

dietr Tutorial

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1: Introduction

This is a tutorial for using the R package **dietr**. **dietr** uses diet or food item data to calculate trophic levels following the procedures described in **TrophLab** (Pauly et al., 2000), which currently is only available as a Microsoft Access database program. Our implementation is very easy to use and extremely fast as users can specify all their data as dataframes. It also differs from TrophLab in that users can specify a taxonomic hierarchy and measure trophic levels from their data at various levels (e.x. individual, population, species, genus, etc.). **dietr** also works well with FishBase (Froese & Pauly, 2018) data and can use diet and food item data obtained in R using the rfishbase package (Boettiger et al., 2012). In addition to estimating trophic levels, **dietr** can also estimate various electivity indices used in diet studies.

For this tutorial we will refer to diet data as quantitative stomach content data where the proportion of prey items are known (e.x. percent volume or weight of prey items, be cautious using percent frequency as various literature suggest it may be misleading). In this case, trophic level is simply estimated by adding one to the sum of trophic levels of the prey items consumed weighted by their contribution to the diet. The trophic level of consumer i ($Troph_i$) is defined by the equation below:

$$Troph_i = 1 + \sum_{j=1}^G DC_{ij} \times Troph_j$$

Here $Troph_j$ is the trophic level of the j th prey item consumed in the diet of i , DC_{ij} is the fraction of prey item j in the diet of i , and G is the number of prey species in the diet.

As estimates of prey trophic levels may not be exact, the standard error around the estimate of the trophic level can be calculated with equation 2. TrophLab gets around this by assigning a standard error for each trophic level and using the below equation to measure the standard error (which they also refer to as the omnivory index). Here, the standard error of the focal species i , $s.e._i$, is calculated as the square root of the sum of the product of the standard errors of the prey items $s.e._j$ to the second weighted by their respective contribution in the diet, DC_{ij} over 100.

$$s.e._i = \sqrt{\frac{\sum_{j=1}^G DC_{ij} * s.e._j^2}{100}}$$

For estimating trophic levels from food items found in the diet that don't have proportions, a random sampling and ranking of the food items is used to get an estimate of the trophic level. The simulated proportion of prey items for calculating trophic level, P is calculated using the following equation:

$$\log_{10}P = 2 - 1.9\log_{10}R - 0.16\log_{10}G$$

Here, R is the rank of the food item and G is the number of food items, up to 10. If more than 10 food items are listed, then a subsample of ten are randomly selected. The trophic level is then calculated using the following equation:

$$Troph = \sum (P_i * Troph_i) / \sum P_i$$

Here, P_i is the simulated proportion of the prey item i in the diet and $Troph_i$ is the trophic level of prey item i . This procedure is repeated n times and the mean of these n simulations is taken as the trophic level. In cases where only a single prey item is in the diet, the procedure is much simpler and the estimated trophic level is simply calculated by adding 1 to the trophic level of the single prey item.

The estimate of the standard error around the trophic level from food item data is defined below. Here, the standard error is the square root of the sum of the product of the standard error of a prey item squared and the contribution of the prey item minus 1 divided by the sum of the contribution of the prey items, P , minus the total number of prey items, G . For food item data, the random sampling routine and calculation to estimate trophic level and standard error is repeated 100 times, with the final trophic level and standard error being the mean of these 100 calculations.

$$s.e_i = \sqrt{\frac{(s.e_1)^2 * (P_1 - 1) + (s.e_2)^2 * (P_2 - 1) \dots (s.e_G)^2 * (P_G - 1)}{\sum P - G}}$$

In this tutorial we will cover the basics of how to use **dietr** to measure trophic levels. We will first discuss how to read in data from FishBase using **rfishbase** and how to pass that data into **dietr**. We will then show how one can input their own raw data and use the package to estimate trophic levels.

