Package ‘isoWater’

July 21, 2021

Type   Package
Title   Discovery, Retrieval, and Analysis of Water Isotope Data
Version 1.0.1
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Imports R2jags, abind, R2WinBUGS, doParallel, foreach, httr, jsonlite
Depends R (>= 3.5)
Suggests knitr, rmarkdown, testthat (>= 3.0.0), covr, curl
VignetteBuilder knitr
License GPL-3
Language en-US
Encoding UTF-8
RoxygenNote 7.1.1
NeedsCompilation no
Author Gabe Bowen [aut, cre]
Repository CRAN
Date/Publication 2021-07-20 23:40:02 UTC

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iso

Iso Object

Description

Creates objects of type "iso"

Usage

iso(H, O, Hsd, Osd, HOc = 0)

Arguments

H numeric. Hydrogen isotope value or vector of hydrogen isotope values.
O numeric. Oxygen isotope value or vector of oxygen isotope values.
Hsd numeric. 1 standard deviation uncertainty of H (value or vector of values).
Osd numeric. 1 standard deviation uncertainty of O (value or vector of values).
HOc numeric. Covariance of H and O uncertainties.

Value

Returns an object of class “iso”, a data.frame containing the provided values.

Examples

obs = iso(-30, -5, 2, 0.2, 0.3)
str(obs)
isoWater

Discovery, Retrieval, and Analysis of Water Isotope Data

Description


Author(s)

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mixSource

Water Source as a Mixture

Description

Given isotopic compositions of two or more potential sources, generate a posterior sample of source mixtures conditioned on one or more sample values.

Usage

```r
mixSource(obs, sources, slope, prior = rep(1,nrow(sources)), shp = 1, ngens = 1e5, ncores = 1)
```

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>obs</td>
<td><em>iso</em> object containing isotope values for one or more samples.</td>
</tr>
<tr>
<td>sources</td>
<td><em>iso</em> object containing isotope values for two or more sources which may have contributed to the observed samples.</td>
</tr>
<tr>
<td>slope</td>
<td>numeric. Vector of length two specifying prior parameters for the evaporation line slope (mean, standard deviation).</td>
</tr>
<tr>
<td>prior</td>
<td>numeric. Vector of length equal to the number of sources, giving prior estimates of relative contributions of different sources.</td>
</tr>
<tr>
<td>shp</td>
<td>numeric. Shape parameter constant used in specifying prior estimates of source contributions (see Details).</td>
</tr>
<tr>
<td>ngens</td>
<td>integer. Number of posterior samples to obtain (per chain).</td>
</tr>
<tr>
<td>ncores</td>
<td>integer. Number of cores to use for parallel processing.</td>
</tr>
</tbody>
</table>
Details

The Dirichlet distribution is used to represent the fractional contribution of each source. The prior estimate is a Dirichlet where the shape parameter for source \( i \) is given by \( \text{prior}[i] / \min(\text{prior}) \times \text{shp} \).

If \( n\text{cores} = 1 \), three chains will be run on a single core. If \( n\text{cores} > 1 \), \( n\text{cores} \) chains will be run in parallel on \( n\text{cores} \) cores.

Value

Returns an object of class “mixSource”, a list containing:

- **summary** matrix. Summary table of JAGS MCMC results, including parameter posterior distributions and convergence statistics.
- **results** data.frame. Posterior samples of model parameters.
  - **mixture_d2H** Hydrogen isotopic composition of unevaporated source mixture.
  - **mixture_d18O** Oxygen isotopic composition of unevaporated source mixture.
  - **sX_fraction** Fractional contribution of each source.
  - \( S \) Evaporation line slope.
  - \( E \) Evaporation index, equal to the difference between the sample and unevaporated source mixture oxygen isotope values.

