

Package ‘HiClimR’

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Type Package

Version 2.2.1

Title Hierarchical Climate Regionalization

Description

A tool for Hierarchical Climate Regionalization applicable to any correlation-based clustering. It adds several features and a new clustering method (called, 'regional' linkage) to hierarchical clustering in R ('hclust' function in 'stats' library): data regridding, coarsening spatial resolution, geographic masking, contiguity-constrained clustering, data filtering by mean and/or variance thresholds, data preprocessing (detrending, standardization, and PCA), faster correlation function with preliminary big data support, different clustering methods, hybrid hierarchical clustering, multivariate clustering (MVC), cluster validation, visualization of regionalization results, and exporting region map and mean timeseries into NetCDF-4 file.

The technical details are described in Badr et al. (2015) <[doi:10.1007/s12145-015-0221-7](https://doi.org/10.1007/s12145-015-0221-7)>.

License GPL-3

URL <https://hsbadr.github.io/HiClimR/>,
<https://github.com/hsbadr/HiClimR>

BugReports <https://github.com/hsbadr/HiClimR/issues>

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coarseR	<i>Coarsening spatial resolution for gridded data</i>
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Description

`coarseR` is a helper function that helps coarsening spatial resolution of the input matrix for the `HiClimR` function.

Usage

```
coarseR(x = x, lon = lon, lat = lat, lonStep = 1, latStep = 1, verbose = TRUE)
```

Arguments

`x` an (N rows by M columns) matrix of 'double' values: N objects (spatial points or stations) to be clustered by M observations (temporal points or years). For gridded data, the N objects should be created from the original matrix `x0` using `as.vector(t(x0))`, where `x0` is an (n rows by m columns) matrix, `n = length(unique(lon))` and `m = length(unique(lat))`.

lon	a vector of longitudes with length N. For gridded data, the length may have the value (n) provided that $n * m = N$ where $n = \text{length}(\text{unique}(\text{lon}))$ and $m = \text{length}(\text{unique}(\text{lat}))$.
lat	a vector of latitudes with length N or m. See lon.
lonStep	an integer greater than or equal to 1 for longitude step to coarsen gridded data in the longitudinal direction. If lonStep = 1, gridded data will not be coarsened in the longitudinal direction (the default). If lonStep = 2, every other grid in longitudinal direction will be retained.
latStep	an integer greater than or equal to 1 for latitude step to coarsen gridded data in the latitudinal direction. If latStep = 1, gridded data will not be coarsened in the latitudinal direction (the default). If latStep = 2, every other grid in latitudinal direction will be retained. lonStep and latStep are independent so that user can optionally apply different coarsening level to each dimension.
verbose	logical to print processing information if verbose = TRUE.

Details

For high-resolution data, the computational and memory requirements may not be met on old machines. This function enables the user to use coarser data in any spatial dimension: longitude, latitude, or both. It is available for testing or running HiClimR package on old computers or machines with small memory resources. The rows of output matrix (x component) will be also named by longitude and latitude coordinates. If lonStep = 1 and latStep = 1, `coarseR` function will just rename rows of matrix x.

Value

A list with the following components:

lon	longitude mesh vector for the coarsened data.
lat	latitude mesh vector for the coarsened data.
rownum	original row numbers for the coarsened data.
x	coarsened data of the input data matrix x.

Author(s)

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References

- Hamada S. Badr, Zaitchik, B. F., and Dezfuli, A. K. (2015): A Tool for Hierarchical Climate Regionalization, *Earth Science Informatics*, **8**(4), 949-958, doi: [10.1007/s1214501502217](https://doi.org/10.1007/s1214501502217).
- Hamada S. Badr, Zaitchik, B. F., and Dezfuli, A. K. (2014): Hierarchical Climate Regionalization, *Comprehensive R Archive Network (CRAN)*, <https://cran.r-project.org/package=HiClimR>.

See Also

[HiClimR](#), [HiClimR2nc](#), [validClimR](#), [geogMask](#), [coarseR](#), [fastCor](#), [grid2D](#) and [minSigCor](#).

Examples

```
require(HiClimR)

## Load test case data
x <- TestCase$x

## Generate longitude and latitude mesh vectors
xGrid <- grid2D(lon = unique(TestCase$lon), lat = unique(TestCase$lat))
lon <- c(xGrid$lon)
lat <- c(xGrid$lat)

## Coarsening spatial resolution
xc <- coarseR(x = x, lon = lon, lat = lat, lonStep = 2, latStep = 2)
lon <- xc$lon
lat <- xc$lat
x <- xc$x
```

fastCor

Fast correlation for large matrices

Description

`fastCor` is a helper function that compute Pearson correlation matrix for `HiClimR` and `validClimR` functions. It is similar to `cor` function in R but uses a faster implementation on 64-bit machines (an optimized BLAS library is highly recommended). `fastCor` also uses a memory-efficient algorithm that allows for splitting the data matrix and only compute the upper-triangular part of the correlation matrix. It can be used to compute correlation matrix for the columns of any data matrix.

Usage

```
fastCor(xt, nSplit = 1, upperTri = FALSE, optBLAS = FALSE, verbose = TRUE)
```

Arguments

<code>xt</code>	an (M rows by N columns) matrix of 'double' values: N objects (spatial points or stations) to be clustered by M observations (temporal points or years). It is the transpose of the input matrix <code>x</code> required for <code>HiClimR</code> and <code>validClimR</code> functions.
<code>nSplit</code>	integer number greater than or equal to one, to split the data matrix into <code>nSplit</code> splits of the total number of columns <code>ncol(xt)</code> . If <code>nSplit = 1</code> , the default method will be used to compute correlation matrix for the full data matrix (no splits). If <code>nSplit > 1</code> , the correlation matrix (or the upper-triangular part if <code>upperTri = TRUE</code>) will be allocated and filled with the computed correlation sub-matrix for each split. the first <code>n-1</code> splits have equal size while the last split may include any remaining columns. This is used with <code>upperTri = TRUE</code> to compute only the upper-triangular part of the correlation matrix. The maximum number of splits <code>nSplitMax = floor(N / 2)</code> makes splits with 2 columns; if <code>nSplit > nSplitMax</code> , <code>nSplitMax</code> will be used. Very large number of splits

nSplit makes computation slower but it could handle big data or if the available memory is not enough to allocate the correlation matrix, which helps in solving the “Error: cannot allocate vector of size...” memory limitation problem. It is recommended to start with a small number of splits. If the data is very large compared to the physical memory, it is highly recommended to use a 64-Bit machine with enough memory resources and/or use coarsening feature for gridded data by setting lonStep > 1 and latStep > 1.

upperTri	logical to compute only the upper-triangular half of the correlation matrix if upperTri = TRUE and nSplit > 1., which includes all required info since the correlation/dissimilarity matrix is symmetric. This almost halves memory use, which can be very important for big data.
optBLAS	logical to use optimized BLAS library if installed and optBLAS = TRUE only on 64-bit machines.
verbose	logical to print processing information if verbose = TRUE.

Details

The `fastCor` function computes the correlation matrix by calling the cross product function in the Basic Linear Algebra Subroutines (BLAS) library used by R. A significant performance improvement can be achieved when building R on 64-bit machines with an optimized BLAS library, such as *ATLAS*, *OpenBLAS*, or the commercial *Intel MKL*. For big data, the memory required to allocate the square matrix of correlations may exceed the total amount of physical memory available resulting in “Error: cannot allocate vector of size...”. `fastCor` allows for splitting the data matrix into `nSplit` splits and only computes the upper-triangular part of the correlation matrix with `upperTri = TRUE`. This almost halves memory use, which can be very important for big data. If `nSplit > 1`, the correlation matrix (or the upper-triangular part if `upperTri = TRUE`) will be allocated and filled with computed correlation sub-matrix for each split. the first `n-1` splits have equal size while the last split may include any remaining columns.

Value

An (N rows by N columns) correlation matrix.

Author(s)

Hamada S. Badr <badr@jhu.edu>, Benjamin F. Zaitchik <zaitchik@jhu.edu>, and Amin K. Dezfuli <amin.dezfuli@nasa.gov>.

