# Package 'bbsAssistant'

July 5, 2019

July 3, 2019
<b>Fitle</b> Helper Functions for Obtaining Data and Information from the North American Breeding Bird Survey (BBS)
Version 0.0.0.9000
Description Provides various helper functions for getting Breeding Bird Survey (BBS) data and metadata from the North American Breeding Bird Survey website.  This package will allow users to facilitate the ease with which BBS data are accessed and imported into R.
<b>Depends</b> R (>= $3.5.3$ )
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export\_bbsFeathers Save BBS dataframe as a feather file on disk.

## **Description**

Save the BBS data frame as a feather file to disk for easier import.

## Usage

```
export_bbsFeathers(dataIn, newDir = NULL, filename)
```

#### **Arguments**

dataIn Data to be saved to local disk (e.g., /bbsData/). This data frame is a product

of merging route information and raw count data. Data frame should have at least the columns dplyr::select(year, countrynum, statenum, route, bcr, latitude,

longitude, aou, stoptotal).

newDir Where to save the BBS data as feathers. If not specified will default to /bbsData/.

filename Name of the new filename (e.g., 'arizona.zip'). This should be the same filename

used to import the .zip from FTP (i.e, it should end with ".zip") This function

will replace .zip with .feather

# Description

Downloads all species-regions annual abundances indices from various .csv files on the BBS analysis results page. See https://www.mbr-pwrc.usgs.gov/bbs/BBSAnalysis\_Results\_Metadata\_1966\_2017\_v3.xml for metadata for analysis results and trend estiamtes.

#### Usage

```
get_analysis_results(url = NULL, analysis = "trend.ests")
```

# **Arguments**

url Web link to the .csv file for annual abundance indices. Default = NULL.

analysis Default = "trend.ests". Must be one of :c("annual.inds.2016", "trend.ests", "core.trend.revised.2017",

"core.2017", "expanded.2017", "core.twedt.revised.2017")

get\_bbsData 3

get_bbsData Download USGS Breeding Bird Survey data to file and import into the environment.	get_bbsData	
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# Description

This function downloads a select subset or all the BBS data from the FTP server using the .txt file downloaded from get\_regions() or using specified regions (e.g. Florida, Florida.zip). The data are saved to a temporary folder. This function was adapted from **oharar/rBBS** package. Note: this function requires an internet connection. If the bbs data are on file, please specify in the parameter file.

# Usage

```
get_bbsData(file,
    dir = "ftp://ftpext.usgs.gov/pub/er/md/laurel/BBS/DataFiles/States/",
    year = NULL, aou = NULL, countrynum = NULL, states = NULL)
```

# **Arguments**

file	One file name including the .zip extension ("stateX.zip"). Preferably download a single state at a time, otherwise run time will take >1 minutes.
dir	URL to the StatesFiles.
year	Vector of years. Default = NULL (all years).
aou	Vector of AOU numeric codes. Default = NULL (all species). (For species list visit the BBS FTP site.
countrynum	Vector of country ID #'s. Default = NULL (all countryNums).
states	Vector of state names Default = NULL (all states).

# Value

If download successful, a dataframe with the results.

# **Examples**

```
# download all species and years from Nebraska.
## Not run:
NE <- getDataBBS(file = "Nebrask.zip")
## End(Not run)</pre>
```

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```
get_credibility_trends
```

Get data credibility and trends from the BBS analysis results

# Description

The BBS provides regional and state-wide species population trend estimates. They also include a 'data credibility' rating (low, medium, or high deficiencies) for each species-region combination. This function uses web scraping to capture population trends and their credibility ratings.

# Usage

```
get_credibility_trends(url = "https://www.mbr-pwrc.usgs.gov/cgi-bin/atlasa15.pl?FLA&2&15&csrfmi
```

# **Arguments**

url

Web address of the region or state for which species' population trends and credibility ratings are to be scraped. Default example is 'Kansas, USA'.

get\_regions

Get the list of available data regions from the BBS website

# **Description**

Read in list of regions (State/Prov/TerrName), from RegionCodes.txt, and then extract list of where the 10-stop data is kept

# Usage

```
get_regions(source = "ftp://ftpext.usgs.gov/pub/er/md/laurel/BBS/DataFiles/",
ZipFiles = TRUE, bbsDir = NULL)
```

## **Arguments**

source The website or location of the BBS data files.

bbsDir The location of the local folder containing the bbs raw data. If this folder DNE,

a new folder will be created in the working directory.

get\_routeInfo 5

get_routeInfo Download route information from USGS server	get_routeInfo	Download route information from USGS server	
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# Description

This function downloads information about route location from the BBS FTP server. This function was adapted from **oharar/rBBS** package.

# Usage

