

Forest Modeling Exercise

Process-based and empirical modeling exercises

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1 Overview

This document contains the instructions and solutions of the modeling group assignments. First, you will find the section for the process model approach (Section 2) and after that the section for the empirical modelling (Section 3). This document is very long because it describes all the exercises and solutions step by step.

You will work per groups in the four groups established at the beginning of the week. Each of the four groups will be divided internally in four subgroups (A-D). Each subgroup (A-D) inside each group (1-4) will address one question that will have to be answered using one approach. In the table below you can find a link to the section to each question e.g. group 1 subgroup A will answer the questions "How are biodiversity indices changing in time and across the simulated scenario(s) on Plot 1?" using the approach based on the model iLand and will have the full instructions and solution in Section 2.2.6.

Please, always ask your coach if you do not know what is the section where you should be working in.

Group	Subgroup Question	Question	Approach	Go to
1	A	How are biodiversity indices changing in time and across the simulated scenario(s) on Plot 1?	iLand	Section 2.2.6
1	В	How the species distribution and total living biomass C content changing in time on Plot 1?	iLand	Section 2.2.8

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Group	Subgroup Question	Question	Approach	Go to
1	C	Does a more diverse forest in structure and composition have more Bryophites species?	GLM	Section 3.4.4
1	D	Is the number of Bryophites species affected by forest management type and the forest structural diversity?	GLM	Section 3.4.5
2	A	How are biodiversity indices changing in time and across the simulated scenario(s) on Plot 2?	iLand	Section 2.2.8
2	В	How the species distribution and total living biomass C content changing in time on Plot 2?	iLand	Section 2.2.9
2	C	Does a more diverse forest in structure and composition have more bird species?	GLM	Section 3.4.6

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Group	Subgroup Question	Question	Approach	Go to
2	D	Is the number of bird species	GLM	Section 3.4.7
		affected by		
		forest		
		management		
		type and the		
		forest		
		structural		
		diversity?		
3	A	How are	iLand	Section 2.2.10
		biodiversity		
		indices		
		changing in		
		time and across the		
		simulated		
		scenario(s) on		
		Plot 3?		
3	В	How the	iLand	Section 2.2.11
		species		
		distribution		
		and total living		
		biomass C		
		content		
		changing in		
3	$^{\mathrm{C}}$	time on Plot 3? Is the presence	BRT	Section 3.4.8
3	C	of the Great	DUI	Section 5.4.6
		spotted		
		woodpecker		
		affected by		
		forest density?		
3	D	Is the presence	BRT	Section 3.4.9
		of the Great		
		spotted		
		woodpecker		
		affected by		
		forest		
		diversity?		

1 Overview

	Subgroup			
Group	Question	Question	Approach	Go to
4	A	How are biodiversity indices changing in time and across the simulated scenario(s) on Plot 4?	iLand	Section 2.2.12
4	В	How the species distribution and total living biomass C content changing in time on Plot 4?	iLand	Section 2.2.13
4	С	Is the presence of the Eurasian treecreeper affected by forest density?	BRT	Section 3.4.10
4	D	Is the presence of the Eurasian treecreeper affected by forest management?	BRT	Section 3.4.11

1.1 Data and model

We provide all data, model and scripts for the excercises. However if you will use the model iLand for your research in the future please get in contact with the model developers at the Technical University Münich: Werner Rammer (werner.rammer@tum.de) and Rupert Seidl (rupert.seidl@tum.de). Regarding the biodiversity data used in the empirical modeling processed in the empirical modeling part, in case of further usage please contact Jeňýk Hofmeister at the Czech University of Life Sciences Prague (jenyk.hofmeister@email.cz).

2.1 iLand model

2.1.1 General

iLand is an ecosystem model that simulates forest landscape dynamics, including growth and regeneration, disturbance dynamics, and management in a spatially explicit manner. The main entity in the model is a tree, for which the demographic processes are simulated. Processes at the stand and landscape scale constrain the dynamics of individual trees and thus allow for the scaling of tree-scale processes to large areas. The model explicitly simulates tree competition for resources such as light, water, and nutrients. A light use efficiency approach is used to simulate the production physiology. Carbon starvation is used as a process oriented indicator of tree stress, which can result from competition for resources as well as suboptimal environmental conditions for tree growth (e.g., drought).

iLand's mechanistic representation of forest disturbances and vegetation dynamics, as well as the climatic sensitivity of these processes, makes it well suited for the research of disturbance dynamics under climate change. Additionally, flexible implementation of management operations, which include planting after harvests or natural disturbances, thinning, harvesting, and post-disturbance salvaging, allows for testing the effects of various disturbance management strategies.

2.1.2 Input and outputs of iLand

iLand needs information in specific formats about environment (mainly climate and soil) and trees. Daily climate data is needed (https://iland-model.org/ClimateData) for minimum, maximum temperature, precipitation, vapor pressure deficit and radiation. Soil depth and soil texture (sand, silt, clay %) information are needed. The soil is represented as a one-layer bucket, no depth-dependent information is needed. The model needs species-specific parameters (https://iland-model.org/species+parameter) that are describing the behavior of each tree species (growth, mortality, competition etc.). Currently there are 31 species parametrized for iLand, and the species specific parameters

are stored in a database that is an input for the model. The parameters were previously calibrated for European conditions (species_param_europe.sqlite). iLand uses short codes for species derived from the latin names (e.g. Picea Abies - piab, Fagus Sylvatica - fasy, ..) Information on the initial forest is also required to start a simulation.

As iLand is a landscape scale model, both climate, soil and tree information need to be set spatially. For larger areas environmental maps can be set using a 100x100m resolution grid, a so called resource grid with resource units, and tree information can be set by maximum 10x10m resolution grid on a stand grid using stand IDs. There are more options how to initialize a landscape in iLand depending on our spatial scales and available data (https://iland-model.org/initialization).

In our case now we will use iLand in a stand-scale mode (called torus), where we simulate only one 100x100m pixel, 1 ha forest area. We will use single-tree initialization method, where we set the x-y coordinates of the trees inside the 100x100m area, tree species, dbh and height of each tree. From inventory we have data for the a circle with 13.82m radius. We generated trees for the 1ha area based on the plot-data. See details later.

The output of the model is an sqlite database file. The output produced by the model is highly versatile and needs to be set in advance of the simulation which output tables we would like to get.

2.1.3 How to work with the model

Simulations are driven by one xml file, the project file. After starting iland.exe, we should call the project file, load the initial forest and then run the simulation for the required length of period. iLand comes with a graphical interface where you can visualize single trees and many other things: https://iland-model.org/iLand+viewer

2.2 Excercises

Your group will work with the plot which you measured and worked with in the previous days (Plot1-Plot4). The 2 process based subgroups will do the same modeling experiment, just addressing different questions based on the results (see the 2 questions below).

You will simulate the growth and development using the process-based model, iLand. The task is to simulate your study plot for 100 years under reference conditions and one/two simplified scenarios that you could define upon your ideas. You can define scenarios assuming different temperature, precipitation and CO2 concentration levels (see more details how to do it later below). The questions that you should address forming 2 sub-groups are the following:

- A) How biodiversity indices are changing in time and across the simulated scenario(s)? (Calculate indices)
- B) How the species distribution and total living biomass C content changing in time? Compare 0 year and 100 year status in the reference case and in the case of your scenario(s)!

2.2.1 Getting started

Navigate to this folder and download the materials for the process-based modeling exercise: https://drive.google.com/drive/folders/1o2_K3ZN-SnPR44UzZoHj9lFZQu-vU3wg?usp=drive link

Go to the *iLand_simulations* folder. Here you will find several folders that are the part of the model, others are containing input data for the model and one folder is dedicated for the outputs. Additionally, you will find *iland.exe* and *.xml* files. The *.xml* file is called *project file*. This file is driving the settings of simulations, such as input/output file names, output variables and many other settings. This file is what you need to run in the model.

The xml file structure follows a tree structure, e.g:

The iLand project file consists of five main sections:

- system: settings like file path, database locations, or logging
- model:main settings of the models: extent (world), site specific setting, used climate, model initialization and management
- modules: settings related to the disturbance modules of iLand
- output: definition of which outputs should be created by the model
- user: this section can be used for user-defined settings

More details about the project file structure and meaning of each record can be found here: https://iland-model.org/project+file

We prepared one project file for each group (for each simulation plot). You can right away run it, and then copy-paste and modify it to specify your own scenarios for alternative simulations. You can open a project file in any text editor, but we recommend to use Notepad++ for this (https://notepad-plus-plus.org/downloads/).