References

- Hamada S. Badr, Zaitchik, B. F. and Dezfuli, A. K. (2015): A Tool for Hierarchical Climate Regionalization, *Earth Science Informatics*, **8**(4), 949-958, doi: [10.1007/s1214501502217](https://doi.org/10.1007/s1214501502217).
- Hamada S. Badr, Zaitchik, B. F. and Dezfuli, A. K. (2014): Hierarchical Climate Regionalization, *Comprehensive R Archive Network (CRAN)*, <https://cran.r-project.org/package=HiClimR>.

See Also

[HiClimR](#), [HiClimR2nc](#), [validClimR](#), [geogMask](#), [coarseR](#), [fastCor](#), [grid2D](#) and [minSigCor](#).

Examples

```

require(HiClimR)

## Load test case data
x <- TestCase$x

## Use fastCor function to compute the correlation matrix
t0 <- proc.time() ; xcor <- fastCor(t(x)) ; proc.time() - t0
## compare with cor function
t0 <- proc.time() ; xcor0 <- cor(t(x)) ; proc.time() - t0

## Not run:

## Split the data into 10 splits and return upper-triangular half only
xcor10 <- fastCor(t(x), nSplit = 10, upperTri = TRUE)

## End(Not run)

```

geogMask

Geographic mask from longitude and latitude

Description

`geogMask` is a helper function that preprocess input for the `HiClimR` via `geogMask` parameter.

Usage

```

geogMask(continent = NULL, region = NULL, country = NULL, lon = NULL, lat = NULL,
          InDispute = TRUE, verbose = TRUE, plot = FALSE, colPalette = NULL, pch = 15, cex = 1)

```

Arguments

<code>continent</code>	NULL or a string (or array of strings) to specify continent name(s): only one of continent, region, or country should be specified. Valid list of continent names can be obtained by running <code>geogMask()</code> .
<code>region</code>	NULL or a string (or array of strings) to specify region name(s): only one of continent, region, or country should be specified. Valid list of region names can be obtained by running <code>geogMask()</code> .
<code>country</code>	NULL or a string (or array of strings) to specify country ISO3 character code(s): only one of continent, region, or country should be specified. Valid list of country ISO3 character code(s) can be obtained by running <code>geogMask()</code> .
<code>lon</code>	a vector of longitudes with length N. Longitudes takes values from -180 to 180 (not 0 to 360). For gridded data, the length may have the value (n) provided that $n * m = N$ where $n = \text{length}(\text{unique}(\text{lon}))$ and $m = \text{length}(\text{unique}(\text{lat}))$.
<code>lat</code>	a vector of latitudes with length N or m. See <code>lon</code> .

InDispute	a logical: should the areas in dispute be considered for geographic masking by country? If InDispute = TRUE (the default), areas in dispute will be considered as a part of the country.
verbose	logical to print processing information if verbose = TRUE.
plot	logical to call the plotting method if plot = TRUE.
colPalette	a color palette or a list of colors such as that generated by <code>rainbow</code> , <code>heat.colors</code> , <code>topo.colors</code> , <code>terrain.colors</code> or similar functions.
pch	Either an integer specifying a symbol or a single character to be used as the default in plotting points. See points for possible values.
cex	A numerical value giving the amount by which plotting symbols should be magnified relative to the default = 1.

Details

In some applications, a user may want to focus on an area that is a mask-defined subset of the full dataset. For instance, the NASA Tropical Rainfall Measuring Mission (TRMM) data covers ocean and land, while a researcher might be interested in the precipitation variability only over land, a country, or a list of countries (e.g., Nile Basin countries). This masking capability is supported by the `geogMask` helper function. It requires the longitude (`lon`) and latitude (`lat`) vectors together with a string (or array of strings) to specify continent name(s), region name(s), or country ISO3 character code(s) via either `continent`, `region`, or `country` parameters. Valid values for them can be obtained by running `geogMask()`. World mask data is based on the HIU Large Scale International Boundaries (LSIB) data (<https://hiu.state.gov/data>).

Value

A vector of indices for the spatial elements to be masked, as required by `HiClimR`.

Author(s)

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References

Hamada S. Badr, Zaitchik, B. F. and Dezfuli, A. K. (2015): A Tool for Hierarchical Climate Regionalization, *Earth Science Informatics*, **8**(4), 949-958, doi: [10.1007/s1214501502217](https://doi.org/10.1007/s1214501502217).

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See Also

[HiClimR](#), [HiClimR2nc](#), [validClimR](#), [geogMask](#), [coarseR](#), [fastCor](#), [grid2D](#) and [minSigCor](#).

Examples

```

require(HiClimR)

## Load test case data
x <- TestCase$x

## Generate longitude and latitude mesh vectors
xGrid <- grid2D(lon = unique(TestCase$lon), lat = unique(TestCase$lat))
lon <- c(xGrid$lon)
lat <- c(xGrid$lat)

## Check the valid options for geographic masking
geogMask()

## geographic mask for Africa
gMask <- geogMask(continent = "Africa", lon = lon, lat = lat, plot = TRUE)

```

grid2D

Generate longitude and latitude grid matrices

Description

`grid2D` is a helper function that generates longitude and latitude rectangular mesh from short longitude and latitude vectors in gridded data.

Usage

```
grid2D(lon = lon, lat = lat)
```

Arguments

lon	a vector of longitudes with length N. Longitudes takes values from -180 to 180 (not 0 to 360). For gridded data, the length may have the value (n) provided that $n * m = N$ where $n = \text{length}(\text{unique}(\text{lon}))$ and $m = \text{length}(\text{unique}(\text{lat}))$.
lat	a vector of latitudes with length N or m. See lon.

Details

`grid2D` function convert the long latitude and longitude vectors to a rectangular two-dimensional grid for visualization and geographic masking purposes for gridded data in HiClimR package.

Value

A list with the following components:

lon	an (n rows by m columns) matrix of 'double' values for longitude mesh grid, or a vector with length $n * m$.
lat	an (n rows by m columns) matrix of 'double' values for latitude mesh grid, or a vector with length $n * m$.

Author(s)

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References

Hamada S. Badr, Zaitchik, B. F. and Dezfuli, A. K. (2015): A Tool for Hierarchical Climate Regionalization, *Earth Science Informatics*, **8**(4), 949-958, doi: [10.1007/s1214501502217](https://doi.org/10.1007/s1214501502217).

Hamada S. Badr, Zaitchik, B. F. and Dezfuli, A. K. (2014): Hierarchical Climate Regionalization, *Comprehensive R Archive Network (CRAN)*, <https://cran.r-project.org/package=HiClimR>.

See Also

[HiClimR](#), [HiClimR2nc](#), [validClimR](#), [geogMask](#), [coarseR](#), [fastCor](#), [grid2D](#) and [minSigCor](#).

Examples

```
require(HiClimR)

## Load test case data
x <- TestCase$x

## Generate longitude and latitude mesh vectors
xGrid <- grid2D(lon = unique(TestCase$lon), lat = unique(TestCase$lat))
lon <- c(xGrid$lon)
lat <- c(xGrid$lat)
```

HiClimR

Hierarchical Climate Regionalization

Description

HiClimR is a tool for **Hierarchical Climate Regionalization** applicable to any correlation-based clustering. Climate regionalization is the process of dividing an area into smaller regions that are homogeneous with respect to a specified climatic metric. Several features are added to facilitate the applications of climate regionalization (or spatiotemporal analysis in general) and to implement cluster validation with an objective tree cutting to find an optimal number of clusters for a user-specified confidence level. These include options for preprocessing and postprocessing as well as efficient code execution for large datasets and options for splitting big data and computing only the upper-triangular half of the correlation/dissimilarity matrix to overcome memory limitations. Hybrid hierarchical clustering reconstructs the upper part of the tree above a cut to get the best of the available methods. Multivariate clustering (MVC) provides options for filtering all variables before preprocessing, detrending and standardization of each variable, and applying weights for the preprocessed variables. The correlation distance for MVC represents the (weighted) average of distances between all variables.

