \)

**Examples**

```r
## The sample is composed of 5 singletons, two doubletons, and one tripleton species.
f = c(5, 2, 1)
f.to.X(f)
```

---

**HerpetologicalData**

*Abundance of herpetofauna in the conserved and human disturbed areas of Mexico (Suazo-Ortuno et al. 2008)*

---

**Description**

Suazo-Ortuno et al. (2008) studied how the conversion of tropical forest to agricultural mosaic influenced herpetofaunal distribution and community structure in conserved and human disturbed forest areas of neotropical Mexico.

Therefore, the dataset used here, the first and second columns represented species abundance in the conserved and disturbed areas, respectively.
Usage

data(HerpetologicalData)

Author(s)

Youhua Chen & Tsung-Jen Shen

References


See Also

CanadaMite

Examples

data(HerpetologicalData)
## two columns represent two samples of species abundance data
X.merge = HerpetologicalData
head(X.merge)

Pred.abundance.rare  Abundance-based data: predicting the number of new rare species

Description

Abundance-based prediction on the number of new rare species using a Bayesian-weight and two unweighted estimators along with their bootstrap standard errors and 95% bootstrap confidence intervals.

Usage

Pred.abundance.rare(boot.rep = 100, f = NULL, xi = NULL, m, k.show = 3)
Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>boot.rep</td>
<td>Replicate number of the bootstrapping procedure</td>
</tr>
<tr>
<td>f</td>
<td>A vector of species frequency counts, i.e., the number of singleton species (only one individual observed in the sample), the number of doubleton species (two individuals observed in the sample), and so forth.</td>
</tr>
<tr>
<td>xi</td>
<td>A vector of species abundance data, i.e., the number of individuals of species 1, the number of individuals of species 2, and so forth.</td>
</tr>
<tr>
<td>m</td>
<td>The number of individuals of an additional sample</td>
</tr>
<tr>
<td>k.show</td>
<td>Display the estimating result of the numbers of extremely rare species with abundance ( \leq k.\text{show} ) in the additional sample</td>
</tr>
</tbody>
</table>

Value

Estimating results including point estimate, bootstrap standard error, and 95 % bootstrap confidence interval for each of three methods (a Bayesian-weight and two unweighted estimators)

Author(s)

Youhua Chen & Tsung-Jen Shen

References


See Also

Pred.incidence.rare

Examples

```r
## As an example, Herpetological assemblage data are used here.
data(HerpetologicalData)
## two columns represent two samples of species abundance data
X.merge = HerpetologicalData
## the first column is treated as the original sample
X.col1 = X.merge[,1]
## the second column is treated as the additional sample
X.col2 = X.merge[,2]
x1 = X.col1
## Convert species abundance data to species frequency counts data
f = x.to.f(x1)
## the number of individuals of the additional sample
m = sum(X.col2)
Pred.abundance.rare(f = f, m = m)
```
Description

Bayesian-weight estimator for predicting the number of new rare species using abundance data as inputs

Usage

Pred.Fk.BW(f, m, b, k.show = 3)

Arguments

f  A vector of species frequency counts, i.e., the number of singleton species (only one individual observed in the sample), the number of doubleton species (two individuals observed in the sample), and so forth.

m  The number of individuals of an additional sample

b  A vector of two estimated parameters for obtaining the estimated relative species abundances by Chao et al.'s (2015) method.

k.show  Display the estimating result of the numbers of extremely rare species with abundance <= k.show in the additional sample

Value

The numbers of new rare species with abundance <= k.show are estimated by the abundance-based Bayesian-weight estimator and returned.

Author(s)

Youhua Chen & Tsung-Jen Shen

References


See Also

Pred.Qk.BW
Examples

```r
## As an example, Herpetological assemblage data are used here.
data(HerpetologicalData)
## two columns represent two samples of species abundance data
X.merge = HerpetologicalData
## the first column is treated as the original sample
X.col1 = X.merge[,1]
## the second column is treated as the additional sample
X.col2 = X.merge[,2]
Xi = X.col1
## Convert species abundance data to species frequency counts data
f = X.to.f(Xi)
## the number of individuals of the additional sample
m = sum(X.col2)
b = DetAbu(x=Xi, zero=FALSE)
Pred.Fk.BW(f=f, m=m, b=b)
```

Description

Abundance-based unweighted naive estimator for predicting the number of new rare species

Usage

```
Pred.Fk.Naive(f, m, k.show = 3)
```

Arguments

- `f`: A vector of species frequency counts, i.e., the number of singleton species (only one individual observed in the sample), the number of doubleton species (two individuals observed in the sample), and so forth.
- `m`: The number of individuals of an additional sample
- `k.show`: Display the estimating result of the numbers of extremely rare species with abundance <= k.show in the additional sample.

Value

The numbers of new rare species with abundance <= k.show are estimated by the abundance-based unweighted naive estimator and returned.