2.2.2 Explore input files

The following input files are needed and prepared for you to run the model: This is all prepared, YOU DO NOT HAVE TO CHANGE ANYTHING:

• Species parameter file: database/species_param_europe.sqlite Containing the parameters for each species that are driving growth, mortality, establishment and other processes. https://iland-model.org/species+parameter

Set in project file:

• Climate file: database/E-OBSv27_Roznik_46.05_14.45_1961-1990.sqlite Daily climate data for the area based on the E-OBS dataset. https://iland-model.org/ClimateData https://surfobs.climate.copernicus.eu/dataaccess/access_eobs.php#datafiles We are using the same climate input for all plots. The E-OBS data is representative for a 0.1x0.1 degree resolution gridbox

Set in project file:

```
<database>
     <in>species_param_europe.sqlite</in>
     <out>Output_plot1_4deg_30percdrier.sqlite</out>
     <climate>E-OBSv27_Roznik_46.05_14.45_1961-1990.sqlite</climate> <!--- HERE
</database>
```

• Environmental file: gis/Environment.txt

The structure of this file is very flexible, more or less columns can be set. If we would have a landscape with larger area with different environments, we could set

more lines into this file. For now we have only one simulation pixel with 1ha area. Here we set which climate table to use from the climate file (there could be more tables, but now we have only one), what is the soil texture, soil depth. The column names are referring to specific lines in the project file.

For example: if we set here "model.site.pctSand", the model will use this sand % value instead of the value that we have in the project file under:

Set in project file:

```
<environmentGrid>gis/environment_grid.asc/environmentGrid>
<environmentFile>gis/Environment.txt/environmentFile> <!---- HERE</pre>
```

• Environment grid: gis/environment_grid.asc This would specify the map of different environments. Now we have only 1 pixel, with the id of 1. In the previous file we had also id column, this specifys we want to use that environment for our pixel.

Set in project file:

```
<environmentGrid>gis/environment_grid.asc/environmentGrid> <!---- HERE
<environmentFile>gis/Environment.txt/environmentFile>
```

• Tree initialization file: init/Tree_init_plot_1.txt This is the list of the tree with x,y coordinates, species, dbh, height and optionally with age. (age=0 means, there is no data on age, model will generate) There are other options to set initial trees/landscape (e.g. using distributions, number of trees in a given dbh range, or using outputs of another simulation) More details: https://iland-model.org/initialize+trees

Here we initialize trees taller than 4 meters. Trees smaller than 4m are handled as sapling cohorts in separated input file. Now we do not put there saplings and we do not have a sapling file.

We assume now no-management scenario for all simulations. Generally iLand has a very sophisticated so called "agent based management engine" functionality for large land-scapes, but also simple management activities are possible for stand-scale simulations. However, for the sake of simplicity in these examples we do not simulate any management activities. This need to keep in mind during the interpretation of the results.

NOTE: in the graphical interface you also wont see trees smaller than 4m, because they are not simulated as individuals rather in cohorts. But as regeneration will work, after a while as they grow above 4m, they become individual trees and we can see them also visually and they have own properties (dbh, height, age,...https://iland-model.org/tree+variables). More details on saplings: https://iland-model.org/sapling+growth+and+competition

Setting the output tables: In the output section you can set which tables you want to enable. Here you can find more info on the structure and variables of each table: https://iland-model.org/Outputs

To answer the questions the most straight forward way is to look the landscape output and use the $total_carbon_kg$ column (total carbon in living biomass (aboveground compartments and roots) of all living trees (including regeneration layer) (kg/ha)). This output table aggregates on the level of landscape x species. Values are always aggregated per hectare. The output is created after the growth of the year. We pre-set the project file to have this output. Additionally we will work with the tree output, where individual trees are recorded with position.

```
<output>
<tree>
<enabled>true</enabled>
</tree>
...
```

```
<landscape>
<enabled>true</enabled>
</landscape>
</output>
```

2.2.3 Run the model

- Double click on iland.exe and wait until the graphical interface opens.
- Load the required project file on the left-hand-side under *iLand project file* by browsing on your computer after clicking on the folder icon next to the box.
- After you selected a project file, click on the "Create Model" on the top-left part of the window with an icon of a Globe.
- Wait until the simulation area shown in the middle.
- Try out some visualizations on the right hand-side "Visualization options". e.g individual trees + color by species. Here you can explore the initial stage of the 1 ha simulation area.
- To run the model click on Run Model icon on the top and give 100 years to simulate.
- On the bottom of the window you can follow how fast the model is running and see if it is finished.
- Stage of the last year remains in the simulation area, and you can go to the folder output to check your output file.
- When you want to run another simulation, first click on "Destroy", then start the process again by "Creating Model" for the same project file or loading new project file.

2.2.4 Working with the output:

The output database has a .sqlite extension. You can explore outputs:

- Open and browse it in the *DB Browser* software that was on the software list. (https://sqlitebrowser.org/)
- Load the file into R, where you can make additional data analyses needed to answer the questions in the exercise.

```
file <-
    here::here("model/iLand_simulations/output/Output_plot1.sqlite")

sqlite.driver <- RSQLite::dbDriver("SQLite")
db1 <-</pre>
```

```
RSQLite::dbConnect(sqlite.driver, dbname = file) # connect to the file
tabs <- RSQLite::dbListTables(db1) # explore the tables in the file
print(tabs)

landscape <- RSQLite::dbReadTable(db1, "landscape")

RSQLite::dbDisconnect(db1)

summary(landscape)
head(landscape)</pre>
```

There are different output tables in iLand and all have specific structure, column and spatial representation of the records. The landscape output gives information on the whole landscape, but here we will work with one 1 ha pixel that is our "whole" landscape for now. Here species specific information is also available for each year. More about the meaning of the columns in the landscape output you can find here: https://iland-model.org/Outputs#Landscape_aggregates_per_species Not always all outputs are produced, in the project file we can set which output tables we want to enable, and those will be created.

2.2.5 Run alternative scenarios

The prepared simulation is using the 30years climate of 1961-1990, and repeating during the simulation. (The model copy paste the 30year time series after each other to fill up the simulation period) For CO2 concentration 400ppm is given. Your task is to run 2 extra alternative scenarios where you modify the output data. For example assuming 800ppm, and 4degree warming, or keep CO2 on 400ppm and decrease precipitation by 20%.

All of these modification can be done in the project file directly, you do not need to modify input data files!

For this, go to the climate section and modify the values there, and save as the project file as your alternative scenario.

```
<randomSamplingList></randomSamplingList>
</climate>
```

Note that the same CO2 concentration will be used during the whole simulation period (there is a possibility to give annually changing CO2 concentration using an external file, but for now we won't use it for simplicity). The temperature shift is applied by the model on each day's temperature values. The precipitation shift is a scaler that the model applies on each day's precipitation amount. (0.8 gives 20% decrease)

DO NOT FORGET TO CHANGE THE OUTPUT FILE NAME AS WELL for your alternative scenario, otherwise it will overwrite the previous output. You can do it in this section in the beginning of the project file:

```
<database>
<in>species_param_europe.sqlite</in>
<out>Output_plot1_4deg_20percdrier.sqlite</out> <!----- HERE
<climate>E-OBSv27_Roznik_46.05_14.45_1961-1990.sqlite</climate>
</database>
```

Running the alternative scenario follows the same steps as before.

2.2.6 Question A - Group 1

• How are biodiversity indices changing in time and across the simulated scenario(s) on Plot 1?

Run the model as it is described in the previous chapters to have at least 2 simulations completed: one with reference conditions and one scenario with some changes in the environment. Check if you have the output files in the output folder: <code>iLand_simulations/output/</code>. If you manage to simulate 2 scenarios, you can expand the code to have file3, and 3rd from each variable, or make the comparison in 2 step always comparing one scenario to the reference conditions at the time.

Open the *summerSchoolExerciseProcessBased.Rproj* project in R studio and start working there.

Read in the *tree* output table from two outputs you have for your plot:

```
path1 <- "model/iLand_simulations/output/Output_plot1.sqlite"
path2 <- "model/iLand_simulations/output/Output_plot1_4deg_2Opercdrier.sqlite"
file1 <- here::here(path1)
file2 <- here::here(path2)</pre>
```

```
# Give some name for the three simulations.
name1 <- "reference"</pre>
name2 <- "4deg_20percdrier"</pre>
# Read in data using the RSQLite package
sqlite.driver <- RSQLite::dbDriver("SQLite")</pre>
db1 <- RSQLite::dbConnect(sqlite.driver, dbname = file1) # connect to the file1
tables.in.the.file <- RSQLite::dbListTables(db1) # explore the tables in the file1
print(tables.in.the.file)
[1] "carbon"
                         "carbonflow"
                                              "dynamicstand"
[4] "landscape"
                         "landscape_removed" "runinfo"
[7] "stand"
                         "tree"
# We will work with "tree" table and tree-scale data:
tree1 <- RSQLite::dbReadTable(db1, "tree")</pre>
RSQLite::dbDisconnect(db1) # disconnect to the file1
# READ IN DATA FROM THE SECOND FILE: ----
db2 <- RSQLite::dbConnect(sqlite.driver, dbname = file2) # connect to the file2
tree2 <- RSQLite::dbReadTable(db2, "tree")</pre>
RSQLite::dbDisconnect(db2)
```

Merge the data from the two files together and make a column which tells which comes from which simulation, and study the table. The column "run" will support the plotting where we can use facets in ggplot.