2: Installation

2.1: Installation From CRAN

In order to install the stable CRAN version of the **dietr** package:

```
install.packages("dietr")
```

2.2: Installation of Development Version From GitHub

While we recommend use of the stable CRAN version of this package, we recommend using the package **devtools** to temporarily install the development version of the package from GitHub if for any reason you wish to use it:

#1. Install 'devtools' if you do not already have it installed:

```
install.packages("devtools")
```

#2. Load the 'devtools' package and temporarily install the development version of
#'dietr' from GitHub:

```
library(devtools)
```

```
dev_mode(on=T)
```

```
install_github("sborstein/dietr") # install the package from GitHub
```

```
library(dietr)# load the package
```

#3. Leave developers mode after using the development version of 'dietr' so it will not
#remain on your system permanently.

```
dev_mode(on=F)
```

3: Using dietr

To load **dietr** and all of its functions/data:

```
library(dietr)
```

3.0: Basic data necessary to run dietr

To run **dietr** you will need the following data as inputs. First is diet or food item data, which our functions call in **DietTroph** and **FoodTroph** as **DietItems** and **FoodItems** respectively. These should be organized from most inclusive to least inclusive from left to right and have column names.

The second is a dataframe of trophic levels of the prey, or as we name them in the functions, **PreyValues**. We include a few different datasets with prey values users can use, though you can also supply your own. **FishBasePreyVals** are the values FishBase/TrophLab use to calculate trophic levels. **CortesPreyVals** are standardized diet prey values for sharks from Cortes, 1999. Both of these can be loaded into R from **dietr** using the **data()** function:

```
data(FishBasePreyVals)#Load the Fishbase trophic levels of prey items
#Standardized trophic levels of prey items for elasmobranchs
data(CortesPreyVals)#Load the Cortes (1992)
```

The last data object we need is a data frame we call **Taxonomy**. Columns of this dataframe should move from least inclusive to most inclusive from left to right. This data is used to assign individuals to groups for measuring hierarchical trophic levels (ex. trophic levels for an individuals, populations, species, etc.). This can be as simple as just a single column data frame if you only want to measure trophic levels for each individual and not at a higher level.

3.1: Using dietr to calculate trophic levels from FishBase diet data

Our first example will use FishBase diet data. Unfortunately, this was easier to do with previous versions of **rfishbase** where you could specify the taxa you wanted data for from the **diet** function and this function had to be re-written to work with **rfishbase** 3.0. In version 3.0 and above, the **rfishbase** separated the function into two different functions that are not intuitively useful as one returns the actual diet data (**diet_items**) and the other returns the metadata for the diet records (**diet**). Unfortunately, one must manually merge these records together to see what species belong to which diet data and users can no longer specify a single taxon to retrieve data for. **dietr**'s **ConvertFishBaseDiet** function does the merging of these two data sets for you and allows you to exclude life history stages while converting the data into a format that can be used to calculate trophic levels in **dietr**. However, as **rfishbase** no longer allows you to specify species to return, the function downloads all available diet data, thus, users will need to filter out their focal taxa.

The function has only one argument, **ExcludeStage**, in which users specify if they want to exclude a stage (ex. larvae) or not (in which case **ExcludeStage** = **NULL**). It returns a list of two data frames. The first, named **DietItems** contains the diet items while the second, **Taxonomy**, contains the taxonomy for calculating hierarchical trophic levels. Below, we will get FishBase diet for use with **dietr** while removing records from immature specimens.

```
# Convert Fishbase Diet Data and exclude juvenile and larval records
my.diets <- ConvertFishbaseDiet(ExcludeStage=c("recruits/juv.", "larvae"))
```

We can see that **my.diets** object we created is a list containing two data frames. The first one, called **DietItems**, contains the information about diet composition, with **FoodI**, **FoodII**, **FoodIII**, **Stage** and **DietPercent** being the fields with descriptors for the various diets. In this case, each diet item has its own row in the data frame. The first column, **Individual**, contains the fish base diet reference number unique to that study. By including this, one can have multiple studies of the same species and then pool the data using the **Taxonomy**, which is the second data frame in our list. The **Species** column in the **DietItems** data frame contains the species name. For example, from the **my.diets\$DietItems** object we created above, we can see that the first record in our dataset *Merlangius merlangus* with the diet record number of 4 ate eight different items, mollusks, squids/cuttlefish, bony fish, n.a./other annelids, and a variety of other invertebrates. If we look at **my.diets\$Taxonomy** we will see it is a data frame of two columns. For FishBase data, our function returns each individual diet study for a species in the column **Individual**. The next column has the respective species name. This information will then be used to calculate the trophic levels for each individual diet record and then for all records belonging to that species.