```
get_routeInfo(routesFile = "routes.zip",
  routesDir = "ftp://ftpext.usgs.gov/pub/er/md/laurel/BBS/DataFiles/",
  RouteTypeID = 1, Stratum = NULL, BCR = NULL)
```

# **Arguments**

RouteTypeID	One or more numbers indicating route substrate (1=roadside;2=water;3=offroad; Default = 1, roadside only).
Stratum	A vector of BBS physiographic stratum codes by which to filter the routes.
BCR	A vector of Bird Conservation Region codes where by which to filter the routes.
routeDir	Location of the routes.zip folder Should be in DatFiles folder (default).
routeFile	Name of the route information file. Usually "routes.zip".

# Value

If download successful, a dataframe with the results.

# **Examples**

```
# download BBS route data.
## Not run:
RouteInfo <- get_routeInfo()
## End(Not run)</pre>
```

get_speciesList	Download species names
gct_spccrcstrst	Downioud species names

# Description

Read in list of species names, from SpeciesList.txt, and then extract list of where the data is kept

# Usage

```
get_speciesList(Dir = "ftp://ftpext.usgs.gov/pub/er/md/laurel/BBS/DataFiles/")
```

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## **Arguments**

Dir

ftp URL for directory with data files

## Value

A dataframe

get\_unzip

Downloads and unzips a zip archive to file.

## **Description**

Downloads and unzips a zip archive to file.

# Usage

```
get_unzip(ZipName, FileName)
```

# **Arguments**

ZipName file to download FileName file to unzip to

## **Details**

Used internally. If ZipName begins with 'http' or 'ftp', then download and unzip to Filename and return as a dataframe. Otherwise, unzip Zipname and return as a data.frame.

# Value

A dataframe

import\_bbsData

A wrapper function for downloading raw data, filtering by species names/AOU codes, and merging with route information.

## **Description**

This function will import BBS data which has already been downloaded to file. Some parts of this function were borrowed from **oharar/rBBS**.

## Usage

```
import_bbsData(file, dir, year = NULL, aou = NULL, countrynum = NULL,
  states = NULL, routesFile = "routes.zip",
  routesDir = "ftp://ftpext.usgs.gov/pub/er/md/laurel/BBS/DataFiles/",
  RouteTypeID = 1, Stratum = NULL, BCR = NULL)
```

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## **Arguments**

file The name of the zipfile to be downloaded from dir

dir http:// pr ftp:/ link to bbs data location (here: state files)

year Vector of years. Default = NULL (all years).

aou Vector of AOU #s Default = NULL (all species).

countrynum Vector of country ID #'s. Default = NULL (all countryNums).

states Vector of state names Default = NULL (all states).

RouteTypeID One or more numbers indicating route substrate (1=roadside;2=water;3=off-

road; Default = 1, roadside only).

Stratum A vector of BBS physiographic stratum codes by which to filter the routes.

BCR A vector of Bird Conservation Region codes where by which to filter the routes.

zipFileNames One or more file names ("state.zip"), char vector. Preferably download a single

state at a time, otherwise run time will take >1 minutes.

countryName Vector of country names. Default = NULL (all countryNames).

regions Vector of regionCodes. Default = NULL (all regions).

routeDir Location of the routes.zip folder Should be in DatFiles folder (default).

routeFile Name of the route information file. Usually "routes.zip".

# Description

Load the BBS data feathers into R.

# Usage

import\_bbsFeathers(newDir, filename)

# **Arguments**

newDir Where the BBS feathers are saved.

filename Name of the feather filename (e.g., 'arizona.zip' or 'arizona'). This function

will replace .zip with .feather when necessary.

8 subset\_speciesList

subset_speciesList	Subset the BBS data by species, functional traits, and/or body mass.  Default is null.

# Description

This function allows the user to subset the BBS data by specific species using AOU numbers.

# Usage

```
subset_speciesList(myData, subset.by = NULL, aou.ind = NULL,
  order.ind = NULL, fam.ind = NULL)
```

# Arguments

myData	A data frame containing the BBS data. Must contain the column "aou".
aou.ind	Numeric or vector of numeric values of the AOU codes. These are the species you want to REMOVE from analysis.
order.ind	Character or vector of characters of taxonomic orders to remove
fam.ind	Character or vector of characters of taxonomic family to remove

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