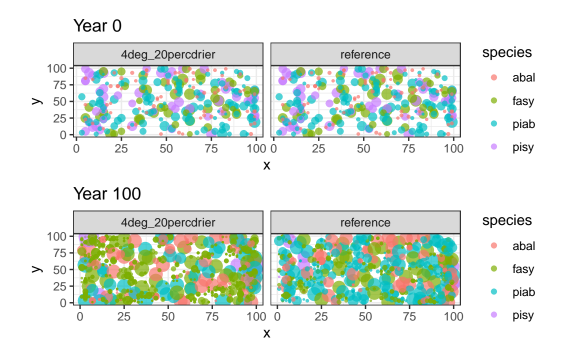
```
height basalArea volume_m3 age
 year ru rid species id x y
                                   dbh
1
    0
                fasy 1 79 51 69.45111 39.46735 0.3788334 6.474022 493
2
    0
      0
                fasy 2 69 91 68.95086 39.18307 0.3733956 6.335131 489
                fasy 3 59 83 68.42681 38.88527 0.3677414 6.191780 486
3
    0 0
           1
4
    0 0
           1
                piab 4 99 19 68.36907 45.96828 0.3671210 7.138516 492
5
                fasy 5 63 61 66.87022 38.00070 0.3512007 5.778762 475
    0
    0
                fasy 6 55 61 65.80000 37.39252 0.3400492 5.505722 467
 \tt leafArea\_m2~foliageMass~stemMass~branchMass~fineRootMass~coarseRootMass
```

```
322.7507
                 29.34097 3099.621
                                      378.6909
                                                    22.00573
                                                                   337.2626
1
2
     318.8086
                 28.98260 3049.604
                                      372.4465
                                                    21.73695
                                                                   331.6535
3
     314.7003
                 28.60912 2997.693
                                      365.9680
                                                    21.45684
                                                                   325.8348
4
                 69.61134 2192.064
     295.8482
                                      365.2582
                                                    52.20850
                                                                   526.4032
5
     302.6273
                 27.51158 2846.413
                                      347.1027
                                                    20.63368
                                                                   308.8961
                 26.76726 2744.921
                                      334.4585
                                                                   297.5477
6
     294.4398
                                                    20.07544
        lri lightResponse stressIndex reserve_kg treeFlags
                                                                   run
1 0.2646746
                                         51.34669
                        0
                                     0
                                                           0 reference
2 0.5870413
                        0
                                     0
                                         50.71954
                                                           0 reference
3 0.3931382
                                         50.06596
                        0
                                     0
                                                           0 reference
4 1.0000000
                        0
                                       121.81984
                                                           0 reference
5 0.3406834
                         0
                                         48.14526
                                     0
                                                           0 reference
6 0.4422086
                         0
                                         46.84270
                                                           0 reference
```

To explore a bit the simulation, plot the tree locations at the beginning and the end of the simulation! For plotting we use the colors for the species, and the size of a circles to show dbh differences. Here we divided dbh with the value 20 just for visualization, not to have too huge circles shading each others completely, but still see the trees.

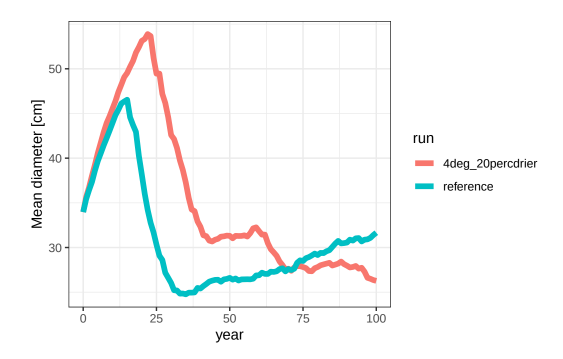
```
tree0 <- tree |> dplyr::filter(year == 0)
g1 <- ggplot2::ggplot(tree0, ggplot2::aes(x = x, y = y, color = species)) +
ggplot2::geom_point(size = tree0$dbh / 20, alpha = 0.7) +
ggplot2::ggtitle("Year 0") +
ggplot2::facet_wrap(~run) + ggplot2::theme_bw()

tree100 <- tree |> dplyr::filter(year == 100)
g2 <- ggplot2::ggplot(tree100, ggplot2::aes(x = x, y = y, color = species)) +
ggplot2::geom_point(size = tree100$dbh / 20, alpha = 0.7) +
ggplot2::ggtitle("Year 100") +
ggplot2::facet_wrap( ~run) + ggplot2::theme_bw()
gridExtra::grid.arrange(g1, g2, ncol = 1)</pre>
```



We can see that the species composition has changed and the forest become more dense. We can even identify the same trees at the same location.

Visualize some changes in time, for example the mean diameter of the trees!



We can see that there is an initial increase in mean dbh, then a drop after 20/25 years. This is due to the growing regeneration layer that is produced by the seeds of the existing trees. In the initial year we do not have regeneration layer in the simulations (it is possible to put there, but we do not have it now). And until the small trees grow up to higher than 4 m they are not appearing in the individual *tree* output as they are handled in cohorts.

To study biodiversity aspects, we can calculate biodiversity indicators. Let's use the adiv package for species diversity. First we need to change the database structure to have the number of trees for each species in different columns (pivot_wider) without any additional columns for the adiv package speciesdiv function. We work first only with tree data from one simulation (tree1). You can find the documentation of the adiv package here: "documents/adiv.pdf"

```
# A tibble: 6 x 4
   abal fasy piab pisy
  <int> <int> <int> <int>
1
     68
           50
                120
2
     68
           49
                115
                        49
3
     68
           48
                111
                        48
4
     68
                110
                        47
           48
5
     66
           48
                107
                        46
     65
           48
                105
                        45
```

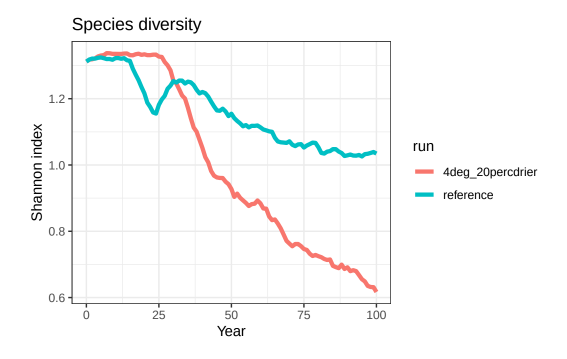
Then we apply the "speciesdiv" function and we add back the year and run columns to have the information.

```
div1 <- data.frame( adiv::speciesdiv(tlong1)) |>
  dplyr::mutate(year = unique(tree1$year), run = name1)
head(div1)
```

```
richness GiniSimpson Simpson Shannon Margalef Menhinick McIntosh year
             0.7095388 3.442800 1.312005 0.5300838 0.2361125 0.4899779
1
2
             0.7131369 3.485983 1.318098 0.5320701 0.2386200 0.4938655
                                                                            1
            0.7150017 3.508792 1.321063 0.5341147 0.2412091 0.4960613
3
             0.7150505 \ 3.509394 \ 1.320996 \ 0.5348097 \ 0.2420910 \ 0.4962264
4
                                                                            3
5
             0.7162956 3.524796 1.323410 0.5369369 0.2447960 0.4978275
                                                                            4
         4
             0.7169397 3.532816 1.324555 0.5383914 0.2466506 0.4987181
                                                                            5
6
        run
1 reference
2 reference
3 reference
4 reference
5 reference
6 reference
```

We make the same steps for the other simulation results (tree2).

We merge them all together and make a plot based on Shannon index.



We see that the values of Shannon index are first stable and then tend to decrease, while the reduction under the 4deg_20percdrier is more substantial than under the reference scenario. Shannon index is calculated as follows:

$$S = -\sum \left(p \ i \ *log(p \ i \) \right)$$

Where p i is the relative abundance of the species i, calculated as the ratio of the abundance of the species i and the abundance of all species. Shannon-index is 0 when we have only one species at the stand.

We can calculate some additional biodiversity indices targeting spatial diversity using the *treespat* package developed by Francesco Chianucci (fchianucci@gmail.com). You can find a short description of the package here: "documents/treespat_package.pdf"

You can install the package using devtools:

```
devtools::install_gitlab('fchianucci/treespat')
```

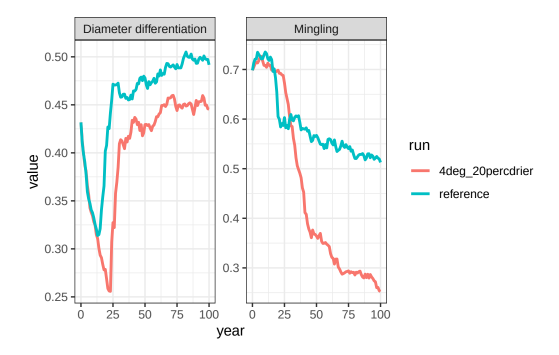
Then, we are ready to use it! Let's calculate these two indices:

- Diameter differentiation (Gadow, 1993): Spatial size inequality defined as the mean of the ratio of smaller and larger plant sizes in the nearest neighbors of a tree. The value of the index increases with increasing average size difference between neighboring trees. 0 is implying that neighboring trees have equal size.
- Mingling (Aguirre et al., 2003): One very intuitive extension of taxonomic species diversity (either richness or abundance) is considering spatial mingling, namely how plants of the same (con-specific neighbors) or different (hetero-specific neighbors) species are arranged in space. The mingling index calculates the proportion of the k nearest neighbors that do not belong to the same species as the reference tree. For example, with four neighbors, the mingling attribute can assume five values, ranging from 0 (all trees are of the same species) to 1 (all trees belong to different species).