[HiClimR](#) is the main function that calls all helper functions. It adds several features and a new clustering method (called, *regional linkage*) to hierarchical clustering in R ([hclust](#) function in *stats*

library): data regridding (`grid2D` function), coarsening spatial resolution (`coarseR` function), geographic masking (`geogMask` function), contiguity-constrained clustering, data filtering by mean and/or variance thresholds, data preprocessing (detrending, standardization, and PCA), faster correlation function (`fastCor` function), hybrid hierarchical clustering, multivariate clustering (MVC), cluster validation (`validClimR` and `minSigCor` functions), and visualization of regionalization results, and exporting region map and mean timeseries into NetCDF-4 file.

Badr et al. (2015) describes the regionalization algorithms, features, and data processing tools included in the package and presents a demonstration application in which the package is used to regionalize Africa on the basis of interannual precipitation variability.

HiClimR is applicable to any correlation-based clustering.

Usage

```
HiClimR(

  # Input data matrix (N spatial elements x M observations)
  x = list(),

  # Geographic coordinates
  lon = NULL, lat = NULL,

  # Coarsening spatial resolution
  lonStep = 1, latStep = 1,

  # Geographic masking:
  geogMask = FALSE, gMask = NULL, continent = NULL, region = NULL, country = NULL,

  # Contiguity constraint:
  contigConst = 0,

  # Data thresholds:
  meanThresh = if (inherits(x, "list")) {
    vector("list", length(x))
  } else {
    list(NULL)
  },
  varThresh = if (inherits(x, "list")) {
    as.list(rep(0, length(x)))
  } else {
    list(0)
  },

  # Data preprocessing:
  detrend = if (inherits(x, "list")) {
    as.list(rep(FALSE, length(x)))
  } else {
    list(FALSE)
  },
```

```

standardize = if (inherits(x, "list")) {
  as.list(rep(FALSE, length(x)))
} else {
  list(FALSE)
},
weightMVC = if (inherits(x, "list")) {
  as.list(rep(1, length(x)))
} else {
  list(1)
},
nPC = NULL,

# Clustering options:
method = "ward", hybrid = FALSE, kH = NULL, members = NULL,

# Big data support:
nSplit = 1, upperTri = TRUE, verbose = TRUE,

# Cluster validation:
validClimR = TRUE, rawStats = TRUE, k = NULL, minSize = 1, alpha = 0.05,

# Graphical options:
plot = TRUE, dendrogram = TRUE, colPalette = NULL,
hang = -1, labels = FALSE, pch = 15, cex = 1

)

```

Arguments

- x** an (N rows by M columns) matrix of 'double' values: N objects (spatial points or stations) to be clustered by M observations (temporal points or years). For gridded data, the N objects should be created from the original matrix x_0 using `as.vector(t(x0))`, where x_0 is an (n rows by m columns) matrix, $n = \text{length}(\text{unique}(\text{lon}))$ and $m = \text{length}(\text{unique}(\text{lat}))$. Zero-variance rows (e.g., stations with zero variability) and/or missing values (e.g., years with missing observations) are allowed. The zero-variance rows and the columns with missing values will be removed. However, it is recommended to take care of both zero-variance rows and missing values before clustering. For Multivariate Clustering (MVC), **x** can be a list of `nvar` matrices for the `nvar` variables (one matrix for each variable). The matrixes in **x** list should have the same number of rows (objects: spatial points or stations) Data preprocessing is specified by lists of `meanThresh`, `varThresh`, `detrend`, and `standardize` with the same length of **x** where `length(x) = nvar`. Each variable is separately preprocessed to allow for all possible options. However, it is strongly recommended to standardize all variables since their magnitude range could be different. Note that: for gridded data, the rows of input data matrix for each variable is ordered by longitudes (check `TestCase$x` for more details).

lon	a vector of longitudes with length N. Longitudes takes values from -180 to 180 (not 0 to 360). For gridded data, the length may have the value (n) provided that $n * m = N$ where $n = \text{length}(\text{unique}(\text{lon}))$ and $m = \text{length}(\text{unique}(\text{lat}))$.
lat	a vector of latitudes with length N or m. See lon.
lonStep	an integer greater than or equal to 1 for longitude step to coarsen gridded data in the longitudinal direction. If lonStep = 1, gridded data will not be coarsened in the longitudinal direction (the default). If lonStep = 2, every other grid in longitudinal direction will be retained.
latStep	an integer greater than or equal to 1 for latitude step to coarsen gridded data in the latitudinal direction. If latStep = 1, gridded data will not be coarsened in the latitudinal direction (the default). If latStep = 2, every other grid in latitudinal direction will be retained. lonStep and latStep are independent so that user can optionally apply different coarsening level to each dimension.
geogMask	a logical: if geogMask = TRUE, <code>geogMask</code> function will be called. Additional arguments are required. It requires the longitude and latitude vector together with a string (or array of strings) to specify continent, region name(s), or country ISO3 character code(s). If gMask != NULL, the provided gmask vector will be used for geographic masking without calling <code>geogMask</code> .
gMask	A vector of indices for the spatial elements to be masked, as required by <code>HiClimR</code> . This is typically an output vector from <code>geogMask</code> function. This helps in saving time when the same geographic mask will be used many times.
continent	NULL or a string (or array of strings) to specify continent name(s): only one of continent, region, or country should be specified. Valid list of continent names can be obtained by running <code>geogMask()</code> .
region	NULL or a string (or array of strings) to specify region name(s): only one of continent, region, or country should be specified. Valid list of region names can be obtained by running <code>geogMask()</code> .
country	NULL or a string (or array of strings) to specify country ISO3 character code(s): only one of continent, region, or country should be specified. Valid list of country ISO3 character code(s) can be obtained by running <code>geogMask()</code> .
contigConst	0 or a positive value for contiguity constraint: This is used to apply weighted geographic distances on the initial dissimilarity matrix between the variable(s) in the data matrix x. It should be a positive real value ($\text{contigConst} > 0$, recommended between 0 and 1) where $\text{contigConst} = 0$ means that clustering is completely based on the data with no constraints for contiguity while $\text{contigConst} > 0$ gives priority to merging contiguous points. Geographic coordinates lon and lat are required to compute the geographic distances. regional linkage method does not support contiguity constraints.
meanThresh	NULL or a threshold for the temporal mean: This is used with <code>varThresh</code> to mask zero- and near-zero-variance data, Observations with mean less than or equal to <code>meanThresh</code> will be removed. If <code>meanThresh = NULL</code> , then the <code>varThresh</code> will be used either to mask zero-variance data by default or by increased variance threshold to mask near-zero-variance data. For Multivariate Clustering (MVC), <code>meanThresh</code> is a list of thresholds with the same length of x where $\text{length}(x) = \text{nvar}$. Each variable is separately preprocessed to allow for all possible options.

