Recreate UK Biodiversity Trends Paper

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This vignette uses the UKBiodiversity R package to recreate the figures generated in the paper "Complexity of biodiversity change revealed through long-term trends of invertebrates, bryophytes and lichens" by Outhwaite et al.

Set up R environment

First you will need to download the dataset used. These data are freely available from the EIDC repository and are described in full in the accompanying data paper: Outhwaite, C. L., Powney, G. D., August, T. A., Chandler, R. E., Rorke, S., Pescott, O. L., ... Isaac, N. J. B. (in press). Annual estimates of occupancy for bryophytes, lichens and invertebrates in the UK (1970 – 2015). *Scientific Data*.

To download the required datasets you will need to register with the website first, then you can follow the instructions to download the data. The complete dataset zip file should be downloaded.

In the repository you will find the following items:

- POSTERIOR_SAMPLES: zip file containing 1 file per species of the posterior samples of average occupancy.
- SUMMARY TABLES: zip file containing 1 file per species of occupancy model output summary tables.
- Dataset_Information: csv file containing input dataset metadata.
- Species_Names: csv file containing information on species names.
- Species_Trends: csv file containing all estimates of species trends as annual percentage growth rates.

For this analysis you will need to unzip the POSTERIOR_SAMPLES folder.

Set your working directory so that the POSTERIOR_SAMPLES folder is a sub-folder within it, you can do this using the function setwd().

You will also need to download the R package called UKBiodiversity from Github. How to do this is detailed below.

First, let's take a look at the POSTERIOR SAMPLES data files.

```
# Where are the posterior sample files?
# These should be in the named subfolder of your working directory.
datadir <- paste0(getwd(), "/POSTERIOR SAMPLES")</pre>
# List the files
files <- list.files(datadir)</pre>
# How many files are there?
length(files) # There should be 5293, one for each species.
## [1] 5293
# Take a look at the file list
head(files)
## [1] "AnnualSpeciesOccupancy_PosteriorSamples_Ants_Formica aquilonia.csv"
## [2] "AnnualSpeciesOccupancy_PosteriorSamples_Ants_Formica cunicularia.csv"
## [3] "AnnualSpeciesOccupancy PosteriorSamples Ants Formica fusca.csv"
## [4] "AnnualSpeciesOccupancy PosteriorSamples Ants Formica lemani.csv"
## [5] "AnnualSpeciesOccupancy_PosteriorSamples_Ants_Formica lugubris.csv"
## [6] "AnnualSpeciesOccupancy_PosteriorSamples_Ants_Formica picea.csv"
# Download the UKBiodiversity R package from GitHub,
# to do this you will first need the devtools package
library(devtools)
# Install the R package from GitHUb
install_github("CharlieOuthwaite/UKBiodiversity", quiet = TRUE)
# Load the package
library(UKBiodiversity)
```

This process should have downloaded the required packages for this analysis.

Generation of group level posterior sample datasets

For some of the analyses we will need a matrix containing all the posterior samples of the species within each of the major groups and for each of the taxonomic level groups. In the repository, the posterior samples for each species are saved as individual csv files, one for each species. Use the combine_posteriors function to make the required combinations of the posteriors by major group and taxa. The function will save the outputs as .rdata files in a sub-folder of the directory that you specify, within a group specific sub-folder. Use the group_level argument to select either "major_group" or "taxa" level datasets.

Note that these functions can take a long time to run (estimated run time below) and may have high memory requirements.

```
# Run the combine_posteriors function providing a data directory and and output directory.
# select the group level you wish to generate datasets for, either "major_group" or "taxa".
# If status is TRUE, progress will be printed within the console.
## First for major groups.
# Runtime of this function: 67 mins (when status = FALSE)
```

```
combine_posteriors(datadir = paste0(getwd(), "/POSTERIOR_SAMPLES"),
                   outdir = getwd(),
                   group_level = "major_group",
                   status = FALSE)
# Take a look at the outputs in the function generated subdirectory
files <- list.files(paste0(getwd(), "/MajorGroups"), pattern = "posterior_samples")</pre>
head(files)
## [1] "ALL posterior samples national.rdata"
## [2] "FRESHWATER SPECIES posterior samples national.rdata"
## [3] "LOWER_PLANTS_posterior_samples_national.rdata"
## [4] "TERRESTRIAL INSECTS posterior samples national.rdata"
## [5] "TERRESTRIAL_NONINSECT_INVERTS_posterior_samples_national.rdata"
# A file has been created for each of the major groups and for all species.
## Then for taxa.
# Runtime for this function: ~ 20 minutes
combine_posteriors(datadir = paste0(getwd(), "/POSTERIOR_SAMPLES"),
                   outdir = getwd(),
                   group_level = "taxa",
                   status = FALSE)
# Take a look at the outputs in the function generated subdirectory
files <- list.files(paste0(getwd(), "/Taxa"), pattern = "posterior_samples")
head(files)
## [1] "Ants posterior samples national.rdata"
## [2] "AquaticBugs_posterior_samples_national.rdata"
## [3] "Bees_posterior_samples_national.rdata"
## [4] "Bryophytes_posterior_samples_national.rdata"
## [5] "Caddisflies posterior samples national.rdata"
## [6] "Carabids posterior samples national.rdata"
# A file has been created for each taxonomic group
```