Calculate the indices based on the reference run, then based on the scenario. For each index we add extra columns with the index name, and run name for further analyses. The max.k parameter is telling how many neighboring trees we want to account for.

```
year
           DIFF index
                                           name
1
     0 0.4317331 DIFF Diameter differentiation reference
2
     1 0.4104274 DIFF Diameter differentiation reference
     2 0.3988705 DIFF Diameter differentiation reference
3
     3 0.3903934 DIFF Diameter differentiation reference
5
    4 0.3775628 DIFF Diameter differentiation reference
     5 0.3595558 DIFF Diameter differentiation reference
ming <- data.frame(treespat::MING(tree1, .x = x, .y = y, .species = species,
        xmax = 100, ymax = 100, max.k = 4, shape='square', .groups=c('year')) |>
        dplyr::mutate(index="MING", name="Mingling", run=name1))
indices1<-rbind(diff |> dplyr::rename(value=DIFF),
ming |> dplyr::rename(value=MINGLING ) )
```

We calculated each index as separate variable and then merged them into indices1. Here I renamed the column names to have it all "value", for easier plotting later on. (each line has the information on which run and which index is the value) We do the same for the scenario simulation, we merge them together and make the plotting.



The diameter differentiation increases with the increasing average difference in diameter between neighboring trees. The value 0 indicates that neighboring trees have equal diameters. The development shows that the index values decrease during the first 20/25 years, after which it steeply increases due to the occurrence of ingrowth and then the values are stabilized. The values for the reference climate scenario are slightly higher than for the climate change scenario indicating slower growth under drier and warmer conditions and/or the differences in tree species composition (proportion of tree species). The mingling index is studying the neighboring trees regarding their species. In both development there is a decrease in the index after the occurrence of regeneration. Using the climate change scenario the index shows steeper decrease.

2.2.7 Question B - Group 1

• How are the species distribution and total living biomass C content changing in time on Plot 1? Compare 0 year and 100 year status in the reference case and in the case of your scenario(s)!

Run the model as it is described in the previous chapters to have at least 2 simulations completed: one with reference conditions and one scenario with some changes in the environment. Check if you have the output files in the output folder: *iLand_simulations/output/*. If you manage to simulate 2 scenarios, you can expand the code to have file3, and 3rd from each variable, or make the comparison in 2 step always comparing

one scenario to the reference conditions at the time. Open the *summerSchoolExercise-ProcessBased.Rproj* project in R studio and start working there.

Read in the *landscape* output table from two outputs you have for your plot:

```
path1 <- "model/iLand_simulations/output/Output_plot1.sqlite"</pre>
path2 <- "model/iLand simulations/output/Output_plot1 4deg_20percdrier.sqlite"</pre>
file1 <- here::here(path1)
file2 <- here::here(path2)
# Give some name for the three simulations.
name1 <- "reference"</pre>
name2 <- "4deg 20percdrier"</pre>
# Read in data using the RSQLite package
sqlite.driver <- RSQLite::dbDriver("SQLite")</pre>
# We will work with "landscape" table and tree-scale data:
db1 <- RSQLite::dbConnect(sqlite.driver, dbname = file1) # connect to the file1
landscape1 <- RSQLite::dbReadTable(db1,"landscape")</pre>
RSQLite::dbDisconnect(db1) # disconnect to the file1
db2 <- RSQLite::dbConnect(sqlite.driver, dbname = file2) # connect to the file2
landscape2 <- RSQLite::dbReadTable(db2,"landscape")</pre>
RSQLite::dbDisconnect(db2)
```

Merge the data from the three files together and make a column which tells which comes from which simulation, and study the table. The column "run" will support the plotting where we can use facets in ggplot.

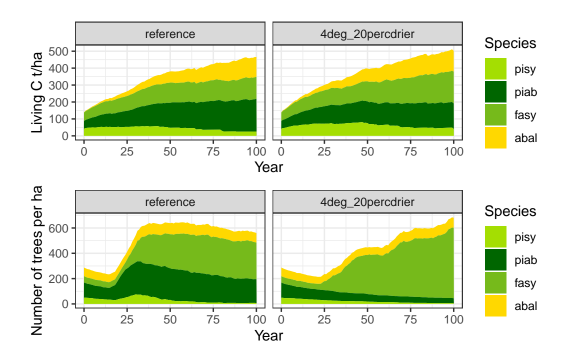
```
landscape<-rbind(landscape1 |> dplyr::mutate(run=name1) ,
  landscape2 |> dplyr::mutate(run=name2))
head(landscape)
```

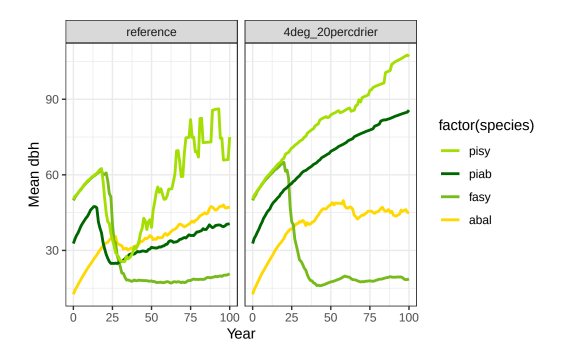
```
year area area_100m species count_ha dbh_avg_cm height_avg_m volume_m3
     0
                                                                 4.33641
1
          1
                    1
                         abal
                                    68
                                         12.55456
                                                      10.04365
2
     0
          1
                                    50
                                         49.86496
                                                      28.33702 136.89579
                    1
                         fasy
3
     0
                                                      22.00713 150.34005
         1
                   1
                         piab
                                   120
                                         32.73141
4
     0
          1
                    1
                                    49
                                         50.45049
                                                      28.66693 126.09230
                         pisy
5
     1
          1
                    1
                         abal
                                    68
                                         14.15550
                                                      11.39836
                                                                 6.12901
```

```
50.95480
                                                        28.84830 141.11347
6
     1
                          fasy
                                     49
                      gwl_m3 basal_area_m2
                                              NPP_kg NPPabove_kg
  total_carbon_kg
         2264.490
                    4.33641
                                 0.8659201
                                               0.000
                                                           0.000 0.09848037
1
                                               0.000
        49667.453 136.89579
                                                           0.000 0.94517453
2
                                10.2322163
3
        45424.641 150.34005
                                12.1157843
                                               0.000
                                                           0.000 1.22598624
4
        42580.564 126.09230
                                 9.9010286
                                               0.000
                                                           0.000 1.13237721
5
                    6.12901
                                 1.0937620 1657.542
                                                        1128.888 0.11701639
         2784.089
        52091.802 143.51049
                                10.4297469 9855.392
                                                        6472.199 0.96978453
6
  cohort_count_ha
                         run
1
                0 reference
2
                0 reference
3
                0 reference
4
                0 reference
5
               31 reference
6
               54 reference
```

We can see that the output is given for each year and each species in our 1ha area. Plot the living carbon (total_carbon_kg) in time, coloring by species. We give a unified species coloring in the beginning. Plot number of trees and mean diameter. Carbon and number of trees can be plotted in an additive way, but for mean dbh we do line plot per species.

```
cols.all=c( "rops"="#e0e0e0", "acpl"="#A9A9A9", "alin"="#696969", "alvi"="#2e2e2e",
 "bepe"="#fadfad","casa"="#7eeadf","coav"="#20c6b6","tipl"="#645394",
 "ulgl"="#311432","saca"="#D8BFD8","soar"="#DDA0DD","soau"="#BA55D3",
 "pice"="#D27D2D","pini"="#a81c07","alg1"="#2ECBE9","tico"="#128FC8",
 "potr"="#00468B", "poni"="#5BAEB7", "frex"="#fe9cb5", "cabe"="#fe6181",
 "acps"="#fe223e","lade"="#FFFE71","abal"="#FFD800", "pisy"="#A4DE02",
 "fasy"="#76BA1B", "piab"="#006600", "quro"="#FF7F00", "qupe"="#FF9900",
 "qupu"="#CC9900")
# We set here the order of the run categories for the table, to have the first
#run first and then the second. (left-right of the plots)
landscape$run <- factor(landscape$run, levels = c(name1, name2))</pre>
# Plot the living carbon content, to have tonnes/ha, we divide total_carbon_kg by 1000.
g1 <- ggplot2::ggplot(landscape, aes(year, total_carbon_kg / 1000,
                                     fill = factor(species))) +
    ggplot2::geom_area() +
    ggplot2::scale fill manual(values = cols.all, guide = guide legend(reverse = TRUE))
   ggplot2::facet_wrap( ~ run, nrow = 1) +
    ggplot2::labs(x = "Year", y = "Living C t/ha", fill = "Species") +
   ggplot2::theme_bw()
```





Note that we do not have management intervention, and in the model trees shorter than 4m are not included in these outputs. However as they grow taller than 4m, model start to handle them as individual trees and including in these outputs which we are looking now.

