# Example usage in R

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#### Introduction

This document gives a first introduction in querying the TGD database such that occurrences and phylogenetic data are properly linked despite input data sets potentially using different taxon labels and different backbone taxonomies The overarching design of the schema is such that there is a table of taxa, which uses the names accepted by either the Mammal Species of the World 3rd edition, the GBIF backbone taxonomy, or both. As such, each record in that table has at least one of the identifiers used by these resources. Subsequently, all variations on these accepted names (that is, alternative spellings, synonyms, etc.) are records in the taxonvariants table, each of which have a foreign key that links back to the accepted name version. Whenever a data set (a tree, a set of occurrences, a trait data set) is imported, the record labels that are used in that data set are looked up in the taxon variants table. Any existing exact match is re-used, and new ones are created as needed. Then, an attempt is made to link new variants to accepted taxon names using taxonomic name reconciliation (as implemented in the taxize package). The upshot of this design is that, if you want to get all occurrences (say) for a given species, you first have to look up that species in the taxa table, then get all the variations on that name in the taxonvariants table, and then all occurrences that link to any of these variants. And, if you want to combine, for example, occurrences and phylogeny, you have to look for the intersection between "taxa that have variants that have occurrences", and "taxa that have variants that have branches in the given tree".

Without further ado, here's what the steps might be. First we connect to the database:

```
library(DBI)
```

```
## Warning: package 'DBI' was built under R version 3.4.4

db_file <- "../data/sql/tgd.db"

tgd <- dbConnect(RSQLite::SQLite(), db_file)</pre>
```

Then we need to look up the ID of the mammal supertree of Bininda-Emonds et al. (2007), which is named .../data/phylogeny/Bininda-emonds\_2007\_mammals.tsv. We query as follows:

```
tree_name <- "../data/phylogeny/Bininda-emonds_2007_mammals.tsv"
query <- 'select tree_id from trees where tree_name="%s"'
tree_id <- dbGetQuery(tgd, sprintf(query, tree_name))$tree_id</pre>
```

Given this tree\_id, we are now going to look for taxa that have taxonvariants that have branches in the tree and that have taxonvariants that have occurrences. This is a fairly complex query all in all because there's a number of joins that need to be done. As follows:

```
query <- 'select distinct
taxa.taxon_name,occurrences.decimal_latitude,occurrences.decimal_longitude
from (((taxonvariants
inner join taxa on taxonvariants.taxon_id = taxa.taxon_id)
inner join occurrences on occurrences.taxonvariant_id = taxonvariants.taxonvariant_id)
inner join branches on branches.taxonvariant_id = taxonvariants.taxonvariant_id)
where branches.tree_id=12 and
taxa.taxon_level="SPECIES"
order by taxon_name'
results <- dbGetQuery(tgd, query)</pre>
```

## Building a data frame

Now that we have a great big table, we should be able to do some potentially interesting analyses. For example, does Rapoport's rule apply to our data? This is an ecogeographical rule that states that latitudinal ranges of plants and animals are generally smaller at lower latitudes than at higher latitudes. So, for this we need the absolute average latitude and the latitudinal range.

The data structure that we are going to want thus needs to have one observation for each taxon, recording 2 variables (absolute average latitude, latitudinal range, longitudinal range). Here now we populate a data frame to do this:

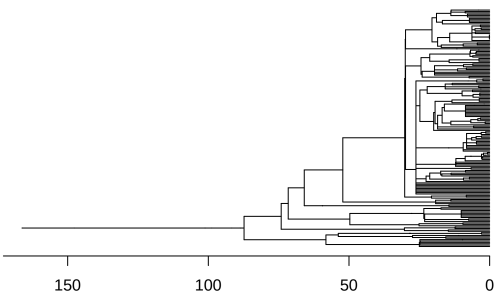
```
# we will use the distinct taxon names (these are the clean, accepted names) as row names
names <- unique(results$taxon_name)</pre>
# we start with a matrix, populate this, and then make it into a data.frame
df <- matrix(ncol=2, nrow=length(names))</pre>
# use i index for the matrix
for ( i in 1:length(names) ) {
    # average absolute latitude
    aal <- mean(abs(results[which(results$taxon name==names[[i]]), "decimal latitude"]))
    # latitudinal range
    lar <- range(results[which(results$taxon_name==names[[i]]),"decimal_latitude"])</pre>
    lar <- lar[[2]] - lar[[1]]</pre>
    # add row to matrix
    df[i,] <- c( aal, lar )</pre>
}
# make data frame, assign taxa as row names, variables as col names
df <- data.frame(df, row.names = names)</pre>
colnames(df) <- c("average absolute latitude", "latitudinal range")</pre>
# remove observations with ranges of O, i.e. single observation
row_sub = apply(df, 1, function(row) all(row !=0))
df <- df[row sub,]</pre>
names <- row.names(df)
```