Author(s)

Youhua Chen & Tsung-Jen Shen
References


See Also

Pred.Qk.Naive

Examples

```r
## As an example, Herpetological assemblage data are used here.
data(HerpetologicalData)
## two columns represent two samples of species abundance data
X.merge = HerpetologicalData
## the first column is treated as the original sample
X.col1 = X.merge[,1]
## the second column is treated as the additional sample
X.col2 = X.merge[,2]
X1 = X.col1
## Convert species abundance data to species frequency counts data
f = X.to.f(X1)
## the number of individuals of the additional sample
m = sum(X.col2)
Pred.Fk.Naive(f=f, m=m)
```

---

**Pred.Fk.unweighted**  
*Abundance-based data: Unweighted estimator*

**Description**

Unweighted estimator based on Chao et al. (2015)'s paper using abundance-based data for predicting the number of new rare species in an additional ecological sample

**Usage**

`Pred.Fk.unweighted(f, m, b, f0, k.show = 3)`

**Arguments**

- **f**: A vector of species frequency counts, i.e., the number of singleton species (only one individual observed in the sample), the number of doubleton species (two individuals observed in the sample), and so forth.
- **m**: The number of individuals of an additional sample
- **b**: A vector of two estimated parameters for obtaining the estimated relative species abundances by Chao et al.'s (2015) method.
- **f0**: The estimated number of unseen species in the original sample by Chao 1 estimator (Chao 1984)
- **k.show**: Display the estimating result of the numbers of extremely rare species with abundance \( \leq k.\text{show} \) in the additional sample
Value

The numbers of new rare species with abundance <= k.show are estimated by the abundance-based unweighted estimator and returned.

Author(s)

Youhua Chen & Tsung-Jen Shen

References


See Also

Pred.Qk.unweighted

Examples

### As an example, Herpetological assemblage data are used here.
```r
data(HerpetologicalData)
## two columns represent two samples of species abundance data
X.merge = HerpetologicalData
## the first column is treated as the original sample
X.col1 = X.merge[,1]
## the second column is treated as the additional sample
X.col2 = X.merge[,2]
X1 = X.col1
## Convert species abundance data to species frequency counts data
f = X.to.f(X1)
## the number of individuals of the additional sample
m = sum(X.col2)
## the estimated number of unseen species in the original sample
b = DetAbu(x=X1, zero=FALSE)
## the estimated number of newly discovered rare species
f0 = SpEst.Chao1.abun(f)-sum(f)
Pred.Fk.unweighted(f=f, m=m, b=b, f0=f0)
```

Pred.incidence.rare  Incidence-based data: predicting the number of new rare species

Description

Incidence-based prediction on the number of new rare species using a Bayesian-weight and two unweighted estimators along with their bootstrap standard errors and 95% bootstrap confidence intervals.
Usage

Pred.incidence.rare(boot.rep = 100, Q = NULL, xi = NULL, nT, u, k.show = 3)

Arguments

boot.rep        Replicate number of the bootstrapping procedure
Q               A vector of species frequency counts, i.e., the number of species detected once (in only one quadrat), the number of species detected twice (in exactly two quadrats), and so forth.
xi              A vector of species incidence counts, i.e., the number of quadrats with species 1, the number of quadrats with species 2, and so forth.
nT              The number of quadrats of the original sample
u               The number of quadrats of an additional sample
k.show          Display the estimating results of the numbers of new rare species detected in the number of quadrats <= k.show in the additional sample

Value

Estimating results including point estimate, bootstrap standard error, and 95 % bootstrap confidence interval for each of three methods (a Bayesian-weight and two unweighted estimators)

Author(s)

Youhua Chen & Tsung-Jen Shen

References


See Also

Pred.abundance.rare

Examples

```r
## As an example, Canadian-mite data are used here.
data(CanadaMite)
## two columns represent two samples of incidence counts
X.merge = CanadaMite
## the first column is treated as the original sample
X.col1 = X.merge[,1]
Xi = X.col1
## Convert species incidence count data to frequency counts data
Q = X.to.f(Xi)
## the number of quadrats in the first sample
nT = 16
## the number of quadrats in the additional sample (i.e., the second column)
u = 16
Pred.incidence.rare(Q=Q, nT=nT, u=u)
```
Description

Bayesian-weight estimator for predicting the number of new rare species using incidence/quadrat data

Usage

`Pred.Qk.BW(Q, nT, u, b, k.show = 3)`

Arguments

- `Q`: A vector of species frequency counts, i.e., the number of species detected once (in only one quadrat), the number of species detected twice (in exactly two quadrats), and so forth.
- `nT`: The number of quadrats of the original sample
- `u`: The number of quadrats of an additional sample
- `b`: A vector of two estimated parameters for obtaining the estimated relative species abundances by Chao et al.'s (2015) method.
- `k.show`: Display the estimating results of the numbers of new rare species detected in the number of quadrats <= k.show in the additional sample

Value

The numbers of new rare species detected in the number of quadrats <= k.show are estimated by the incidence-based Bayesian-weight estimator and returned.