At the end of the simulated 100 years, the total simulated carbon stock in living biomass under climate change scenario was slightly higher than under the reference climate. The proportion of beech and pine increased under drier and warmer climate, while the proportion of spruce was lower, since spruce is in general more susceptible to dry conditions. The changes in the number of trees of individual tree species were more substantial than the changes in carbon stock. The number of spruce trees was substantially reduced under climate change, while under reference climate it was doubled after 30 years and then remain stable until the end of simulation. In contrast, the number of beech trees increased under both scenarios, but the increase was more pronounced under climate change scenario due to the reduction of spruce trees.

Mean dbh of individual species reflected the development of the number of trees. The increase in the number of trees caused the reduction of mean dbh, e.g. of beech under both scenarios between 20 and 35 simulation years. The species-specific graphs for mean dbh indicates that some species do not have successful regeneration under the climate change scenario. Spruce and pine dbh decreased only under reference scenario, when we observed the increase of the number of trees, while under climate change scenario the mean dbh of spruce or pine was growing indicating the growth of remaining trees of the species at the plot. In the case of fir, we observed the continuous increase of dbh due

to the more or less stable number of trees at the plots. While the mean dbh of fir was growing during the whole simulation period, under drier and warmer climate mean dbh of fir did not substantially change after 70 years.

Calculate the stored C amount in the initial year (year==0) and the last year (year==100)!

```
livingC0 <-
    data.frame(
        landscape |> dplyr::filter(year == 0) |>
              dplyr::group_by(run) |>
                  dplyr::summarize(sum.livingC = sum(total_carbon_kg / 1000))
        )
print(livingC0)
```

```
run sum.livingC
1 reference 139.9371
2 4deg_20percdrier 139.9371
```

```
run sum.livingC
1 reference 465.9313
2 4deg_20percdrier 502.2416
```

The initial conditions are same for the runs, but they are ending up at different C levels.

Calculate the stored C amount PER SPECIES in the initial year (year==0) and the last year (year==100)

```
species.livingC0 <-
    data.frame(
    landscape |> dplyr::filter(year == 0) |>
        dplyr::group_by(run, species) |>
        dplyr::summarize(livingC = sum(total_carbon_kg / 1000))
    )
print(species.livingC0)
```

```
run species livingC
1
                      abal 2.26449
        reference
2
        reference
                      fasy 49.66745
3
        reference
                     piab 45.42464
        reference
4
                     pisy 42.58056
5 4deg_20percdrier
                     abal 2.26449
6 4deg_20percdrier
                      fasy 49.66745
7 4deg_20percdrier
                     piab 45.42464
8 4deg_20percdrier
                     pisy 42.58056
species.livingC100 <-</pre>
   data.frame(
        landscape |> dplyr::filter(year == 100) |>
            dplyr::group_by(run, species) |>
            dplyr::summarize(livingC = sum(total_carbon_kg / 1000))
    )
print(species.livingC100)
```

```
run species
                           livingC
                    abal 117.08641
1
        reference
2
        reference
                    fasy 130.23479
3
        reference piab 192.96180
        reference pisy 25.64826
5 4deg_20percdrier abal 123.25491
6 4deg_20percdrier
                   fasy 187.43320
7 4deg_20percdrier
                    piab 150.38585
8 4deg_20percdrier
                    pisy 41.16762
```

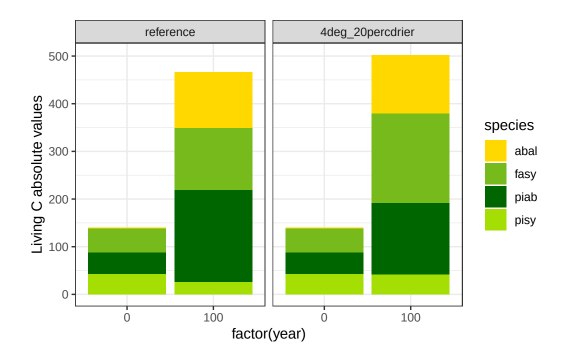
Calculate the species proportions based on the stored C amount in the initial year (year==0) and the last year (year==100) For this we need the total C amount and the species-specific C amount.

```
run species livingC sum.livingC spec.prop year
1 reference abal 2.26449 139.9371 0.01618219 0
2 reference fasy 49.66745 139.9371 0.35492686 0
3 reference piab 45.42464 139.9371 0.32460745 0
```

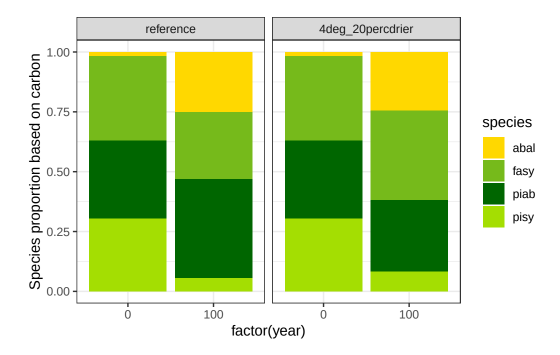
```
pisy 42.58056
                                      139.9371 0.30428349
        reference
                     abal 2.26449
5 4deg_20percdrier
                                      139.9371 0.01618219
                                                            0
                     fasy 49.66745
6 4deg_20percdrier
                                      139.9371 0.35492686
                                                            0
7 4deg_20percdrier
                     piab 45.42464
                                      139.9371 0.32460745
                                                            0
8 4deg_20percdrier
                     pisy 42.58056
                                      139.9371 0.30428349
                                                            0
LC100 <- dplyr::left_join(species.livingC100, livingC100, by = "run")
LC100 <-
   data.frame(LC100 |> dplyr::mutate(spec.prop = livingC / sum.livingC, year =
                                         100))
print(LC100)
```

```
livingC sum.livingC spec.prop year
             run species
                    abal 117.08641
                                    465.9313 0.25129545 100
1
        reference
                   fasy 130.23479
                                    465.9313 0.27951504 100
2
        reference
3
        reference piab 192.96180
                                    465.9313 0.41414220 100
        reference pisy 25.64826
                                    465.9313 0.05504730 100
5 4deg_20percdrier abal 123.25491
                                    502.2416 0.24540961 100
6 4deg_20percdrier fasy 187.43320
                                    502.2416 0.37319331 100
7 4deg_20percdrier
                   piab 150.38585
                                    502.2416 0.29942930 100
8 4deg_20percdrier
                   pisy 41.16762
                                    502.2416 0.08196777 100
```

Put the two tables together and visualize the results!



```
ggplot2::ggplot(LC, aes(fill = species, y = spec.prop, x = factor(year))) +
    ggplot2::geom_bar(position = "stack", stat = "identity") +
    ggplot2::scale_fill_manual(values = cols.all) +
    ggplot2::facet_wrap( ~ run) +
    ggplot2::ylab("Species proportion based on carbon") +
    ggplot2::theme_bw()
```



The living C stock tripled over 100 years due to the accumulation of biomass in trees and no management interventions. Moreover, the proportion of individual tree species changed with fir substantially increasing its share. The living C stock of pine was reduced under the reference climate or remained unchanged under the climate change scenario.

2.2.8 Question A - Group 2

• How are biodiversity indices changing in time and across the simulated scenario(s) on Plot 2?

Run the model as it is described in the previous chapters to have at least 2 simulations completed: one with reference conditions and one scenario with some changes in the environment. Check if you have the output files in the output folder: *iLand_simulations/output/*. If you manage to simulate 2 scenarios, you can expand the code to have file3, and 3rd from each variable, or make the comparison in 2 step always comparing one scenario to the reference conditions at the time.

Open the summerSchoolExerciseProcessBased.Rproj project in R studio and start working there.