These group level files are required for the following functions. Files must be maintained within the folder they are placed by the function.

For the analyses carried out here, rove beetles are excluded since they do not have data from 1970 to 1979. Therefore, there are 5,214 species included in the analyses carried out here. 79 Rove Beetle species are removed.

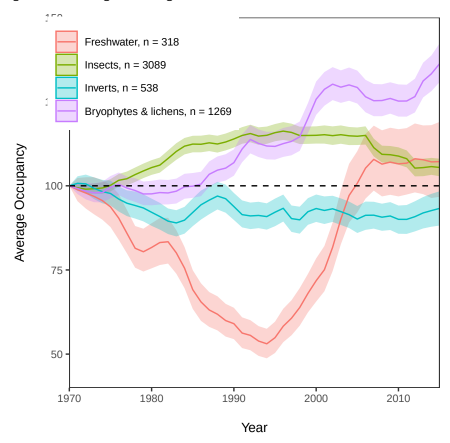
Recreating Figure 1

Figure 1 within the paper presents indicators of average occupancy over time for the four major groups.

Use the generate_fig1 function to determine the group level average change associated indicator values and the plot of Figure 1. The generate_fig1 function will save csv files of the indicator posterior values, the average annual indicator estimates and associated credible interval values, and a pdf file of the plot within a sub-directory of your specified folder called "geomeans".

This function will only work if outputs have already been produced from the "major_group" specification of the combine_posteriors function. These should be in a sub-folder called "MajorGroups" in the outdir that you specified within that function.

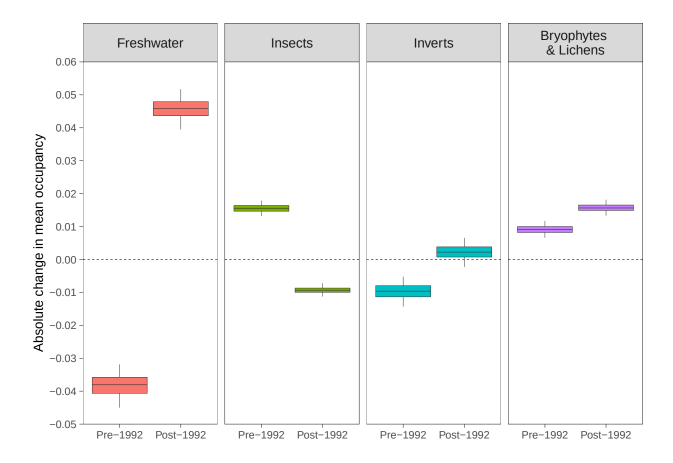
[1] "Warning: overwriting existing files"



Recreating Figure 2

Figure 2 presents box plots of absolute change in average occupancy across 2 halves of the time period assessed for each major group. It shows estimates for 1970 to 1992 and for 1993 to 2015.

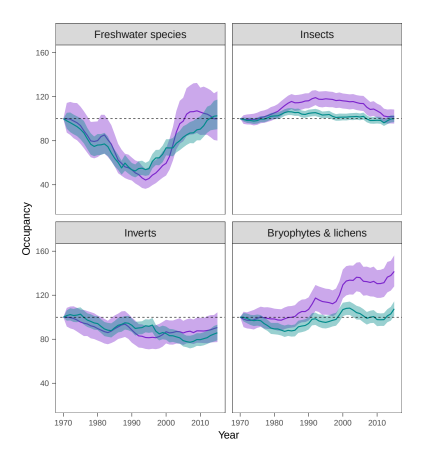
Again, this function uses the major group level posteriors of average occupancy across each group that were generated and saved within the <code>generate_fig1</code> function. These will have been saved within a sub-folder within the <code>/MajorGroups/geomeans</code>. You will need to direct the function to this folder.



