## Appendix S7 - Extracting Migration Status, Diet Diversity and Habitat Diversity

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This appendix describes how to extract information on species' migratory status from BirdLife International as well as calculate indices of habitat diversity, diet diversity and rarity. This data can be accessed in the BBS.occurrences library using data('Bird.traits'). Note, the primary functions for extracting migratory status were adapted from code that was generously provided by Tad Dallas.

Some libraries that are not automatically included in the package will be needed for web-scraping. Check that they are installed and up-to-date

```
library(tidyverse)
library(httr)
library(rvest)
library(purrr)
library(stringr)
library(parallel)
```

Download the Birdtree.org taxonomy file, which will be necessary for searching common names when species' binomials are not recognised by BirdLife. Note, this step requires an internet connection

```
temp <- tempfile()
download.file("https://data.vertlife.org/birdtree/BLIOCPhyloMasterTax.csv",
    temp)
BT.tax <- read.csv(temp)</pre>
```

Load the binary species occurrence data that is included in the BBS.occurrences package and extract the column names

```
library(BBS.occurrences)
data("BBS.occurrences")
sp_names <- colnames(BBS.occurrences)</pre>
```

Here we perform a sanity check to ensure that parallel loading is supported on our machine. Note, the library parallel needs to be loaded for this to function properly. If parallel computation is not desired, just set n\_cores to 1

```
n_cores <- 3
if (n_cores > 1) {
    cl <- makePSOCKcluster(n_cores)</pre>
    setDefaultCluster(cl)
    test_load1 <- try(clusterEvalQ(cl, library(rvest)),</pre>
        silent = TRUE)
    if (class(test_load1) == "try-error") {
        pkgLibs <- unique(c(sub("/rvest$", "", system.file(package = "rvest"))))</pre>
         clusterExport(NULL, c("pkgLibs"), envir = environment())
         clusterEvalQ(cl, .libPaths(pkgLibs))
         test_load2 <- try(clusterEvalQ(cl, library(rvest)),</pre>
             silent = TRUE)
         if (class(test_load2) == "try-error") {
             clusterEvalQ(cl, .libPaths(as.character(.libPaths())))
             test_load3 <- try(clusterEvalQ(cl, library(rvest)),</pre>
                 silent = TRUE)
             if (class(test_load3) == "try-error") {
                 parallel_compliant <- FALSE</pre>
                 stopCluster(cl)
             } else {
                 parallel_compliant <- TRUE</pre>
         } else {
             parallel_compliant <- TRUE</pre>
    } else {
        parallel_compliant <- TRUE</pre>
} else {
    parallel_compliant <- FALSE</pre>
parallel_compliant
```

After confirming parallel capabilities, start the parallel clusters and run the function. Note, this step requires an internet connection and will take some time to process the 303 avian species in the dataset

```
# Export necessary data and variables to each
clusterExport(NULL, c("BT.tax", "sp_names"), envir = environment())
clusterEvalQ(cl, library(rvest))
clusterEvalQ(cl, library(stringr))
clusterEvalQ(cl, library(purrr))
clusterEvalQ(c1, library(xm12))
search_migstatus <- pbapply::pblapply(seq_along(sp_names),</pre>
    function(x) {
        session <- rvest::html_session("http://datazone.birdlife.org/species/search")</pre>
        Sys.sleep(4)
        form <- html_form(session)[[3]]</pre>
        filledform <- set_values(form, kw = sp_names[x])
        session <- submit_form(session, filledform)$url</pre>
        landing_pg <- xml2::read_html(session)</pre>
        sp_link <- rvest::html_nodes(landing_pg, css = "tr") %>%
            rvest::html_attrs() %>% purrr::flatten_chr() %>%
            purrr::keep(~grepl("rowClick", .x)) %>%
            stringr::str_replace("rowClick\\(\\'",
                 "") %>% stringr::str_replace("\\'\)",
        if (identical(sp_link, character(0))) {
            session <- rvest::html_session("http://datazone.birdlife.org/species/search")</pre>
            Sys.sleep(4)
            comm_name <- as.character(BT.tax$English[which(BT.tax$TipLabel ==</pre>
                 paste(sp_names[x]))])
            form <- rvest::html_form(session)[[3]]</pre>
            filledform <- rvest::set_values(form, kw = comm_name)
            session <- rvest::submit_form(session,</pre>
                 filledform) $url
            landing_pg <- xml2::read_html(session)</pre>
            sp_link <- rvest::html_nodes(landing_pg,</pre>
                 css = "tr") %>% rvest::html_attrs() %>%
                 purrr::flatten_chr() %>% purrr::keep(~grepl("rowClick",
                 .x)) %>% stringr::str_replace("rowClick\\(\\'",
                 "") %>% stringr::str_replace("\\'\)",
```

```
# If still not found, return NA
             if (identical(sp_link, character(0))) {
                 output <- data.frame(species = sp_names[x],</pre>
                   Migrate.status = "NA")
             } else {
                 details_link <- paste0("http://datazone.birdlife.org/species/factsheet/",</pre>
                   sp_link[1], "/details/")
                 migrate_status <- xml2::read_html(details_link) %>%
                   rvest::html_nodes(css = "tr:nth-child(1) td:nth-child(2)") %>%
                   rvest::html_text()
                 output <- data.frame(species = sp_names[x],</pre>
                   Migrate.status = migrate_status)
        } else {
             details_link <- paste0("http://datazone.birdlife.org/species/factsheet/",</pre>
                 sp_link[1], "/details/")
            migrate_status <- xml2::read_html(details_link) %>%
                 rvest::html_nodes(css = "tr:nth-child(1) td:nth-child(2)") %>%
                 rvest::html text()
             output <- data.frame(species = sp_names[x],</pre>
                 Migrate.status = migrate_status)
        output
    }, cl = cl)
Bird.mig.status <- do.call(rbind, search_migstatus)</pre>
Bird.mig.status$Migrate.status <- as.character(Bird.mig.status$Migrate.status)</pre>
```

Next, we calculate Shannon Diversity indices to represent species' diet and habitat breadths. This function makes use of the data available in the EltonTraits database, which is the same that we used in Appendix\_S4

```
temp <- tempfile()
download.file("https://ndownloader.figshare.com/files/5631081",
          temp)
Sp_traits <- read.table(temp, header = TRUE, fill = TRUE,
          quote = "\"", stringsAsFactors = FALSE, sep = "\t")
unlink(temp)</pre>
```

Calculate Shannon indices of diet and habitat diversity for each species

Calculate an index of rarity for each species, defined by it's scaled occurrence total across all observations, where larger values indicate a species is more rare

Add the rarity and diversity metrics to the migration dataset

```
Bird.traits = Bird.mig.status %>% dplyr::left_join(Rarity) %>%
    dplyr::left_join(Sp_traits) %>% dplyr::select(species,
    dplyr::everything())
```

Finally, we extract information on species' average clutch size from the Avian Life History Traits Dataset. We then join this variable to our remaining trait variables and save in the analysis folder. This dataframe can be accessed in the BBS.occurrences library using data('Bird.traits').