Read in the *tree* output table from two outputs you have for your plot:

```
path1 <- "model/iLand_simulations/output/Output_plot2.sqlite"</pre>
path2 <-
    "model/iLand simulations/output/Output_plot2 4deg_20percdrier.sqlite"
file1 <- here::here(path1)</pre>
file2 <- here::here(path2)</pre>
# Give some name for the three simulations.
name1 <- "reference"</pre>
name2 <- "4deg_20percdrier"</pre>
# Read in data using the RSQLite package
sqlite.driver <- RSQLite::dbDriver("SQLite")</pre>
db1 <-
    RSQLite::dbConnect(sqlite.driver, dbname = file1) # connect to the file1
tables.in.the.file <-
    RSQLite::dbListTables(db1) # explore the tables in the file1
print(tables.in.the.file)
[1] "carbon"
                         "carbonflow"
                                              "dynamicstand"
[4] "landscape"
                         "landscape_removed" "runinfo"
[7] "stand"
                         "tree"
# We will work with "tree" table and tree-scale data:
tree1 <- RSQLite::dbReadTable(db1, "tree")</pre>
RSQLite::dbDisconnect(db1) # disconnect to the file1
# READ IN DATA FROM THE SECOND FILE: ----
db2 <-
    RSQLite::dbConnect(sqlite.driver, dbname = file2) # connect to the file2
tree2 <- RSQLite::dbReadTable(db2, "tree")</pre>
RSQLite::dbDisconnect(db2)
```

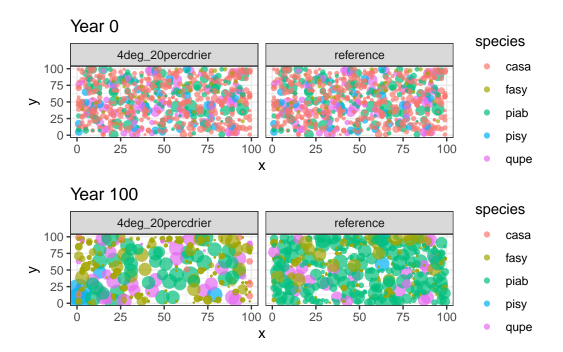
Merge the data from the two files together and make a column which tells which comes from which simulation, and study the table. The column "run" will support the plotting where we can use facets in ggplot.

head(tree)

```
year ru rid species id x y
                                     dbh
                                           height basalArea volume_m3
     0
        0
                 piab
                       1 65 43 85.63804 57.57916 0.5760011 14.029073
1
2
     0
        0
                         9 47 78.57278 40.23281 0.4848798 8.856667 1072
3
                       3 95 39 78.37173 52.69362 0.4824016 10.752442
     0
        0
                                                                        564
4
     0
        0
                       4 95 91 76.88074 51.69115 0.4642212 10.150361
                                                                        553
                                                                        536
5
     0
        0
                 piab 5 79 75 74.49680 50.08829 0.4358782
                                                             9.235103
            1
6
     0
        0
            1
                 qupe 6 89 49 71.96641 36.85005 0.4067706
                                                             6.805241
                                                                        982
  leafArea m2 foliageMass stemMass branchMass fineRootMass coarseRootMass
     420.3857
                 98.91427 3815.740
                                      613.1350
                                                    74.18570
                                                                   986.7341
1
2
     394.8285
                 41.56089 3702.413
                                      365.5260
                                                    31.17067
                                                                  1160.0538
3
     366.0804
                 86.13657 3067.633
                                      500.0222
                                                    64.60242
                                                                   770.4860
4
     355.2738
                 83.59383 2925.986
                                      478.4131
                                                    62.69537
                                                                   730.2825
5
     338.2381
                 79.58543 2707.714
                                      444.9787
                                                    59.68908
                                                                   668.8427
                 35.79670 3147.565
                                      311.7678
     340.0686
                                                    26.84752
                                                                   936.2913
        lri lightResponse stressIndex reserve_kg treeFlags
                                                                   run
1 0.4567858
                         0
                                        173.09998
                                                           0 reference
2 0.2441241
                        0
                                         72.73156
                                                           0 reference
                                     0
3 1.0000000
                        0
                                     0
                                       150.73898
                                                           0 reference
4 1.0000000
                        0
                                     0
                                        146.28922
                                                           0 reference
5 0.4540371
                         0
                                        139.27451
                                     0
                                                           0 reference
6 0.2951740
                         0
                                         62.64423
                                                           0 reference
```

To explore a bit the simulation, plot the tree locations at the beginning and the end of the simulation! For plotting we use the colors for the species, and the size of a circles to show dbh differences. Here we divided dbh with the value 20 just for visualization, not to have too huge circles shading each others completely, but still see the trees.

```
ggplot2::ggtitle("Year 100") +
    ggplot2::facet_wrap(~ run) + ggplot2::theme_bw()
gridExtra::grid.arrange(g1, g2, ncol = 1)
```

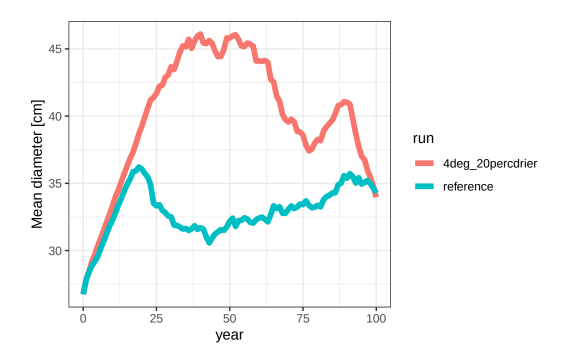


We can see that the species composition has changed and the dbh of the trees seems to be larger, than in the initial year. We can even identify the same trees at the same location.

Visualize some changes in time, for example the mean diameter of the trees!

```
sum.table <- tree |> dplyr::group_by(year, run) |>
    dplyr::summarise(
        N = dplyr::n(),
        MD = mean(dbh, na.rm = TRUE),
        #mean diameter
        BA = sum(basalArea)
    ) #basal area

ggplot2::ggplot(sum.table, ggplot2::aes(x = year, y = MD, color = run)) +
        ggplot2::geom_line(lwd = 2) +
        ggplot2::ylab("Mean diameter [cm]") + ggplot2::theme_bw()
```



We can see that there is an initial increase in mean dbh, then a drop after 20/25 years under reference conditions. Under the climate change scenario, the mean dbh is growing for longer time and has the drop after 60 years. The dbh in the last year are very similar however the development of the simulations were different. The drop is due to the growing regeneration layer that is produced by the seeds of the existing trees. In the initial year we do not have regeneration layer in the simulations (it is possible to put there, but we do not have it now). And until the small trees grow up to higher than 4 m they are not appearing in the individual tree output as they are handled in cohorts.

To study biodiversity aspects, we can calculate biodiversity indicators. Let's use the adiv package for species diversity. First we need to change the database structure to have the number of trees for each species in different columns (pivot_wider) without any additional columns for the adiv package speciesdiv function. We work first only with tree data from one simulation (tree1). You can find the documentation of the adiv package here: "documents/adiv.pdf"

```
# First we need to change the
tlong1 <- tree1 |>
    dplyr::group_by(year, species) |> dplyr::summarise(N = dplyr::n()) |>
    tidyr::pivot_wider(
        id_cols = 'year',
        names_from = 'species',
        values_from = 'N',
        values_fill = 0
```

```
dplyr::ungroup() |>
  dplyr::select(-year)
head(tlong1)
```

```
# A tibble: 6 x 5
   casa fasy piab pisy qupe
  <int> <int> <int> <int> <int>
1
    385
           34
                118
                        32
                              68
2
    358
           34
                        30
                              64
                117
3
    335
           33
                115
                        30
                              63
4
    319
           33
                114
                        30
                              63
5
    310
           32
                112
                        29
                              63
    303
                        29
                              62
           31
                112
```

Then we apply the "speciesdiv" function and we add back the year and run columns to have the information.

```
div1 <- data.frame(adiv::speciesdiv(tlong1)) |>
     dplyr::mutate(year = unique(tree1$year), run = name1)
head(div1)
```

```
richness GiniSimpson Simpson Shannon Margalef Menhinick McIntosh year
1
             0.5836227 2.401668 1.162161 0.6195048 0.1981072 0.3693616
2
         5
             0.5929556 2.456735 1.177198 0.6248128 0.2036157 0.3773673
                                                                            1
3
         5
             0.6039255\ 2.524778\ 1.196665\ 0.6293160\ 0.2083333\ 0.3867710
                                                                            2
4
             0.6136885 2.588585 1.214413 0.6322962 0.2114775 0.3951745
         5
                                                                            3
5
             0.6159951\ 2.604134\ 1.217670\ 0.6346568\ 0.2139802\ 0.3973226
                                                                            4
         5
             0.6185478 2.621560 1.221342 0.6363349 0.2157659 0.3996270
         5
1 reference
2 reference
3 reference
4 reference
5 reference
6 reference
```

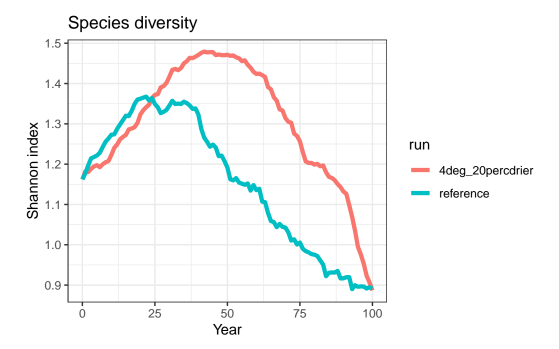
We make the same steps for the other simulation results (tree2).

```
tlong2 <- tree2 |>
    dplyr::group_by(year, species) |> dplyr::summarise(N = n()) |>
    tidyr::pivot_wider(
        id_cols = 'year',
        names_from = 'species',
        values_from = 'N',
        values_fill = 0
    ) |>
    dplyr::ungroup() |>
    dplyr::select(-year)

div2 <- data.frame(adiv::speciesdiv(tlong2)) |>
    dplyr::mutate(year = unique(tree2$year), run = name2)
```

We merge them all together and make a plot based on Shannon index.

```
tlong2 <- tree2 |>
    dplyr::group_by(year, species) |> dplyr::summarise(N = n()) |>
    tidyr::pivot_wider(
        id cols = 'year',
        names_from = 'species',
        values_from = 'N',
        values_fill = 0
    ) |>
    dplyr::ungroup() |>
    dplyr::select(-year)
div2 <- data.frame(adiv::speciesdiv(tlong2)) |>
    dplyr::mutate(year = unique(tree2$year), run = name2)
div <- rbind(div1, div2)</pre>
ggplot2::ggplot(div, aes(year, Shannon, color = run)) +
    ggplot2::geom_line(lwd = 1.5) +
    ggplot2::ggtitle("Species diversity") +
    ggplot2::labs(x = "Year", y = "Shannon index") +
    ggplot2::theme_bw()
```



We see that the values of Shannon index are first increasing and then tend to decrease, while the reduction under the 4deg_20percdrier is starting later than under the reference scenario. Shannon index is calculated as follows:

$$S = -\sum \left(p \ i \ *log(p \ i \) \right)$$

Where p i is the relative abundance of the species i, calculated as the ratio of the abundance of the species i and the abundance of all species. Shannon-index is 0 when we have only one species at the stand.