### Importing phylogenies

We might now be tempted to simply do some correlation tests, but this is actually not a valid approach. Our observations are of species that are related to one another, so part of the commonality in observation values might be due to shared descent. For example, two sister species are probably near each other on the earth, and have a similar range: pseudo-replication. To account for this we need take phylogeny into account. Hence, we need to extract a tree from the database. We will do this by extracting a phylo object (as originally introduced by the ape package). As follows:

```
library(ape)
source('../R/make_phylo.R')
tree <- make.phylo(db_file, tree_id, names)
# at this scale, the tip labels will be cluttered.</pre>
```

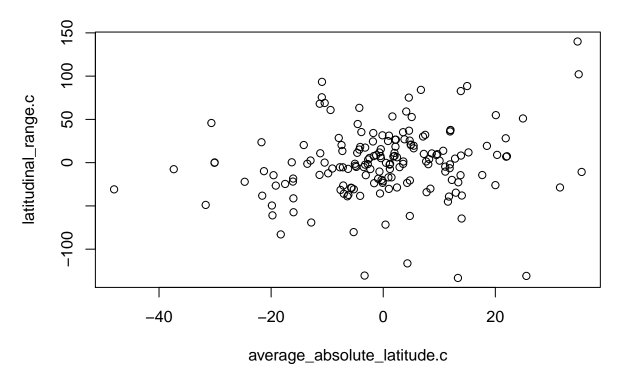
```
# let's just make sure it's an ultrametric tree.
plot(tree, show.tip.label=F)
axisPhylo(side = 1)
```



## Doing a comparative analysis

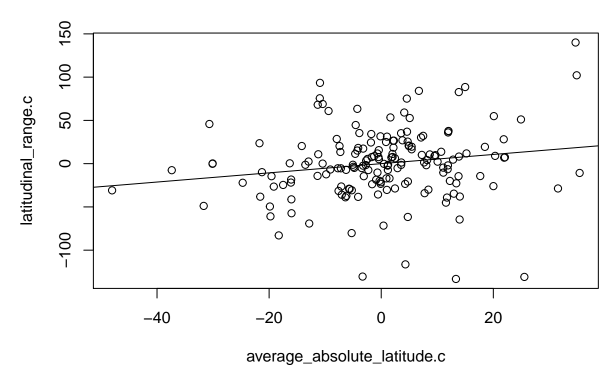
We now have everything in place to do a phylogenetic comparative analysis, e.g. pic in ape. We follow this tutorial:

```
# Because we extract a subtree from the database, it may
# have interior nodes with just one child. Here we
# collapse these.
tree <- collapse.singles(tree)</pre>
# The tree may not be fully bifurcating, althout PIC needs
# this. Hence we arbitrarily resolve it.
tree <- multi2di(tree)</pre>
# create separate vectors
average_absolute_latitude <- df$average_absolute_latitude</pre>
names(average_absolute_latitude) <- row.names(df)</pre>
latitudinal_range <- df$latitudinal_range</pre>
names(latitudinal_range) <- row.names(df)</pre>
# calculate contrasts
average absolute latitude.c <- pic(average absolute latitude, tree, scaled=F)
latitudinal_range.c <- pic(latitudinal_range, tree, scaled=F)</pre>
# produce scatterplot
plot(average_absolute_latitude.c, latitudinal_range.c)
```



The raw plot is suggestive of a positive correlation, so let's do a linear model:

```
regress <- lm(latitudinal_range.c~average_absolute_latitude.c-1)</pre>
summary.lm(regress) # p < 0.05
##
## Call:
## lm(formula = latitudinal_range.c ~ average_absolute_latitude.c -
##
##
## Residuals:
                       Median
                                    ЗQ
                                            Max
##
        Min
                  1Q
## -144.492 -22.751
                       -0.609
                                18.034
                                        121.592
##
## Coefficients:
##
                               Estimate Std. Error t value Pr(>|t|)
## average_absolute_latitude.c
                                 0.5315
                                            0.2305
                                                     2.306
                                                             0.0223 *
##
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 39.84 on 170 degrees of freedom
## Multiple R-squared: 0.03033,
                                    Adjusted R-squared: 0.02463
## F-statistic: 5.318 on 1 and 170 DF, p-value: 0.02231
# plot with regression line
plot(average_absolute_latitude.c, latitudinal_range.c)
abline(regress)
```



It appears that Rapoport's rule indeed applies (p<0.05\*)