Examples

```r
# Prep sources
O = runif(3, -15, -2)
H = O * 8 + 10 + rnorm(3, 0, 6)
sources = iso(H, O, 1, 0.2, 0.17)

# Sample data
obs = iso(-60, -6, 0.5, 0.1, 0)

# Evaporation slope
slope = c(5, 0.3)

# Run and report...likely not converged!
ws = mixSource(obs, sources, slope, ngens = 1e3)
ws$summary

# A traceplot
plot(ws$results$mixture_d18O[1:1000], type = "l")
lines(ws$results$mixture_d18O[1001:2000], col = 2)
lines(ws$results$mixture_d18O[2001:3000], col = 3)
```
Description

Creates objects of type "mwl" containing statistics for a meteoric water line in H and O isotope space.

Usage

mwl(HO, plot = TRUE)

Arguments

HO       data.frame. Hydrogen (column 1) and oxygen (column 2) isotope values for 3 or more water samples.
plot     logical. Plot the data, MWL, and standard error of prediction?

Details

mwl will return an error if fewer than 3 sample values are provided and a warning if fewer than 10 samples are provided or if the correlation coefficient between H and O values is less than 0.7. Sample values should span a broad enough range of isotope values to strongly constrain the MWL.

Model II (reduced major axis) regression is used to accommodate errors on both isotope values.

Value

Returns an object of class “mwl”, a numeric vector containing meteoric water line statistics. See mwlSource.

Examples

O = runif(10, -15, -2)
H = 0 * 8 + 10 + rnorm(10, 0, 6)
MWL = mwl(data.frame(H, O))
str(MWL)
mwlSource

Water Source Using Meteoric Water Line

Description

Given parameters describing a meteoric water line in H-O isotope space, generate a posterior sample of unevaporated source water values conditioned on one or more sample values.

Usage

mwlSource(obs, MWL=c(8.01, 9.57, -8.096, 2564532.2, 5.76, 80672), slope, stype = 1, ngens=1e4, ncores = 1)

Arguments

- obs: iso object containing isotope values for one or more samples.
- MWL: numeric. Vector of length 6 containing parameters describing a meteoric water line (see Details).
- slope: numeric. Vector of length two specifying prior parameters for the evaporation line slope (mean, standard deviation).
- stype: integer. Line statistic used to constrain the source prior: 1 = confidence interval, 2 = prediction interval (see Details).
- ngens: integer. Number of posterior samples to obtain (per chain).
- ncores: integer. Number of cores to use for parallel processing.

Details

The prior distribution of source values is constrained by MWL, which contains the parameters: slope, intercept, average d18O, sum of squares in d18O, root mean square error, and number of samples for an emperically-determined meteoric water line. This object can be created from a H and O isotope dataset using the function mwl. The default values reflect the Global Meteoric Water Line estimated from a global precipitation compilation in Bowen, et al. (2019). stype determines how the source uncertainty about the MWL is calculated; the default (1, confidence interval) is appropriate if the source is best represented as an integrated mixture of the samples defining the MWL, whereas option 2 (prediction interval) is appropriate if the source is best represented as a single sample.

If ncores = 1, three chains will be run on a single core. If ncores > 1, ncores chains will be run in parallel on ncores cores.

Value

Returns an object of class “mwlSource”, a list containing:

- summary: matrix. Summary table of JAGS MCMC results, including parameter posterior distributions and convergence statistics.
- results: data.frame. Posterior samples of model parameters.
**source_d2H**  Hydrogen isotopic composition of unevaporated source.

**source_d18O**  Oxygen isotopic composition of unevaporated source.

**S**  Evaporation line slope.

**E**  Evaporation index, equal to the difference between the sample and unevaporated source mixture oxygen isotope values.

### References


### Examples

```r
# Prep MWL
O = runif(10, -15, -2)
H = O * 8 + 10 + rnorm(10, 0, 6)
MWL = mwl(data.frame(H, O), plot = FALSE)

# Sample data
obs = iso(-60, -6, 0.5, 0.1, 0)

# Evaporation slope
slope = c(5, 0.3)

# Run and report...likely not converged!
ws = mwlSource(obs, MWL, slope, ngens = 1e3)
ws$summary

# A traceplot
plot(ws$results$source_d18O[1:1000], type = "l")
lines(ws$results$source_d18O[1001:2000], col = 2)
lines(ws$results$source_d18O[2001:3000], col = 3)
```