Author(s)

Youhua Chen & Tsung-Jen Shen

References


See Also

`Pred.Fk.BW`
Examples

```r
## As an example, Canadian-mite data are used here.
data(CanadaMite)
## two columns represent two samples of incidence counts
X.merge = CanadaMite
## the first column is treated as the original sample
X.col1 = X.merge[,1]
X1 = X.col1
## Convert species incidence count data to frequency counts data
Q = X.to.f(X1)
## the number of quadrats in the first sample
nT = 16
## the number of quadrats in the additional sample (i.e., the second column)
 u = 16
 b = DetInc(y=X1, nT=nT)
 Pred.Qk.BW(Q=Q, nT=nT, u=u, b=b[1:2])
```

## Pred.Qk.Naive

**Incidence-based data: unweighted naive estimator**

**Description**

Incidence-based unweighted naive estimator for predicting the number of new rare species

**Usage**

```r
Pred.Qk.Naive(nT, u, f, k.show = 3)
```

**Arguments**

- `nT` The number of quadrats of the original sample
- `u` The number of quadrats of an additional sample
- `f` A vector of species frequency counts, i.e., the number of species detected once (in only one quadrat), the number of species detected twice (in exactly two quadrats), and so forth.
- `k.show` Display the estimating results of the numbers of new rare species detected in the number of quadrats <= k.show in the additional sample.

**Value**

The numbers of new rare species detected in the number of quadrats <= k.show are estimated by the incidence-based unweighted naive estimator and returned.

**Author(s)**

Youhua Chen & Tsung-Jen Shen
References


See Also

Pred.Fk.Naive

Examples

```r
## As an example, Canadian-mote data are used here.
data(CanadaMite)
## two columns represent two samples of incidence counts
X.merge = CanadaMite
## the first column is treated as the original sample
X.col1 = X.merge[,1]
X1 = X.col1
## Convert species incidence count data to frequency counts data
Q = X.to.f(X1)
## the number of quadrats in the first sample
nT = 16
## the number of quadrats in the additional sample (i.e., the second column)
u = 16
Pred.Qk.Naive(nT=nT, u=u, f=Q)
```

Description

Unweighted Estimator derived from Chao et al. (2015)’s paper using incidence/quadrat data for predicting the number of new rare species in an additional ecological sample

Usage

```r
Pred.Qk.unweighted(Q, nT, u, b, Q0, k.show = 3)
```

Arguments

- `Q` A vector of species frequency counts, i.e., the number of species detected once (in only one quadrat), the number of species detected twice (in exactly two quadrats), and so forth.
- `nT` The number of quadrats of the original sample
- `u` The number of quadrats of an additional sample
- `b` A vector of two estimated parameters for obtaining the estimated relative species abundances by Chao et al.’s (2015) method.
The estimated number of unseen species in the original sample by Chao 2 estimator (Chao 1987)

Display the estimating results of the numbers of new rare species detected in the number of quadrats <= k.show in the additional sample

The numbers of new rare species detected in the number of quadrats <= k.show are estimated by the incidence-based unweighted estimator derived from Chao et al. (2015)’s paper and returned.

Youhua Chen & Tsung-Jen Shen


See Also

Pred.Qk.unweighted

Examples

```r
## As an example, Canadian-mite data are used here.
data(CanadaMite)
## two columns represent two samples of incidence counts
X.merge = CanadaMite
## the first column is treated as the original sample
X.col1 = X.merge[,1]
Xi = X.col1
## Convert species incidence count data to frequency counts data
Q = X.to.f(Xi)
## the number of quadrats in the first sample
nT = 16
## the number of quadrats in the additional sample (i.e., the second column)
u = 16
b = DetInc(Xi, nT)
Pred.Qk.unweighted(Q=Q, nT=nT, u=u, b=b[1:2], Q0=b[3])
```
**Description**

Chao1 estimator of species richness

**Usage**

```r
SpEst.Chao1.abun(f)
```

**Arguments**

- `f` A vector of species frequency counts, i.e., the number of singleton species (only one individual observed in the sample), the number of doubleton species (two individuals observed in the sample), and so forth.

**Author(s)**

Youhua Chen & Tsung-Jen Shen

**References**


**Examples**

```r
## As an example, Herpetological assemblage data are used here.
data(HerpetologicalData)
## two columns represent two samples of species abundance data
X.merge = HerpetologicalData
## the first column is treated as the original sample
X1 = X.merge[,1]
## Convert species abundance data to species frequency counts data
f = X.to.f(X1)
SpEst.Chao1.abun(f=f)
```

---

**X.to.f**

Data transformation: from species abundance data to species frequency counts data

**Description**

This function is to convert a vector of species abundance data to a vector of species frequency counts data.
Usage

X.to.f(X)

Arguments

X A vector of species abundance data.

Value

Species frequency counts is returned.

Author(s)

Youhua Chen & Tsung-Jen Shen

See Also

f.to.X

Examples

## As an example, Herpetological assemblage data are used here.
data(HerpetologicalData)
## two columns represent two samples of species abundance data
X.merge = HerpetologicalData
## the first column is treated as the original sample
X.col1 = X.merge[,1]
Xi = X.col1
## convert species abundance data to species frequency counts data
X.to.f(Xi)
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