Recreating Figure 3

Figure 3 presents the changes in the quantiles of occupancy over time to assess how rarity and commonness has changed over time. This function estimates the quantile data from the combined posterior samples generated by the combine_posteriors function, saves this within a sub-folder called "quantiles" as csv files and generates the figure.

[1] "Warning: overwriting existing files"

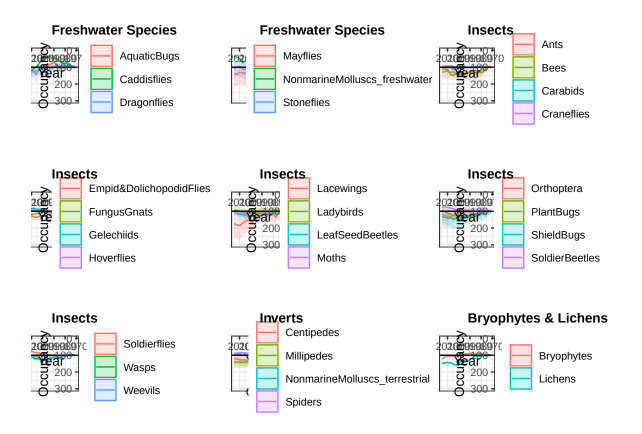


Recreating Figure 4

Figure 4 is a representation of Figure 1, except with a line for each taxonomic group. It uses the taxa level estimates generated from the combine_posteriors function to calculate the indicator posteriors and the average and 95% CI indicator values. The combined samples should be in a sub-folder called "Taxa" which was generated by the combine posteriors function. The outputs are the saved in a sub-folder called "Taxa/geomeans". These are then used to recreate Figure 4 which is saved as a PDF file within that same sub-folder.

So that the trends are more visible, the Freshwater and Insect plots have been split into multiple panels.

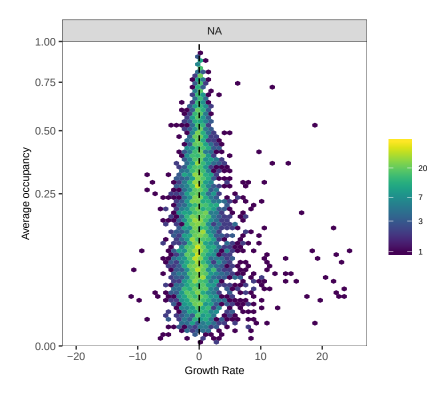
[1] "Warning: overwriting existing files"



Due to the size of this figure it is not clearly visible on this A4 page. See the paper, or create your own for a clearer view!

Recreating Figure 5

Figure 5 presents the average occupancy for each species plotted against its average annual growth rate. Average occupancy across the time period is estimated within the function, growth rates are supplied within the "Species Trends.csv" file downloaded from the repository.



Calculating group and taxa level trends

Within the text are associated group level trends and an average trend estimate for all species since 1970.