We can now measure the trophic level from diet items using the function `DietTroph`. Here, we will specify our `DietItems` and `Taxonomy` using the `cleaned.diets` object we generated above. We will also specify the `PreyValues` as being the included `FishBasePreyVals` that are part of the package. As the columns for the `PreyVals` and `DietItems` use a classification of “FoodI”, “FoodII”, “FoodIII”, “Stage”, we will specify these as a vector for the `PreyClass` argument. For this example, let's just focus on a single species with multiple diet records, *Epinephelus itajara*, which has a total of three records.

```
#Remove Data for Epinephelus itajara
my.diets$DietItems <- my.diets$DietItems[my.diets$DietItems$Species == "Epinephelus itajara",]
my.diets$Taxonomy <- my.diets$Taxonomy[my.diets$Taxonomy$Species == "Epinephelus itajara",]
```

We can now calculate the trophic levels. We will use the trophic levels of prey from `FishBase`, which is a data object that can be loaded named `FishBasePreyVals`. We will specify our diet data and taxonomy with the parameter `DietItems` and `Taxonomy` respectively. We also need to specify how the prey is classified. This is meant to be flexible and tells `dietr` how to relate the values in `DietItems` to the values in `PreyValues`. If we look at the columns of both, we can see they use a classification scheme of FoodI, FoodII, "FoodIIIandStage", so we will input that information in `PreyClass`. We will show the flexibility of this parameter a little later in this vignette in another example. The final parameter, `SumCheck`, checks that the diet data do in fact sum to 100 as would be expected if they are percent composition and will recalculate if they are not. I strongly recommend this always be set to `TRUE`:

```
data(FishBasePreyVals)#load the FishBase prey values that are part of the dietr package
#Calculate trophic level with DietTroph function
my.TL<-DietTroph(DietItems = my.diets$DietItems,PreyValues = FishBasePreyVals,
Taxonomy = my.diets$Taxonomy, PreyClass=c("FoodI","FoodII","FoodIII","Stage"), SumCheck = TRUE)
```

We can see that the `my.TL` object we just created contains a list of length two, each with a data frame, with the names of these data frames matching the `Taxonomy` we provided. We can see that the first data frame in this list, named `Individual` contains the trophic levels for each individual study that we input, while the data frame named `Species` provides the mean trophic level and SE and the number of studies across all individuals for each species.

3.2: Using `dietr` to calculate trophic levels from `FishBase` food item data

Our second example will estimate trophic level from food item data. The process is extremely similar to the above. For this example, let's get some data from `rfishbase` using the `fooditems` function for the same two species we did above.

```
#Get some food item data from rfishbase
my.food<-rfishbase::fooditems(c("Lutjanus apodus","Epinephelus itajara"))
```

In order to use this in `dietr`, we will need to convert it using the function `ConvertFishbaseFood`. This function is basically identical to the `ConvertFishbaseDiet` function we used above, except our data frame is from the `fooditems` function. We will then use `FoodTroph` function to calculate the trophic level. It is important to note that as this method randomly samples food items and ranks them for calculating the trophic level, that each time you run the function you will get a slightly different result, though they should largely be close in value.

```
#Convert FishBase food item data to a format usable for FoodTroph
converted.foods<-ConvertFishbaseFood(my.food)
#Calculate trophic level from food items
my.TL<-FoodTroph(FoodItems = converted.foods$FoodItems,PreyValues = FishBasePreyVals, Taxonomy =
converted.foods$Taxonomy,PreyClass=c("FoodI","FoodII","FoodIII","Stage"), Iter = 100,
SE.Type = "TrophLab")
```

3.3: Using diatr to calculate trophic levels from your own data

`diatr` was written with flexibility in mind so it is easy for users to use their own data or data from non-FishBase sources to calculate trophic levels. In this section, we will discuss how we can use this flexibility to customize trophic level calculations for a dataset. For this example, we will use data from Magalhaes et al., 2015, which analyzed the stomach contents of the cichlid *Herichthys minckleyi* volumetrically.