However, it is strongly recommended to standardize all variables since their magnitude range could be different.

varThresh	zero or a threshold for the temporal variance: This is used with meanThresh to mask zero- and near-zero-variance data, Observations with variance less than or equal to varThresh will be removed. If varThresh = 0, then the zero-variance data will be masked (default). For Multivariate Clustering (MVC), varThresh is a list of thresholds with the same length of x where length(x) = nvar. Each variable is separately preprocessed to allow for all possible options. However, it is strongly recommended to standardize all variables since their magnitude range could be different.
detrend	a logical: should the data be detrended before clustering? Detrending (removing the linear trend) is important when variations from temporal point to another is of interest (e.g., interannual variability). The columns of the data matrix x should be temporally ordered (constant step size) or have appropriate names (e.g., colnames(x) = years[1:M]). For Multivariate Clustering (MVC), detrend is a list of thresholds with the same length of x where length(x) = nvar. Each variable is separately preprocessed to allow for all possible options. However, it is strongly recommended to standardize all variables since their magnitude range could be different.
standardize	a logical: should the data be standardized before clustering? The standardized data makes use of the mean of equally-weighted objects within each cluster (cluster mean = mean of standardized variables within the cluster). Otherwise, the mean of raw data will be used (cluster mean = mean of raw variables within the cluster). The variance of the mean is updated at each agglomeration step. For Multivariate Clustering (MVC), standardize is a list of thresholds with the same length of x where length(x) = nvar. Each variable is separately preprocessed to allow for all possible options. However, it is strongly recommended to standardize all variables since their magnitude range could be different.
weightMVC	a list of positive weights (weightMVC > 0) for Multivariate Clustering (MVC) with the same length of x where length(x) = number of variables. The filtered variables are weighted and combined by column (for each object: spatial points or stations) after preprocessing (detrending and standardization) and before PCA (if requested) and computing the correlation/dissimilarity matrix. The default weight is weightMVC = 1 for all variables.
nPC	NULL or number of principal components (PCs) to be retained. If nPC = NULL, then the raw data will be used for clustering. Otherwise, the data will be filtered and reconstructed using nPC PCs obtained from SVD-based PCA. The detrend and/or standardize options will be applied, if requested, before PCA.
method	the agglomeration method to be used. This should be (an unambiguous abbreviation of) one of "regional", "ward", "single", "complete", "average", "mcquitty", "median" or "centroid". The default is "ward" method.
hybrid	a logical: should the upper part of the tree be reconstructed using regional linkage clustering method? This adds hybrid hierarchical clustering feature to get the best of the available methods. It utilizes the pros of available methods, especially the better overall homogeneity in ward's method and the separation, contiguity, and objective tree cut of regional linkage method. If hybrid = FALSE, only the default clustering using the selected method will be used (i.e.,

	no hybrid clustering). Otherwise, the upper part of the tree will be reconstructed above a cut of <code>kH</code> clusters using regional linkage method. Note: hybrid option is redundant when using regional linkage as the main clustering method.
<code>kH</code>	NULL or an integer for the number of regions/clusters in the upper part of the tree to be reconstructed with regional linkage method, if <code>hybrid = TRUE</code> . If <code>kH = NULL</code> , the tree will be reconstructed for the upper part with the first merging cost larger than the mean merging cost for the entire tree. If hybrid clustering is requested, the updated upper part of the tree will be used for cluster validation, and so <code>kH</code> should be greater than the final number of clusters <code>k</code> , if selected.
<code>members</code>	NULL or a vector with length size of <code>d</code> . See the ‘Details’ section.
<code>nSplit</code>	integer number greater than or equal to one, to split the data matrix into <code>nSplit</code> splits of the total number of columns <code>ncol(xt)</code> . If <code>nSplit = 1</code> , the default method will be used to compute correlation matrix for the full data matrix (no splits). If <code>nSplit > 1</code> , the correlation matrix (or the upper-triangular part if <code>upperTri = TRUE</code>) will be allocated and filled with the computed correlation sub-matrix for each split. the first <code>n-1</code> splits have equal size while the last split may include any remaining columns. This is used with <code>upperTri = TRUE</code> to compute only the upper-triangular part of the correlation matrix. The maximum number of splits <code>nSplitMax = floor(N / 2)</code> makes splits with 2 columns; if <code>nSplit > nSplitMax</code> , <code>nSplitMax</code> will be used. Very large number of splits <code>nSplit</code> makes computation slower but it could handle big data or if the available memory is not enough to allocate the correlation matrix, which helps in solving the “Error: cannot allocate vector of size...” memory limitation problem. It is recommended to start with a small number of splits. If the data is very large compared to the physical memory, it is highly recommended to use a 64-Bit machine with enough memory resources and/or use coarsening feature for gridded data by setting <code>lonStep > 1</code> and <code>latStep > 1</code> .
<code>upperTri</code>	logical to compute only the upper-triangular half of the correlation matrix if <code>upperTri = TRUE</code> and <code>nSplit > 1</code> ., which includes all required info since the correlation/dissimilarity matrix is symmetric. This almost halves memory use, which can be very important for big data.
<code>verbose</code>	logical to print processing information if <code>verbose = TRUE</code> .
<code>validClimR</code>	a logical: If <code>validClimR = TRUE</code> , <code>validClimR</code> will be called to compute validation indices including statistical summary for inter- and intra-cluster correlations. This is computationally expensive. It can also objectively cut the dendrogram tree for regional clustering method, if <code>k = NULL</code> .
<code>rawStats</code>	a logical: should validation indices be computed based on the raw data or PCA-filtered data?
<code>k</code>	NULL or an integer <code>k > 1</code> for the number of regions/clusters. Only for regional linkage method, <code>k = NULL</code> is supported, where the "optimal" number of regions will be used at a user specified significance level <code>alpha</code> . It is required to specify number of clusters <code>k</code> for the other methods, since they are not based on inter-cluster correlation. If <code>k = NULL</code> for these methods (except regional) linkage, the <code>validClimR</code> will be aborted. One can use <code>validClimR</code> function to compute inter-cluster correlation at different number of clusters to objectively cut the tree for the other methods, which could be computationally expensive to cover the entire merging history for large number of spatial elements.

<code>minSize</code>	minimum cluster size. The regional linkage method tend to isolate noisy data in small clusters. The <code>minSize</code> can be used to exclude these very small clusters from the <code>statSum</code> statistical summary, because they are most likely noisy data that need to be checked in a quality control step. The analysis may be then repeated.
<code>alpha</code>	confidence level: the default is <code>alpha = 0.05</code> for 95% confidence level.
<code>plot</code>	logical to call the plotting method if <code>plot = TRUE</code> .
<code>dendrogram</code>	logical to enable or disable dendrogram plotting.
<code>colPalette</code>	a color palette or a list of colors such as that generated by <code>rainbow</code> , <code>heat.colors</code> , <code>topo.colors</code> , <code>terrain.colors</code> or similar functions.
<code>hang</code>	The fraction of the plot height by which labels should hang below the rest of the plot. A negative value will cause the labels to hang down from 0.
<code>labels</code>	A character vector of labels for the leaves of the tree. By default the row names or row numbers of the original data are used. If <code>labels = FALSE</code> no labels at all are plotted.
<code>pch</code>	Either an integer specifying a symbol or a single character to be used as the default in plotting points. See points for possible values.
<code>cex</code>	A numerical value giving the amount by which plotting symbols should be magnified relative to the <code>default = 1</code> .

Details

`HiClimR` function is based on `hclust`, which now uses an optimized algorithm to deal with only the upper/lower triangular-half of the symmetric dissimilarity matrix instead of the old algorithm that uses the full matrix in the merging steps. It performs a hierarchical cluster analysis using Pearson correlation distance dissimilarity for the N objects being clustered. Initially, each object is assigned to its own cluster and then the algorithm proceeds iteratively, at each stage joining the two most similar clusters, continuing until there is just a single cluster. At each stage distances between clusters are recomputed by a dissimilarity update formula according to the particular clustering method being used.

All clustering methods in `hclust` are included. The *regional* linkage method minimizes inter-cluster correlations between cluster means (see Badr et al. 2015). *Ward's* minimum variance method aims at finding compact, spherical clusters. The *complete linkage* method finds similar clusters. The *single linkage* method (which is closely related to the minimal spanning tree) adopts a 'friends of friends' clustering strategy. The other methods can be regarded as aiming for clusters with characteristics somewhere between the single and complete link methods. Note however, that methods "median" and "centroid" are *not* leading to a *monotone distance* measure, or equivalently the resulting dendrograms can have so called *inversions* (which are hard to interpret).

The regional linkage method is explained in the context of a spatiotemporal problem, in which N spatial elements (e.g., weather stations) are divided into k regions, given that each element has a time series of length M . It is based on inter-regional correlation distance between the temporal means of different regions (or elements at the first merging step). It modifies the update formulae of average linkage method by incorporating the standard deviation of the merged region timeseries, which is a function of the correlation between the individual regions, and their standard deviations before merging. It is equal to the average of their standard deviations if and only if the correlation

between the two merged regions is 100%. In this special case, the regional linkage method is reduced to the classic average linkage clustering method.