Trends can be calculated ones the associated posterior values have been generated via the generate_fi1 and generate_fig4 functions for major groups and taxa respectively. Use the group_trends function to calculate the trends relative to 1970 and the associated credible intervals. The credible interval value can be specified within the function, but defaults to 95%.

```
# Use the group_trends function to estimate overall and major group level trends
# and associated credible intervals.
# Runtime of this function: 0.17 seconds
# datadir should point to the main directory in which the "MajorGroups" and "Taxa" subdirectories
#can be found created within the generate_fig1 function.
trends <- group_trends(datadir = getwd(),</pre>
                       interval = 95)
trends
##
                                                     LCI
                                                              UCI
                               Group Mean_change
## 1
                                 ALL
                                          10.924
                                                   8.785
                                                           12.948
```

```
## 2
                 FRESHWATER_SPECIES
                                            7.189
                                                    -3.268
                                                            18.985
## 3
                        LOWER PLANTS
                                           36.269
                                                    30.791
                                                            42.298
                 TERRESTRIAL_INSECTS
## 4
                                            5.456
                                                    2.849
                                                             8.061
## 5
      TERRESTRIAL_NONINSECT_INVERTS
                                           -6.713 -11.804
                                                            -1.628
## 6
                                           31.329
                                                   -6.188
                                                            78.270
                                Ants
```

```
## 7
                         AquaticBugs
                                           18.804
                                                   -6.582
                                                            50.517
## 8
                                            9.402
                                                   -0.883
                                Bees
                                                            21.401
## 9
                          Bryophytes
                                           23.626
                                                    16.939
                                                            30.408
## 10
                         Caddisflies
                                           37.871
                                                    19.702
                                                            58.376
## 11
                            Carabids
                                            0.861
                                                    -9.072
                                                            11.818
                                           21.585
                                                   -5.976
## 12
                          Centipedes
                                                            58.529
## 13
                          Craneflies
                                           -3.963 -11.578
                                                             4.319
## 14
                         Dragonflies
                                           16.952 -15.204
                                                            51.311
## 15
            Empid&DolichopodidFlies
                                           19.772
                                                     7.611
                                                            33.650
##
  16
                         FungusGnats
                                           18.637
                                                     9.239
                                                            28.790
##
  17
                          Gelechiids
                                          -24.428 -35.265 -11.612
                                           -5.175 -14.521
##
  18
                          Hoverflies
                                                             4.182
##
  19
                                           73.504
                                                  10.549 160.431
                           Lacewings
## 20
                           Ladybirds
                                           -0.474 - 29.814
                                                            28.984
## 21
                     LeafSeedBeetles
                                            2.506 -12.262
                                                            18.149
## 22
                             Lichens
                                           47.548
                                                   38.768
                                                            57.119
## 23
                            Mayflies
                                           40.707 -12.892 116.382
##
  24
                          Millipedes
                                           39.799
                                                     7.842
                                                            76.020
## 25
                                                     3.696
                               Moths
                                            8.713
                                                            13.886
##
  26
       NonmarineMolluscs freshwater
                                          -64.156 -70.630 -57.504
##
  27
      NonmarineMolluscs_terrestrial
                                           -8.121 -17.314
                                                             2.709
## 28
                                           20.742 -19.594
                          Orthoptera
                                                            70.178
## 29
                                           -1.736 -10.830
                           PlantBugs
                                                             8.932
## 30
                                                   -7.242
                          ShieldBugs
                                           31.504
                                                            83.588
                                                            -6.172
## 31
                      SoldierBeetles
                                          -26.115 -43.157
##
  32
                        Soldierflies
                                          -32.205 -40.620 -23.178
## 33
                             Spiders
                                          -10.194 -15.984
                                                            -3.855
##
  34
                          Stoneflies
                                            9.051 -34.277
                                                            72.529
## 35
                                            4.512
                                                   -6.847
                               Wasps
                                                            16.901
## 36
                             Weevils
                                           19.885 11.747
                                                            28,702
```

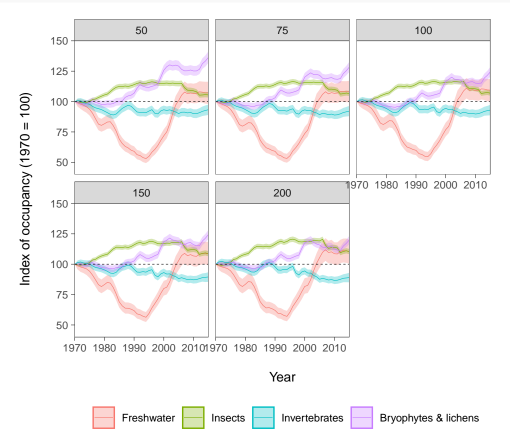
Recreating Supplementary Figure 1

Supplementary Figure 1 presents the occupancy plots for each taxonomic grouping separately within a PDF document. These figures more clearly show the responses of individual groups which may otherwise be difficult to interpret from Figure 4 within the main text.

The PDF version of this plot will be saved in the postdir/geomeans.

Recreating Supplementary Figure 2

Supplementary Figure 2 is a representation of Figure 1 in the main text except varying thresholds have been used to determine which species are included. These thresholds detail the number of records a species must have as a minimum before they are included within the generation of the indicators. The number of records within the raw data is specified within the "Species_Trends.csv" file that was downloaded from the repository. These values are used to subset the species within each group according to thresholds of number of records: 50, 75, 100, 150 or 200 records. In the main text, all species are used which have a minimum of 50 records.



You have now recreated the analyses carried out in the paper by Outhwaite et al using the original data files!