First we can load the data. We will mostly be focusing on the last ten columns, which contain the volumetric proportions of the prey items as well as the first three columns which contain information on the individual fish, the lake it was caught, and the year in which it was caught.

```
data(Herichthys)
```

Note, this is the raw data from their paper and contains other data on morphology and genotypes. As such, not all individuals have diet data associated with them, so we will want to remove them.

```
#Subset out individuals with diet data
HMdat<-Herichthys[Herichthys$total==100,]
```

For this tutorial lets measure four hierarchical trophic levels. Specifically, we will measure trophic levels at the individual, lake by year, lake (across all years), and for the species inclusive of all *Herichthys minckleyi* individuals. To do this, we will need to generate a four column data frame to input as our `Taxonomy` parameter. We can do this by doing the following.

```
#Make a data frame of the individuals, lake by year, and lake.
HMTax<-cbind.data.frame(HMdat$individual,paste(HMdat$lake,HMdat$year),HMdat$lake)
#Name the data frame
colnames(HMTax)<-c("Individual","Lake x Year","Lake (all years)")
#To calculate trophic level for the entire species, add a vecotr to the data frame of the species name
HMTax$species<-"Herichthys minckleyi"
```

Next, we need to organize the data for input with `diatr`. First, lets grab out the data.

```
HMdat<-HMdat[,c("individual","X.Gastrop","X.Insect","X.Fish","X.Zoopl","X.plants","X.algae",
"X.detritus")]
```

Remember that `diatr` requires each row to be a unique diet item per individual and that the first column contains the individual's name. We will need to format our data to work. Currently, we can see that for each individual, there are columns for the prey items, so we will need to take the data in these columns and make each prey item a row.

```
#Repeat the individual name the number of unique prey types (6)
Inds<-rep(x = HMdat$individual, times=6)[order(rep(x = HMdat$individual, times=6))]
#Repeat the number of food typed the length of the number of individuals
FoodTypes<-rep(x = colnames(HMdat[2:7]),times=length(unique(HMTax$Individual)))
#Make a data frame, the length of the individuals with three columns
HM.mat<-as.data.frame(matrix(nrow = length(Inds),ncol = 3))
#Name these columns
colnames(HM.mat)<-c("Individual","FoodItem","Percent")
#Populate the dataframes first column with the individual and the second column with the prey type
HM.mat$Individual<-Inds
HM.mat$FoodItem<-FoodTypes
#Run this for loop to find the diet data based on the individual and then match the diet percentage
#based on the name of the prey type
for(i in 1:nrow(HMdat)){
  rows2match<-which(HM.mat$Individual==HMdat$individual[i])
  HM.mat$Percent[rows2match]<-as.vector(as.numeric(HMdat[i,2:7]))
}
#Remove prey that do not contribute to diets
```

```
HM.mat<-HM.mat[!HM.mat$Percent==0,]
```

We can see that our data frame `HM.mat` is of three columns. The first row has our individual names, the second, the prey name, and the third, the dietary contribution of that prey. We now need to generate the prey values we want to use for calculating trophic levels into a format that can be used with `dietr`. For our example, we can easily do this by creating a data frame, which we will call `PreyMat`. The dimensions of this data frame will be three columns, so we can input the prey name, prey trophic level, and prey SE (if we want to have one, we can also just set them to 0, and functionally not measure it). Note for this example, the values of the prey will be equivalent to those in `FishBasePreyVals`, but as an example, I will show a simple way one could make prey values with vectors.

```
#Create a empty data frame for prey values
PreyMat<-as.data.frame(matrix(ncol = 3,nrow = 6))
#Name the columns something useful
colnames(PreyMat)<-c("FoodItem","TL","SE")
#Add in the prey name to the PreyMat
PreyMat[,1]<-unique(FoodTypes)
#Add in the trophic levels of the prey
PreyMat[,2]<-c(2.37,2.2,3.5,2.1,2,2)
#Add in the SE of the prey
PreyMat[,3]<-c(.58,.4,.8,.3,0,0)
```