If `members` != NULL, then `d` is taken to be a dissimilarity matrix between clusters instead of dissimilarities between singletons and `members` gives the number of observations per cluster. This way the hierarchical cluster algorithm can be ‘started in the middle of the dendrogram’, e.g., in order to reconstruct the part of the tree above a cut (see examples). Dissimilarities between clusters can be efficiently computed (i.e., without `hclust` itself) only for a limited number of distance/linkage combinations, the simplest one being squared Euclidean distance and centroid linkage. In this case the dissimilarities between the clusters are the squared Euclidean distances between cluster means.

In hierarchical cluster displays, a decision is needed at each merge to specify which subtree should go on the left and which on the right. Since, for n observations there are $n - 1$ merges, there are $2^{(n-1)}$ possible orderings for the leaves in a cluster tree, or dendrogram. The algorithm used in `hclust` is to order the subtree so that the tighter cluster is on the left (the last, i.e., most recent, merge of the left subtree is at a lower value than the last merge of the right subtree). Single observations are the tightest clusters possible, and merges involving two observations place them in order by their observation sequence number.

Value

An object of class `HiClimR` and `hclust`, which describes the tree produced by the clustering process. The object is a list with the following components:

<code>merge</code>	an $n - 1$ by 2 matrix. Row i of <code>merge</code> describes the merging of clusters at step i of the clustering. If an element j in the row is negative, then observation $-j$ was merged at this stage. If j is positive then the merge was with the cluster formed at the (earlier) stage j of the algorithm. Thus negative entries in <code>merge</code> indicate agglomerations of singletons, and positive entries indicate agglomerations of non-singletons.
<code>height</code>	a set of $n - 1$ real values (non-decreasing for ultrametric trees). The clustering <i>height</i> : that is, the value of the criterion associated with the clustering method for the particular agglomeration.
<code>order</code>	a vector giving the permutation of the original observations suitable for plotting, in the sense that a cluster plot using this ordering and <code>merge</code> will not have crossings of the branches.
<code>labels</code>	labels for each of the objects being clustered.
<code>method</code>	the cluster method that has been used.
<code>call</code>	the call which produced the result.
<code>dist.method</code>	the distance that has been used to create <code>d</code> (only returned if the distance object has a “method” attribute).
<code>skip</code>	a vector of <code>lonStep</code> and <code>latStep</code> if <code>coarseR = TRUE</code> .
<code>PCA</code>	if <code>nPC</code> != NULL, the eigenvalues, explained variance, and accumulated variance will be stored here.
<code>coords</code>	an (<code>NC</code> rows by 2 columns) matrix of ‘double’ values: longitude and latitude coordinates for the preprocessed data used for clustering, where <code>NC</code> is the number of spatial elements to be clustered after coarsening spatial resolution by <code>lonStep</code> and <code>latStep</code> . It will be returned only if <code>lon</code> and <code>lat</code> vectors were defined. It will be similar to the provided <code>lon</code> and <code>lat</code> if <code>lonStep = 1</code> and <code>latStep = 1</code> .

nvars	number of variables used for multivariate clustering (MVC).
ncols	number of columns for each variable.
data	the preprocessed data used for clustering will be stored here. If <code>rawData = TRUE</code> and <code>nPC != NULL</code> , the preprocessed data without filtering (PCA) will be returned here.
mask	a vector of the indices of masked spatial points by both geographic masking and data thresholds.
treeH	An object of class <code>hclust</code> , which describes the upper part of the tree reconstructed by the hybrid clustering process if <code>hybrid = TRUE</code> .

If `validClimR = TRUE`, an object of class `HiClimR`, which produces indices for validating the tree produced by the clustering process, will be merged in the dendrogram tree above. This object is a list with the following components:

cutLevel	the minimum significant correlation used for objective tree cut together with the corresponding confidence level.
clustMean	the cluster means which are the region's mean timeseries for all selected regions.
clustSize	cluster sizes for all selected regions.
clustFlag	a flag 0 or 1 to indicate the cluster used in <code>statSum</code> validation indices (<code>interCor</code> , <code>intraCor</code> , <code>diffCor</code> , and <code>statSum</code>), based on <code>minSize</code> minimum cluster size. If <code>clustFlag = 0</code> , the cluster has been excluded because its size is less than the <code>minSize</code> minimum cluster size. The sum of <code>clustFlag</code> elements represents the selected number clusters.
interCor	inter-cluster correlations for all selected regions. It is the inter-cluster correlations between cluster means. The maximum inter-cluster correlation is a measure for separation or contiguity, and it is used for objective tree cut (to find the "optimal" number of clusters).
intraCor	intra-cluster correlations for all selected regions. It is the intra-cluster correlations between the mean of each cluster and its members. The average intra-cluster correlation is a weighted average for all clusters, and it is a measure for homogeneity.
diffCor	difference between intra-cluster correlation and maximum inter-cluster correlation for all selected regions.
statSum	overall statistical summary for <code>interCluster</code> , <code>intraCor</code> , and <code>diffCor</code> .
region	ordered regions vector of size N number of spatial elements for the selected number of clusters, after excluding the small clusters defined by <code>minSize</code> argument.
regionID	ordered regions ID vector of length equals the selected number of clusters, after excluding the small clusters defined by <code>minSize</code> argument. It helps in mapping ordered regions and their actual names before ordering. Only the <code>region</code> component uses ordered ID, while other components use the names used during the clustering process.

There are `print`, `plot` and `identify` (see `identify.hclust`) methods and `rect.hclust()` functions for `hclust` objects.

Author(s)

Hamada S. Badr <badr@jhu.edu>, Benjamin F. Zaitchik <zaitchik@jhu.edu>, and Amin K. Dezfuli <amin.dezfuli@nasa.gov>. HiClimR is a modification of `hclust` function, which is based on Fortran code contributed to STATLIB by F. Murtagh.

References

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- Source Code: <https://github.com/hsbadr/HiClimR>

See Also

[HiClimR](#), [HiClimR2nc](#), [validClimR](#), [geogMask](#), [coarseR](#), [fastCor](#), [grid2D](#), [minSigCor](#), [identify.hclust](#), [rect.hclust](#), [cutree](#), [dendrogram](#), and [kmeans](#).

Examples

```
require(HiClimR)

#-----#
# Typical use of HiClimR for single-variate clustering: #
#-----#

## Load the test data included/loaded in the package (1 degree resolution)
x <- TestCase$x
lon <- TestCase$lon
lat <- TestCase$lat

## Generate/check longitude and latitude mesh vectors for gridded data
xGrid <- grid2D(lon = unique(TestCase$lon), lat = unique(TestCase$lat))
lon <- c(xGrid$lon)
```

```

lat <- c(xGrid$lat)

## Single-Variate Hierarchical Climate Regionalization
y <- HiClimR(x, lon = lon, lat = lat, lonStep = 1, latStep = 1, geogMask = FALSE,
  continent = "Africa", meanThresh = 10, varThresh = 0, detrend = TRUE,
  standardize = TRUE, nPC = NULL, method = "ward", hybrid = FALSE, kH = NULL,
  members = NULL, nSplit = 1, upperTri = TRUE, verbose = TRUE,
  validClimR = TRUE, k = 12, minSize = 1, alpha = 0.01,
  plot = TRUE, colPalette = NULL, hang = -1, labels = FALSE)

## For more examples: https://github.com/hsbadr/HiClimR#examples

## Not run:

#-----#
# Additional Examples:                               #
#-----#

## Use Ward's method
y <- HiClimR(x, lon = lon, lat = lat, lonStep = 1, latStep = 1, geogMask = FALSE,
  continent = "Africa", meanThresh = 10, varThresh = 0, detrend = TRUE,
  standardize = TRUE, nPC = NULL, method = "ward", hybrid = FALSE, kH = NULL,
  members = NULL, nSplit = 1, upperTri = TRUE, verbose = TRUE,
  validClimR = TRUE, k = 12, minSize = 1, alpha = 0.01,
  plot = TRUE, colPalette = NULL, hang = -1, labels = FALSE)

## Use data splitting for big data
y <- HiClimR(x, lon = lon, lat = lat, lonStep = 1, latStep = 1, geogMask = FALSE,
  continent = "Africa", meanThresh = 10, varThresh = 0, detrend = TRUE,
  standardize = TRUE, nPC = NULL, method = "ward", hybrid = TRUE, kH = NULL,
  members = NULL, nSplit = 10, upperTri = TRUE, verbose = TRUE,
  validClimR = TRUE, k = 12, minSize = 1, alpha = 0.01,
  plot = TRUE, colPalette = NULL, hang = -1, labels = FALSE)

## Use hybrid Ward-Regional method
y <- HiClimR(x, lon = lon, lat = lat, lonStep = 1, latStep = 1, geogMask = FALSE,
  continent = "Africa", meanThresh = 10, varThresh = 0, detrend = TRUE,
  standardize = TRUE, nPC = NULL, method = "ward", hybrid = TRUE, kH = NULL,
  members = NULL, nSplit = 1, upperTri = TRUE, verbose = TRUE,
  validClimR = TRUE, k = 12, minSize = 1, alpha = 0.01,
  plot = TRUE, colPalette = NULL, hang = -1, labels = FALSE)
## Check sensitivity to kH for the hybrid method above

#-----#
# Typical use of HiClimR for multivariate clustering:   #
#-----#

## Load the test data included/loaded in the package (1 degree resolution)
x1 <- TestCase$x
lon <- TestCase$lon
lat <- TestCase$lat