### wiDB_data

**Obtain data from wiDB using a query**

#### Description

Obtain data from wiDB using a query

#### Usage

```r
wiDB_data(minLat = NULL, maxLat = NULL, minLong = NULL, maxLong = NULL, minElev = NULL, maxElev = NULL, minDate = NULL, maxDate = NULL, countries = NULL, states = NULL, types = NULL, projects = NULL, fields = NULL, tmpdir = tempdir(), clean = TRUE)
```
Arguments

- **minLat** numeric. Minimum latitude for query region, in decimal degrees. South negative.
- **maxLat** numeric. Maximum latitude for query region, in decimal degrees. South negative.
- **minLong** numeric. Minimum longitude for query region, in decimal degrees. West negative.
- **maxLong** numeric. Maximum longitude for query region, in decimal degrees. West negative.
- **minElev** numeric. Minimum elevation for query. Meters.
- **minDate** character. Minimum date for query. Format: "YYYY-MM-DD"
- **maxDate** character. Maximum date for query. Format: "YYYY-MM-DD"
- **countries** character. Vector of one or more two-letter country codes for query.
- **states** character. Vector of one or more two-letter state or province codes for query.
- **types** character. Vector of one or more sample types for query. See vocabulary in the wiDB template.
- **projects** character. Vector of one or more project codes for query.
- **fields** character. Vector of one or more data fields to return from database. If omitted, returns all default fields. See here for details.
- **tmpdir** character. Directory path to use for unpacking data object.
- **clean** logical. Remove working files after data object is unpacked?

Details

One or more arguments must be provided.

Value

named list. See here for details.

- **data** dataframe. Data records for isotope samples returned by query.
- **projects** dataframe. Provenance information associated with samples returned by query.

Note that some data are embargoed or have been shared under a license that prohibits redistribution. In dataframe **data** values of 9999 indicate that a measurement is available but can’t be obtained directly from the wiDB. Project information in **projects** can be used to contact or visit the primary data source to learn about access to these data. Values of -9999 in **data** indicate no measurement.
Examples

# Download data for US precipitation in the 1990s
vals = wiDB_data(minDate = "1990-01-01", maxDate = "2000-01-01",
                  countries = "US", types = "Precipitation")

# Download data for US Rivers and streams, requesting a subset of data fields
vals = wiDB_data(minDate = "1980-01-01", maxDate = "2000-01-01",
                  countries = "US", types = "River_or_stream",
                  fields = "Site_Name,Latitude,Longitude,d2H")

---

**wiDB_sites**

Obtain information on wiDB sites using a query

**Description**

Obtain information on wiDB sites using a query

**Usage**

```r
wiDB_sites(minLat = NULL, maxLat = NULL, minLong = NULL, maxLong = NULL,
            minElev = NULL, maxElev = NULL, minDate = NULL, maxDate = NULL,
            countries = NULL, states = NULL, types = NULL, projects = NULL)
```

**Arguments**

- `minLat`: numeric. Minimum latitude for query region, in decimal degrees. South negative.
- `maxLat`: numeric. Maximum latitude for query region, in decimal degrees. South negative.
- `minDate`: character. Minimum date for query. Format: "YYYY-MM-DD"
- `maxDate`: character. Maximum date for query. Format: "YYYY-MM-DD"
- `countries`: character. Vector of one or more two-letter country codes for query.
- `states`: character. Vector of one or more two-letter state or province codes for query.
- `types`: character. Vector of one or more sample types for query. See vocabulary in the `wiDB tempate`.
- `projects`: character. Vector of one or more project codes for query.
Details

One or more arguments must be provided.

Value

dataframe. Contains location and summary information for all wiDB sites returned by query. See here for details.

Examples

#Find all sites with tap water data since September, 2019
sites = wiDB_sites(minDate = "2019-09-01", types = "Tap")

wiDB_values
Obtain value lists for categorical fields in wiDB

Description

Obtain value lists for categorical fields in wiDB

Usage

wiDB_values(fields)

Arguments

fields character. One or more field names for which to obtain value lists. Limited to: "countries", "states", "types", and "projects".

Value

named list. Each element is a vector or dataframe containing values for the named field.

Examples

#List all projects in the wiDB
wiDB_values("projects")
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