We now have all our needed information to calculate trophic levels (`DietItems`, `Taxonomy`, and `PreyValues`). We can now call `dietr`'s `DietTroph` function and calculate trophic levels at the individual, lake by year, lake, and species level we defined in our `Taxonomy` data frame.

```
HM.TL<-DietTroph(DietItems = HM.mat,PreyValues = PreyMat, PreyClass = "FoodItem",Taxonomy = HMtax,
SumCheck = TRUE)
```

We can see that the object returned `HM.TL` is a list that has a length of four (one for each of our specified levels in taxonomy). Each element of the list is a data frame containing the trophic level calculations at the respective hierarchical levels.

3.4: Electivity Indices

While `dietr` can estimate trophic levels from food item and diet composition data as highlighted above, it can also measure a number of popular electivity indices used in studies of trophic ecology. The `dietr` function `Electivity` implements Ivlev's (1961), Strauss' (1979), Jacob's Q and D (1974), Chesson's (1983)(Which is similar to Manly's Alpha (1974)), and Vanderploeg & Scavia (1979) electivity indices. `Electivity` takes two data frames as input. One should contain the relative abundance of consumed prey (argument `Diet`) and the other should contain the relative abundance of available prey items in the habitat (argument `Available`). These should be formatted as such. The data frame for `Diet` should have each individual row as a diet record. The first column should contain the name of the diet record, the second, a identifier that links that record to an identifier in the first row in the data frame for `Available` (i.e. a habitat name, year, etc.). All following columns should contain a unique prey item and its relative abundance in the diet. The `Available` data frame contains similar data, however, in this data frame, the first column contains the identifier for the available prey relative abundance and all remaining columns should represent prey items available in the habitat and match those in `Diet`.

To highlight an example of how to use this function and how data should be formatted for it, we can load data from Horn, 1982. This is a dataset containing data on consumption and availability of macroalgae prey for two species of Stichaeidae in two different years. We can load it as such and see the format of the data by doing the following.

```
data(Horn1982)# load data
Horn1982$Consumed#See data for prey consumption
Horn1982$Available#See data for available prey
```

We can see here that we are identifying available prey here by year and month in the first column of `Horn1982$Available` and that all remaining columns are prey items available during those two time periods. In `Horn1982$Consumed` we can see that the first column is the name of the record and the second column matches a record in the first column of `Horn1982$Available` to link it to prey availability. We can then run one or any of the indices using the following code and arguments. We will define the consumed prey with the `Diet` argument as `Horn1982$Consumed`, `Available` prey as `Horn1982$Available`, and we will denote we want to calculate all of the indices using the `Indices` argument. For calculating Jacobs' Q, we will use the argument `LogQ` which takes \log_{10} of Q, which is recommended. We will also denote that we do not want to account for prey depletion in the calculation of Chesson's index. If set to true, Chesson's case 2 will be calculated.

```
my.indices <- Electivity(Diet = Horn1982$Consumed, Available = Horn1982$Available, Indices =
  c("ForageRatio", "Ivlev", "Strauss", "JacobsQ", "JacobsD", "Chesson", "VanderploegScavia"), LogQ = TRUE,
  Depleting = FALSE)
```

We can see that the `Electivity` function returned a list of data frames. Each data frame contains a calculated index and is named respective of the index calculated. Do not that in some cases of these calculations, certain calculations may yield `NaN` values or `Inf` values, for instance due to division by 0 or log of zero or negative number. These typically arise when a prey item is either present in the habitat but not consumed, present in the diet but not in the habitat, or where they are both absent from a record. In these cases, `NA` is returned.

4: Final Comments

Further information on the functions and their usage can be found in the helpfiles `help(package=dietr)`. For any further issues and questions send an email with subject 'dietr support' to `sam@borstein.com` or post to the issues section on GitHub(<https://github.com/sborstein/dietr/issues>).

5: References

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