```

```

## Generate/check longitude and latitude mesh vectors for gridded data
xGrid <- grid2D(lon = unique(TestCase$lon), lat = unique(TestCase$lat))
lon <- c(xGrid$lon)
lat <- c(xGrid$lat)

## Test if we can replicate single-variate region map with repeated variable
y <- HiClimR(x=list(x1, x1), lon = lon, lat = lat, lonStep = 1, latStep = 1,
  geogMask = FALSE, continent = "Africa", meanThresh = list(10, 10),
  varThresh = list(0, 0), detrend = list(TRUE, TRUE), standardize = list(TRUE, TRUE),
  nPC = NULL, method = "ward", hybrid = FALSE, KH = NULL,
  members = NULL, nSplit = 1, upperTri = TRUE, verbose = TRUE,
  validClimR = TRUE, k = 12, minSize = 1, alpha = 0.01,
  plot = TRUE, colPalette = NULL, hang = -1, labels = FALSE)

## Generate a random matrix with the same number of rows
x2 <- matrix(rnorm(nrow(x1) * 100, mean=0, sd=1), nrow(x1), 100)

## Multivariate Hierarchical Climate Regionalization
y <- HiClimR(x=list(x1, x2), lon = lon, lat = lat, lonStep = 1, latStep = 1,
  geogMask = FALSE, continent = "Africa", meanThresh = list(10, NULL),
  varThresh = list(0, 0), detrend = list(TRUE, FALSE), standardize = list(TRUE, TRUE),
  weightMVC = list(1, 1), nPC = NULL, method = "ward", hybrid = FALSE, KH = NULL,
  members = NULL, nSplit = 1, upperTri = TRUE, verbose = TRUE,
  validClimR = TRUE, k = 12, minSize = 1, alpha = 0.01,
  plot = TRUE, colPalette = NULL, hang = -1, labels = FALSE)
## You can apply all clustering methods and options

#-----#
# Miscellaneous examples to provide more information about functionality and usage #
# of the helper functions that can be used separately or for other applications. #
#-----#

## Load test case data
x <- TestCase$x

## Generate longitude and latitude mesh vectors
xGrid <- grid2D(lon = unique(TestCase$lon), lat = unique(TestCase$lat))
lon <- c(xGrid$lon)
lat <- c(xGrid$lat)

## Coarsening spatial resolution
xc <- coarseR(x = x, lon = lon, lat = lat, lonStep = 2, latStep = 2)
lon <- xc$lon
lat <- xc$lat
x <- xc$x

## Use fastCor function to compute the correlation matrix
t0 <- proc.time(); xcor <- fastCor(t(x)); proc.time() - t0
## compare with cor function
t0 <- proc.time(); xcor0 <- cor(t(x)); proc.time() - t0

## Check the valid options for geographic masking
geogMask()

```

```

## geographic mask for Africa
gMask <- geogMask(continent = "Africa", lon = lon, lat = lat, plot = TRUE,
  colPalette = NULL)

## Hierarchical Climate Regionalization Without geographic masking
y <- HiClimR(x, lon = lon, lat = lat, lonStep = 1, latStep = 1, geogMask = FALSE,
  continent = "Africa", meanThresh = 10, varThresh = 0, detrend = TRUE,
  standardize = TRUE, nPC = NULL, method = "ward", hybrid = FALSE, kH = NULL,
  members = NULL, nSplit = 1, upperTri = TRUE, verbose = TRUE,
  validClimR = TRUE, k = 12, minSize = 1, alpha = 0.01,
  plot = TRUE, colPalette = NULL, hang = -1, labels = FALSE)

## With geographic masking (you may specify the mask produced above to save time)
y <- HiClimR(x, lon = lon, lat = lat, lonStep = 1, latStep = 1, geogMask = TRUE,
  continent = "Africa", meanThresh = 10, varThresh = 0, detrend = TRUE,
  standardize = TRUE, nPC = NULL, method = "ward", hybrid = FALSE, kH = NULL,
  members = NULL, nSplit = 1, upperTri = TRUE, verbose = TRUE,
  validClimR = TRUE, k = 12, minSize = 1, alpha = 0.01,
  plot = TRUE, colPalette = NULL, hang = -1, labels = FALSE)

## With geographic masking and contiguity constraint
## Change contigConst as appropriate
y <- HiClimR(x, lon = lon, lat = lat, lonStep = 1, latStep = 1, geogMask = TRUE,
  continent = "Africa", contigConst = 1, meanThresh = 10, varThresh = 0, detrend = TRUE,
  standardize = TRUE, nPC = NULL, method = "ward", hybrid = FALSE, kH = NULL,
  members = NULL, nSplit = 1, upperTri = TRUE, verbose = TRUE,
  validClimR = TRUE, k = 12, minSize = 1, alpha = 0.01,
  plot = TRUE, colPalette = NULL, hang = -1, labels = FALSE)

## Find minimum significant correlation at 95
rMin <- minSigCor(n = nrow(x), alpha = 0.05, r = seq(0, 1, by = 1e-06))

## Validtion of Hierarchical Climate Regionalization
z <- validClimR(y, k = 12, minSize = 1, alpha = 0.01,
  plot = TRUE, colPalette = NULL)

## Apply minimum cluster size (minSize = 25)
z <- validClimR(y, k = 12, minSize = 25, alpha = 0.01,
  plot = TRUE, colPalette = NULL)

## The optimal number of clusters, including small clusters
k <- length(z$clustFlag)

## The selected number of clusters, after excluding small clusters (if minSize > 1)
ks <- sum(z$clustFlag)

## Dendrogram plot
plot(y, hang = -1, labels = FALSE)

## Tree cut
cutTree <- cutree(y, k = k)
table(cutTree)

```

```

## Visualization for gridded data
RegionsMap <- matrix(y$region, nrow = length(unique(y$coords[, 1])), byrow = TRUE)
colPalette <- colorRampPalette(c("#00007F", "blue", "#007FFF", "cyan",
  "#7FFF7F", "yellow", "#FF7F00", "red", "#F00000"))
image(unique(y$coords[, 1]), unique(y$coords[, 2]), RegionsMap, col = colPalette(ks))

## Visualization for gridded or ungridded data
plot(y$coords[, 1], y$coords[, 2], col = colPalette(max(Regions, na.rm = TRUE))[y$region],
  pch = 15, cex = 1)

## Export region map and mean timeseries into NetCDF-4 file
y.nc <- HiClimR2nc(y=y, ncfile="HiClimR.nc", timeunit="years", dataunit="mm")
## The NetCDF-4 file is still open to add other variables or close it
nc_close(y.nc)

## End(Not run)

```

HiClimR2nc

Export NetCDF-4 file for Hierarchical Climate Regionalization

Description

`HiClimR2nc` is a helper function that exports region map and mean timeseries into NetCDF-4 file, using the `ncdf4` package.