We can calculate some additional biodiversity indices targeting spatial diversity using the *treespat* package developed by Francesco Chianucci (fchianucci@gmail.com). You can find a short description of the package here: "documents/treespat_package.pdf"

You can install the package using devtools:

Then, we are ready to use it! Let's calculate these two indices:

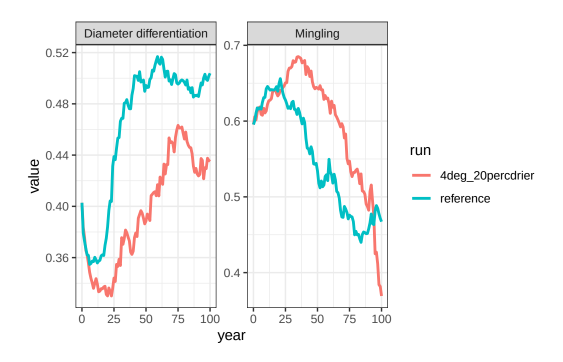
• Diameter differentiation (Gadow, 1993): Spatial size inequality defined as the mean of the ratio of smaller and larger plant sizes in the nearest neighbors of a tree. The value of the index increases with increasing average size difference between neighboring trees. 0 is implying that neighboring trees have equal size.

• Mingling (Aguirre et al., 2003): One very intuitive extension of taxonomic species diversity (either richness or abundance) is considering spatial mingling, namely how plants of the same (con-specific neighbors) or different (hetero-specific neighbors) species are arranged in space. The mingling index calculates the proportion of the k nearest neighbors that do not belong to the same species as the reference tree. For example, with four neighbors, the mingling attribute can assume five values, ranging from 0 (all trees are of the same species) to 1 (all trees belong to different species).

Calculate the indices based on the reference run, then based on the scenario. For each index we add extra columns with the index name, and run name for further analyses. The max.k parameter is telling how many neighboring trees we want to account for.

```
library(treespat)
# diameter differentiation:
# mingling
diff <- data.frame(treespat::DIFF(tree1, .x = x, .y = y,.mark = dbh, xmax = 100,
                  ymax = 100, max.k = 4, shape='square', .groups=c('year')) |>
         dplyr::mutate(index="DIFF", name="Diameter differentiation", run=name1))
head(diff)
            DIFF index
  year
                                           name
                                                      run
     0 0.4027813 DIFF Diameter differentiation reference
1
2
     1 0.3793126 DIFF Diameter differentiation reference
3
     2 0.3728755 DIFF Diameter differentiation reference
     3 0.3672772 DIFF Diameter differentiation reference
     4 0.3621167 DIFF Diameter differentiation reference
5
     5 0.3617524 DIFF Diameter differentiation reference
ming <- data.frame(treespat::MING(tree1, .x = x, .y = y, .species = species,
        xmax = 100, ymax = 100, max.k = 4, shape='square', .groups=c('year')) |>
        dplyr::mutate(index="MING", name="Mingling", run=name1))
indices1<-rbind(diff |> dplyr::rename(value=DIFF),
 ming |> dplyr::rename(value=MINGLING ) )
```

We calculated each index as separate variable and then merged them into indices1. Here I renamed the column names to have it all "value", for easier plotting later on. (each line has the information on which run and which index is the value) We do the same for the scenario simulation, we merge them together and make the plotting.



The diameter differentiation increases with the increasing average difference in diameter between neighboring trees. The value 0 indicates that neighboring trees have equal

diameters. The development shows that the index values decrease during the first circa 10 years, after which it steeply increases due to the occurrence of ingrowth and then the values are stabilized. The increase is not that steep for the climate change scenario that is in accordance with the mean dbh graph. The values for the reference climate scenario are higher than for the climate change scenario indicating slower growth under drier and warmer conditions and/or the differences in tree species composition (proportion of tree species). The mingling index is studying the neighboring trees regarding their species. In both development there is a decrease in the index after the occurrence of regeneration. Using the climate change scenario the index shows steeper decrease and ending up at lower values after 100 years.

2.2.9 Question B - Group 2

• How are the species distribution and total living biomass C content changing in time on Plot 2? Compare 0 year and 100 year status in the reference case and in the case of your scenario(s)!

Run the model as it is described in the previous chapters to have at least 2 simulations completed: one with reference conditions and one scenario with some changes in the environment. Check if you have the output files in the output folder: <code>iLand_simulations/output/</code>. If you manage to simulate 2 scenarios, you can expand the code to have file3, and 3rd from each variable, or make the comparison in 2 step always comparing one scenario to the reference conditions at the time. Open the <code>summerSchoolExercise-ProcessBased.Rproj</code> project in R studio and start working there.

Read in the *landscape* output table from two outputs you have for your plot:

```
path1 <- "model/iLand_simulations/output/Output_plot2.sqlite"
path2 <-
     "model/iLand_simulations/output/Output_plot2_4deg_2Opercdrier.sqlite"

file1 <- here::here(path1)
file2 <- here::here(path2)

# Give some name for the three simulations.
name1 <- "reference"
name2 <- "4deg_2Opercdrier"

# Read in data using the RSQLite package
sqlite.driver <- RSQLite::dbDriver("SQLite")

# We will work with "landscape" table and tree-scale data:
db1 <-</pre>
```

```
RSQLite::dbConnect(sqlite.driver, dbname = file1) # connect to the file1
landscape1 <- RSQLite::dbReadTable(db1, "landscape")
RSQLite::dbDisconnect(db1) # disconnect to the file1

db2 <-
    RSQLite::dbConnect(sqlite.driver, dbname = file2) # connect to the file2
landscape2 <- RSQLite::dbReadTable(db2, "landscape")
RSQLite::dbDisconnect(db2)</pre>
```

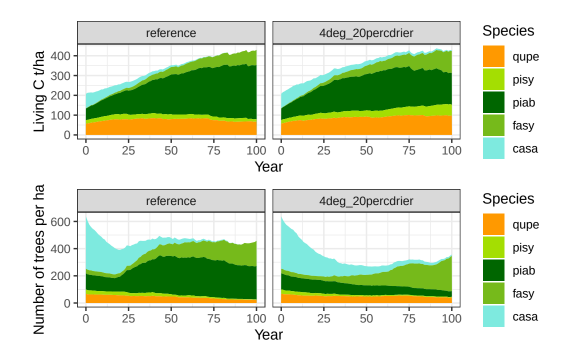
Merge the data from the three files together and make a column which tells which comes from which simulation, and study the table. The column "run" will support the plotting where we can use facets in ggplot.

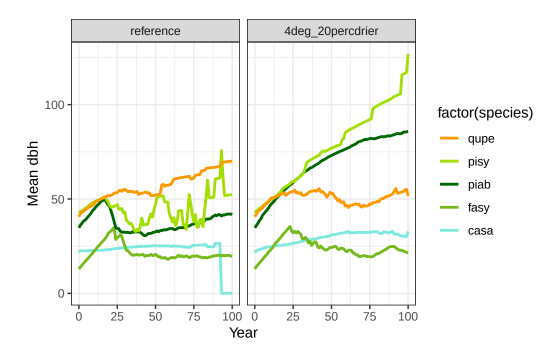
```
year area area_100m species count_ha dbh_avg_cm height_avg_m volume_m3
1
     0
          1
                     1
                          casa
                                     385
                                           21.73276
                                                         18.052312 139.177831
2
     0
          1
                                                          7.340003
                                                                     1.600727
                     1
                          fasy
                                      34
                                            12.91628
3
     0
          1
                     1
                          piab
                                     118
                                           34.75135
                                                        23.365242 215.003187
4
     0
          1
                     1
                                           42.76543
                                                         24.300133
                                                                    49.013727
                          pisy
                                      32
5
     0
          1
                     1
                                           40.62035
                                                        20.799453 118.797078
                          qupe
                                      68
6
     1
          1
                     1
                                     358
                                           22.49524
                                                        18.644885 138.166090
                          casa
  total_carbon_kg
                       gwl_m3 basal_area_m2
                                                NPP_kg NPPabove_kg
                                                                            LAI
        73246.102 139.177831
                                  16.1066092
                                                 0.000
                                                              0.000 1.92032457
1
2
                     1.600727
                                                 0.000
         1636.821
                                   0.4644299
                                                              0.000 0.06445078
3
        58083.681 215.003187
                                  14.4589881
                                                 0.000
                                                              0.000 1.36948128
4
                  49.013727
                                                              0.000 0.51041807
        18250.374
                                   4.6099225
                                                 0.000
5
        56880.735 118.797078
                                  10.0244600
                                                 0.000
                                                              0.000 0.94624017
6
        73118.915 141.336043
                                  15.8220002 8338.849
                                                           5192.833 1.87298787
  cohort_count_ha
                         run
1
                 0 reference
2
                 0 reference
3
                 0 reference
4
                 0 reference
5
                 0 reference
6
                 0 reference
```

