## Updating and checking the taxonomy

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The following R code illustrates how to update the taxonomy used in COMADRE using the R package taxize (Chamberlain & Szöcs 2013) to query the Catalogue of Life (http://www.catalogueoflife.org) with the names used by the source publication authors in COMADRE.

The function classification in taxize allows the programmatic querying of databases including the Catalogue of Life. For example:

```
library(taxize)
classification("Daphnia magna", db = 'col')
> Retrieving data for taxon 'Daphnia magna'
> $`Daphnia magna`
>
                                                          id
             name
                      rank
> 1
         Animalia kingdom 5ede24b0534ebd5e1f552d5b9f874a6a
> 2
       Arthropoda phylum 89ac18bfcf1654a9662a600ba06bb494
> 3 Branchiopoda
                     class 1793fd966478a2d1c959221e2a711569
> 4
     Diplostraca
                     order 04df00f2834eac8dc22966b299226a13
> 5
    Not assigned
                    family
                                                    28669362
> 6
          Daphnia
                     genus 1acee7e4189fdad4c8abf00900254b85
    Ctenodaphnia subgenus
                                                    28908195
> 8 Daphnia magna
                   species e44e0a35c2cdb085859074a9fe73abdf
> attr(,"class")
> [1] "classification"
> attr(,"db")
```

One can capitalise on this for COMADRE.

First load the required package, taxize.

## library(taxize)

> [1] "col"

Now make a new vector called SpeciesBinomial by concatenate the accepted genus (GenusAccepted) and accepted species epithet (SpeciesEpithetAccepted) together. This is necessary, rather than simply using SpeciesAccepted, because SpeciesAccepted retains the infra-specific information which is not used by the following code:

```
comadre$metadata$SpeciesBinomial <-
paste(comadre$metadata$GenusAccepted, comadre$metadata$SpeciesEpithetAccepted)</pre>
```

Some species do not have an epithet (e.g. *Tribolium* sp.), for these the epithet is listed as NA. Therefore, to search the Catalogue of Life effectively, the NA needs to be removed using gsub:

Because species appear in the database numerous times it is advisable for efficiency reasons to make a unique subset of the data:

```
temp <- unique(comadre$metadata[, c("SpeciesBinomial",
"GenusAccepted", "Family", "Order", "Class", "Phylum", "Kingdom")])</pre>
```

This dataset is still quite large (333 rows), so here I will just obtain the information for the first 5 rows.

```
temp <- temp[1:5,]
```

This is accomplished using the classification function which repeatedly queries the Catalogue of Life for each entry. Note that whenever there is an uncertainty as to which species is intended, the software prompts the user to select a species from a list.

```
x <- classification(temp$SpeciesBinomial,db='col')
> Retrieving data for taxon 'Acipenser fulvescens'
> Retrieving data for taxon 'Acipenser transmontanus'
> Retrieving data for taxon 'Ambloplites rupestris'
> Retrieving data for taxon 'Ammocrypta pellucida'
> Retrieving data for taxon 'Ammocrypta pellucida'
```

The object produced by this code is a list with elements corresponding to each species. Thus x[[1]] will return the first species etc. One can also address the list by element name, which is defined by the query that was submitted e.g. x[["Acipenser fulvescens"]]

## x[[1]]

```
>
                    name
> 1
                Animalia kingdom 5ede24b0534ebd5e1f552d5b9f874a6a
> 2
                Chordata phylum 4313bc7637e1fc1feb316a4dea2b668b
                           class 655300e5b96a3dc5e76e02175c3aa191
> 3
          Actinopterygii
> 4
        Acipenseriformes
                           order 1b6074738c3c562418eee137a0369bdb
> 5
           Acipenseridae
                          family 5a7e5d2984e865f1aff640138c760d04
                           genus b506984ec1a4c2f6191b842dcbce8fa7
> 6
               Acipenser
> 7 Acipenser fulvescens species ebc713afa86464a0cfb313955a1db149
```

Thus, to acquire the taxonomic Order of the species of interest one would use:

```
x[[1]] name [x[[1]] rank == "order"]
```

```
> [1] "Acipenseriformes"
```

One can run through this list object in a loop to compare the information held in COMADRE, with that produced by the query to Catalogue of Life. For example, to check the Orders of the 5 species held in temp

```
for(i in 1:nrow(temp)){
n <- temp$SpeciesBinomial[i]
dat <- x[[n]]
CoL_Order <- dat$name[dat$rank == "order"]
if(temp$Order[i] == CoL_Order){temp$OrderOK[i] <- "OK"
} else {
   temp$OrderOK[i] <- "Differs!"}
}</pre>
```

which will add an extra column called OrderOK to the temp data frame:

## temp

```
>
            SpeciesBinomial GenusAccepted
                                                 Family
                                                                   Order
> 1
       Acipenser fulvescens
                                Acipenser Acipenseridae Acipenseriformes
> 14 Acipenser transmontanus
                                Acipenser Acipenseridae Acipenseriformes
      Ambloplites rupestris Ambloplites Centrarchidae
> 15
                                                            Perciformes
> 18
       Ammocrypta pellucida Ammocrypta
                                               Percidae
                                                            Perciformes
> 20
         Amphiprion percula
                               Amphiprion Pomacentridae
                                                            Perciformes
             Class Phylum Kingdom OrderOK
> 1 Actinopterygii Chordata Animalia
                                          OK
> 14 Actinopterygii Chordata Animalia
                                          OK
> 15 Actinopterygii Chordata Animalia
                                          OK
> 18 Actinopterygii Chordata Animalia
                                          OK
> 20 Actinopterygii Chordata Animalia
                                          OK
```

This approach can naturally be repeated for each part of the taxonomy.