Usage

```
HiClimR2nc(y = NULL, ncfile = "HiClimR.nc", timeunit = "", dataunit = "")
```

Arguments

<code>y</code>	a dendrogram tree produced by <code>HiClimR</code> .
<code>ncfile</code>	Path and name of the NetCDF-4 file to be created.
<code>timeunit</code>	an optional character string for the time units, A zero-length string (default: <code>timeunit=""</code>) removes units attribute.
<code>dataunit</code>	an optional character string for the data units, A zero-length string (default: <code>timeunit=""</code>) removes units attribute.

Details

`HiClimR2nc` function exports region map and mean timeseries from `HiClimR` tree into NetCDF-4 file, using the `ncdf4` package. The NetCDF-4 file will be open to add other variables, if needed. It is important to close the created file using `nc_close`, which flushes any unwritten data to disk.

Value

An object of class `ncdf4`, which has the fields described in `nc_open`.

Author(s)

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References

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Hamada S. Badr, Zaitchik, B. F. and Dezfuli, A. K. (2014): Hierarchical Climate Regionalization, *Comprehensive R Archive Network (CRAN)*, <https://cran.r-project.org/package=HiClimR>.

See Also

[HiClimR](#), [HiClimR2nc](#), [validClimR](#), [geogMask](#), [coarseR](#), [fastCor](#), [grid2D](#) and [minSigCor](#).

Examples

```
require(HiClimR)
require(ncdf4)

## Load test case data
x <- TestCase$x

## Generate longitude and latitude mesh vectors
xGrid <- grid2D(lon = unique(TestCase$lon), lat = unique(TestCase$lat))
lon <- c(xGrid$lon)
lat <- c(xGrid$lat)

## Hierarchical Climate Regionalization
y <- HiClimR(x, lon = lon, lat = lat, lonStep = 1, latStep = 1, geogMask = FALSE,
  continent = "Africa", meanThresh = 10, varThresh = 0, detrend = TRUE,
  standardize = TRUE, nPC = NULL, method = "ward", hybrid = FALSE,
  kH = NULL, members = NULL, validClimR = TRUE, k = 12, minSize = 1,
  alpha = 0.01, plot = TRUE, colPalette = NULL, hang = -1, labels = FALSE)

## Not run:

## Export region map and mean timeseries into NetCDF-4 file
y.nc <- HiClimR2nc(y=y, ncfile="HiClimR.nc", timeunit="years", dataunit="mm")
## The NetCDF-4 file is still open to add other variables or close it
nc_close(y.nc)

## End(Not run)
```

`minSigCor`*Minimum significant correlation for a sample size*

Description

`minSigCor` is a helper function that estimates the minimum significant correlation for a sample size n at a confidence level defined by the argument `alpha`.

Usage

```
minSigCor(n = 41, alpha = 0.05, r = seq(0, 1, by = 1e-6))
```

Arguments

<code>n</code>	sample size or the length of a timeseries vector.
<code>alpha</code>	confidence level: the default is <code>alpha = 0.05</code> for 95% confidence level.
<code>r</code>	a vector of values from 0 to 1 to search for the minimum significant correlation for the user-specified sample size n at confidence level <code>alpha</code> . This should be a subset of the valid positive correlation range 0-1. The default is to search for the minimum significant correlation in the complete range 0-1 with a very fine step of $1e-6$. For faster computations, the user may set a shorter range with larger step (e.g., <code>seq(0.1, 0.5, by=1e-3)</code>).

Details

`minSigCor` function estimates the minimum significant correlation for a sample size (number of observations or temporal points in a timeseries) at a certain confidence level selected by the argument `alpha` and an optional search range `r`. It is called by `validClimR` function objective tree cut based on the specified confidence level.

Value

A positive value between 0 and 1 for the estimated the minimum significant correlation.

Author(s)

Hamada S. Badr <badr@jhu.edu>, Benjamin F. Zaitchik <zaitchik@jhu.edu>, and Amin K. Dezfuli <amin.dezfuli@nasa.gov>.

References

Hamada S. Badr, Zaitchik, B. F. and Dezfuli, A. K. (2015): A Tool for Hierarchical Climate Regionalization, *Earth Science Informatics*, **8**(4), 949-958, doi: [10.1007/s1214501502217](https://doi.org/10.1007/s1214501502217).

Hamada S. Badr, Zaitchik, B. F. and Dezfuli, A. K. (2014): Hierarchical Climate Regionalization, *Comprehensive R Archive Network (CRAN)*, <https://cran.r-project.org/package=HiClimR>.

See Also

[HiClimR](#), [HiClimR2nc](#), [validClimR](#), [geogMask](#), [coarseR](#), [fastCor](#), [grid2D](#) and [minSigCor](#).

Examples

```
require(HiClimR)

## Find minimum significant correlation at 95% confidence level
rMin <- minSigCor(n = 41, alpha = 0.05, r = seq(0, 1, by = 1e-06))
```

TestCase

Test Data for Functionality Demonstration of HiClimR Package

Description

This data is a subset of University of East Anglia Climatic Research Unit (CRU) TS (timeseries) precipitation dataset version 3.2.

Usage

```
data(TestCase)
```

Format

TestCase is a list of three components: x, lon, and lat. x is an (6400 rows by 41 columns) matrix as required for [HiClimR](#) function. The rows represent spatial points (or stations), while the columns represent observations (temporal points or years). lon and lat are vectors of length 80 for unique longitudes and latitudes coordinates, where $80 * 80 = 6400$ for this gridded data.

Details

CRU TS 3.21 data (1901-2012) is monthly gridded precipitation with 0.5 degree resolution. This test data is a subset with 1 degree resolution for African precipitation in January, 1949-1989.

Source

Climatic Research Unit (CRU) time-series datasets of variations in climate with variations in other phenomena.

References

- Hamada S. Badr, Zaitchik, B. F. and Dezfuli, A. K. (2015): A Tool for Hierarchical Climate Regionalization, *Earth Science Informatics*, **8**(4), 949-958, doi: [10.1007/s1214501502217](https://doi.org/10.1007/s1214501502217).
- Hamada S. Badr, Zaitchik, B. F. and Dezfuli, A. K. (2014): Hierarchical Climate Regionalization, *Comprehensive R Archive Network (CRAN)*, <https://cran.r-project.org/package=HiClimR>.
- Harris, I., Jones, P. D., Osborn, T. J., and Lister, D. H. (2014): Updated high-resolution grids of monthly climatic observations - the CRU TS3.10 Dataset, *International journal of climatology*, **34**, 623-642, doi: [10.1002/joc.3711](https://doi.org/10.1002/joc.3711).

Examples

```
require(HiClimR)

x <- TestCase$x
dim(x)
colnames(x)
```

validClimR

Validation of Hierarchical Climate Regionalization

Description

`validClimR` computes indices for cluster validation, and an objective tree cut for regional linkage clustering method.

Usage

```
validClimR(y = NULL, k = NULL, minSize = 1, alpha = 0.05, verbose = TRUE,
           plot = FALSE, colPalette = NULL, pch = 15, cex = 1)
```

Arguments

<code>y</code>	a dendrogram tree produced by <code>HiClimR</code> .
<code>k</code>	NULL or a n integer $k > 1$ for the number of regions/clusters. Only for regional linkage method, $k = \text{NULL}$ is supported, where the "optimal" number of regions will be used at a user specified significance level α . It is required to specify number of clusters k for the other methods, since they are not based on inter-cluster correlation. If $k = \text{NULL}$ for these methods (except regional) linkage, the <code>validClimR</code> will be aborted. One can use <code>validClimR</code> function to compute inter-cluster correlation at different number of clusters to objectively cut the tree for the other methods, which could be computationally expensive to cover the entire merging history for large number of spatial elements.
<code>minSize</code>	minimum cluster size. The regional linkage method tend to isolate noisy data in small clusters. The <code>minSize</code> can be used to exclude these very small clusters from the <code>statSum</code> statistical summary, because they are most likely noisy data that need to be checked in a quality control step. The analysis may be then repeated.
<code>alpha</code>	confidence level: the default is $\alpha = 0.05$ for 95% confidence level.
<code>verbose</code>	logical to print processing information if <code>verbose = TRUE</code> .
<code>plot</code>	logical to call the plotting method if <code>plot = TRUE</code> .
<code>colPalette</code>	a color palette or a list of colors such as that generated by <code>rainbow</code> , <code>heat.colors</code> , <code>topo.colors</code> , <code>terrain.colors</code> or similar functions.
<code>pch</code>	Either an integer specifying a symbol or a single character to be used as the default in plotting points. See <code>points</code> for possible values.
<code>cex</code>	A numerical value giving the amount by which plotting symbols should be magnified relative to the default <code>t = 1</code> .

Details

The `validClimR` function is used for validation of a dendrogram tree produced by `HiClimR`, by computing detailed statistical information for each cluster about cluster means, sizes, intra- and inter-cluster correlations, and overall summary. It requires the preprocessed data matrix and the tree from `HiClimR` function as inputs. An optional parameter can be used to validate clustering for a selected number of clusters `k`. If `k = NULL`, the default which supports only the regional linkage method, objective cutting of the tree to find the optimal number of clusters will be applied based on a user specified significance level (`alpha` parameter). In regional linkage method, noisy spatial elements are isolated in very small-size clusters or individuals since they do not correlate well with any other elements. They can be excluded from the validation indices (`interCor`, `intraCor`, `diffCor`, and `statSum`), based on `minSize` minimum cluster size. The excluded clusters are identified in the output of `validClimR` in `clustFlag`, which takes a value of 1 for selected clusters or 0 for excluded clusters. The sum of `clustFlag` elements represents the selected number clusters. This should be followed by a quality control step before repeating the analysis.

Value

An object of class `HiClimR` which produces indices for validating the tree produced by the clustering process. The object is a list with the following components:

<code>cutLevel</code>	the minimum significant correlation used for objective tree cut together with the corresponding confidence level.
<code>clustMean</code>	the cluster means which are the region's mean timeseries for all selected regions.
<code>clustSize</code>	cluster sizes for all selected regions.
<code>clustFlag</code>	a flag 0 or 1 to indicate the cluster used in <code>statSum</code> validation indices (<code>interCor</code> , <code>intraCor</code> , <code>diffCor</code> , and <code>statSum</code>), based on <code>minSize</code> minimum cluster size. If <code>clustFlag = 0</code> , the cluster has been excluded because its size is less than the <code>minSize</code> minimum cluster size. The sum of <code>clustFlag</code> elements represents the selected number clusters.
<code>interCor</code>	inter-cluster correlations for all selected regions. It is the inter-cluster correlations between cluster means. The maximum inter-cluster correlation is a measure for separation or contiguity, and it is used for objective tree cut (to find the "optimal" number of clusters).
<code>intraCor</code>	intra-cluster correlations for all selected regions. It is the intra-cluster correlations between the mean of each cluster and its members. The average intra-cluster correlation is a weighted average for all clusters, and it is a measure for homogeneity.
<code>diffCor</code>	difference between intra-cluster correlation and maximum inter-cluster correlation for all selected regions.
<code>statSum</code>	overall statistical summary for <code>interCluster</code> , <code>intraCor</code> , and <code>diffCor</code> .
<code>region</code>	ordered regions vector of size <code>N</code> number of spatial elements for the selected number of clusters, after excluding the small clusters defined by <code>minSize</code> argument.
<code>regionID</code>	ordered regions ID vector of length equals the selected number of clusters, after excluding the small clusters defined by <code>minSize</code> argument. It helps in mapping ordered regions and their actual names before ordering. Only the <code>region</code> component uses ordered ID, while other components use the names used during the clustering process.

Author(s)

Hamada S. Badr <badr@jhu.edu>, Benjamin F. Zaitchik <zaitchik@jhu.edu>, and Amin K. Dezfuli <amin.dezfuli@nasa.gov>. **HiClimR** is a modification of **hclust** function, which is based on Fortran code contributed to STATLIB by F. Murtagh.

References

Hamada S. Badr, Zaitchik, B. F. and Dezfuli, A. K. (2015): A Tool for Hierarchical Climate Regionalization, *Earth Science Informatics*, **8**(4), 949-958, doi: [10.1007/s1214501502217](https://doi.org/10.1007/s1214501502217).

Hamada S. Badr, Zaitchik, B. F. and Dezfuli, A. K. (2014): Hierarchical Climate Regionalization, *Comprehensive R Archive Network (CRAN)*, <https://cran.r-project.org/package=HiClimR>.

See Also

[HiClimR](#), [HiClimR2nc](#), [validClimR](#), [geogMask](#), [coarseR](#), [fastCor](#), [grid2D](#) and [minSigCor](#).

Examples

```
require(HiClimR)

## Load test case data
x <- TestCase$x

## Generate longitude and latitude mesh vectors
xGrid <- grid2D(lon = unique(TestCase$lon), lat = unique(TestCase$lat))
lon <- c(xGrid$lon)
lat <- c(xGrid$lat)

## Hierarchical Climate Regionalization
y <- HiClimR(x, lon = lon, lat = lat, lonStep = 1, latStep = 1, geogMask = FALSE,
  continent = "Africa", meanThresh = 10, varThresh = 0, detrend = TRUE,
  standardize = TRUE, nPC = NULL, method = "ward", hybrid = FALSE,
  kH = NULL, members = NULL, validClimR = TRUE, k = 12, minSize = 1,
  alpha = 0.01, plot = TRUE, colPalette = NULL, hang = -1, labels = FALSE)

## Validation of Hierarchical Climate Regionalization
z <- validClimR(y, k = 12, minSize = 1, alpha = 0.01, plot = TRUE)

## Use a specified number of clusters (k = 12)
z <- validClimR(y, k = 12, minSize = 1, alpha = 0.01, plot = TRUE)

## Apply minimum cluster size (minSize = 25)
z <- validClimR(y, k = 12, minSize = 25, alpha = 0.01, plot = TRUE)

## The optimal number of clusters, including small clusters
k <- length(z$clustFlag)

## The selected number of clusters, after excluding small clusters (if minSize > 1)
ks <- sum(z$clustFlag)
```

WorldMask

World Mask for Geographic Masking in HiClimR

Description

This data is used for geographic masking by `geogMask` function in HiClimR package.

Usage

```
data(WorldMask)
```

Format

WorldMask is a list with two components: `info` and `mask`. `info` is an (284 rows by 10 columns) matrix. The rows are for areas or countries while the columns are for codes required by `geogMask`. `mask` is an (3601 rows by 1801 columns) matrix with integer values from 1 to 284 for the areas defined in `info`.

Details

This data is used internally by `geogMask` function for geographic masking in HiClimR package. The user is advised to refer to the function manual for more details. The world mask is available in 0.1 degree (10 km) resolution. The `info` data provides information for continents, regions, and country codes).

Source

The data are based on the Humanitarian Information Unit (HIU) Large Scale International Boundaries (LSIB) dataset.

References

Hamada S. Badr, Zaitchik, B. F. and Dezfuli, A. K. (2015): A Tool for Hierarchical Climate Regionalization, *Earth Science Informatics*, **8**(4), 949-958, doi: [10.1007/s1214501502217](https://doi.org/10.1007/s1214501502217).

Hamada S. Badr, Zaitchik, B. F. and Dezfuli, A. K. (2014): Hierarchical Climate Regionalization, *Comprehensive R Archive Network (CRAN)*, <https://cran.r-project.org/package=HiClimR>.

LSIB Data: <https://hiu.state.gov/data/>.

Examples

```
require(HiClimR)
```

```
geogMask()
```

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