We can see that the output is given for each year and each species in our 1ha area. Plot the living carbon (total_carbon_kg) in time, coloring by species. We give a unified

species coloring in the beginning. Plot number of trees and mean diameter. Carbon and number of trees can be plotted in an additive way, but for mean dbh we do line plot per species.

```
cols.all=c( "rops"="#e0e0e0", "acpl"="#A9A9A9", "alin"="#696969", "alvi"="#2e2e2e",
 "bepe"="#fadfad", "casa"="#7eeadf", "coav"="#20c6b6", "tipl"="#645394",
 "ulgl"="#311432", "saca"="#D8BFD8", "soar"="#DDA0DD", "soau"="#BA55D3",
 "pice"="#D27D2D", "pini"="#a81c07", "alg1"="#2ECBE9", "tico"="#128FC8",
 "potr"="#00468B", "poni"="#5BAEB7", "frex"="#fe9cb5", "cabe"="#fe6181",
 "acps"="#fe223e","lade"="#FFFE71","abal"="#FFD800", "pisy"="#A4DE02",
 "fasy"="#76BA1B", "piab"="#006600", "quro"="#FF7F00", "qupe"="#FF9900",
 "qupu"="#CC9900")
# We set here the order of the run categories for the table, to have the first
#run first and then the second. (left-right of the plots)
landscape$run <- factor(landscape$run, levels = c(name1, name2))</pre>
# Plot the living carbon content, to have tonnes/ha, we divide total carbon kg by 1000.
g1 <-
    ggplot2::ggplot(landscape, aes(year, total_carbon_kg / 1000 , fill = factor(species))
    ggplot2::geom_area() +
    ggplot2::scale_fill_manual(values = cols.all, guide = guide_legend(reverse =
                                                                             TRUE)) +
    ggplot2::facet_wrap( ~ run, nrow = 1) +
    ggplot2::labs(x = "Year", y = "Living C t/ha", fill = "Species") +
    ggplot2::theme_bw()
g2 <-
    ggplot2::ggplot(landscape, aes(year, count_ha , fill = factor(species))) +
    ggplot2::geom_area() +
    ggplot2::scale_fill_manual(values = cols.all, guide = guide_legend(reverse =
                                                                             TRUE)) +
    ggplot2::facet_wrap( ~ run, nrow = 1) +
    ggplot2::labs(x = "Year", y = "Number of trees per ha", fill = "Species") +
    ggplot2::theme_bw()
gridExtra::grid.arrange(g1, g2, ncol = 1)
```





Note that we do not have management intervention, and in the model trees shorter than 4m are not included in these outputs. However as they grow taller than 4m, model start to handle them as individual trees and including in these outputs which we are looking now.

Castanea sativa gradually decreased during the simulation period under both scenarios. The reduction resulted from the mortality of chestnut trees, indicating their aging. By the end of the simulation period, chestnut almost completely disappeared from the stand due to the increased number of trees of other species at the plot, and hence higher intertree competition, especially under the reference climate, which hindered the chestnut regeneration and its survival. Mean dbh of chestnut is zero at the end of the simulation period under the reference climate indicating that the species completely disappeared from the stand due to the high stand density, and inter-tree competition.

Calculate the stored C amount in the initial year (year==0) and the last year (year==100)!

```
livingC0 <- data.frame(
    landscape |> dplyr::filter(year == 0) |>
        dplyr::group_by(run) |>
        dplyr::summarize(sum.livingC = sum(total_carbon_kg / 1000))
)
print(livingC0)
```

The initial conditions are same for the runs, but they are ending up at different C levels.

Calculate the stored C amount PER SPECIES in the initial year (year==0) and the last year (year==100)

```
species.livingC0 <-
    data.frame(
        landscape |> dplyr::filter(year == 0) |>
              dplyr::group_by(run, species) |>
              dplyr::summarize(livingC = sum(total_carbon_kg / 1000))
    )
print(species.livingC0)
```

```
run species
                           livingC
                     casa 73.246102
1
         reference
2
         reference
                     fasy 1.636821
3
         reference piab 58.083681
         reference pisy 18.250374
4
         reference qupe 56.880735
5
6 4deg_20percdrier
                     casa 73.246102
7 4deg_20percdrier fasy 1.636821
8 4deg_20percdrier piab 58.083681
9 4deg_20percdrier
                     pisy 18.250374
10 4deg_20percdrier
                     qupe 56.880735
```

```
species.livingC100 <-
    data.frame(
    landscape |> dplyr::filter(year == 100) |>
        dplyr::group_by(run, species) |>
        dplyr::summarize(livingC = sum(total_carbon_kg / 1000))
    )
print(species.livingC100)
```

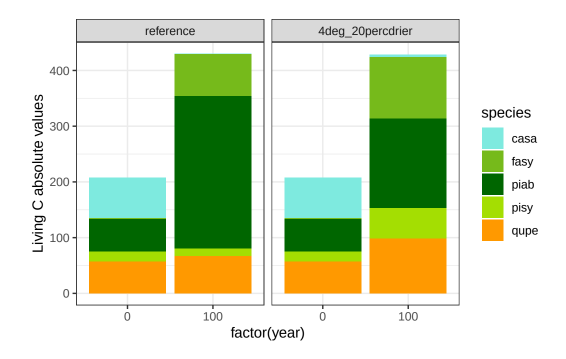
```
run species
                                livingC
                               0.00000
1
          reference
                       casa
2
          reference
                       fasv
                             75.469157
3
          reference
                       piab 274.331862
4
          reference
                       pisy
                              12.701789
5
          reference
                       qupe
                              67.044909
6 4deg_20percdrier
                       casa
                               4.546279
7
 4deg_20percdrier
                       fasy 110.708431
8 4deg_20percdrier
                       piab 159.773037
9 4deg_20percdrier
                       pisy
                             55.439370
10 4deg_20percdrier
                       qupe
                              97.802295
```

Calculate the species proportions based on the stored C amount in the initial year (year==0) and the last year (year==100) For this we need the total C amount and the species-specific C amount.

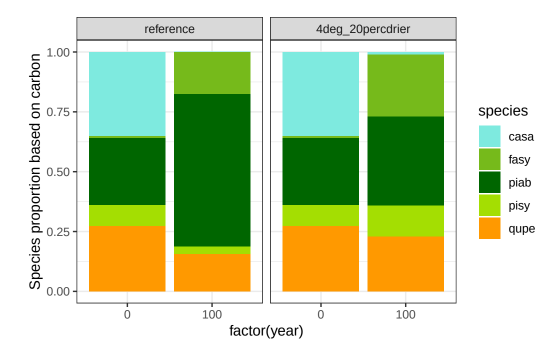
```
livingC sum.livingC
                run species
                                                     spec.prop year
1
          reference
                       casa 73.246102
                                          208.0977 0.351979370
                                                                  0
2
          reference
                       fasy 1.636821
                                          208.0977 0.007865637
                                                                  0
3
                       piab 58.083681
          reference
                                          208.0977 0.279117344
                                                                  0
4
          reference
                       pisy 18.250374
                                          208.0977 0.087700985
                                                                  0
5
          reference
                       qupe 56.880735
                                          208.0977 0.273336665
                                                                  0
6 4deg_20percdrier
                       casa 73.246102
                                          208.0977 0.351979370
                                                                  0
7 4deg_20percdrier
                       fasy 1.636821
                                          208.0977 0.007865637
                                                                  0
8 4deg_20percdrier
                       piab 58.083681
                                          208.0977 0.279117344
                                                                  0
9 4deg_20percdrier
                                          208.0977 0.087700985
                       pisy 18.250374
                                                                  0
10 4deg_20percdrier
                                          208.0977 0.273336665
                       qupe 56.880735
                                                                  0
```

```
run species
                           livingC sum.livingC spec.prop year
                                      429.5477 0.00000000
1
         reference
                   casa
                           0.000000
                                                        100
2
                    fasy 75.469157
                                      429.5477 0.17569447
                                                         100
        reference
3
                    piab 274.331862
                                      429.5477 0.63865282 100
        reference
4
                                      429.5477 0.02957015 100
        reference pisy 12.701789
5
        reference
                    qupe 67.044909 429.5477 0.15608256 100
                                    428.2694 0.01061547 100
                           4.546279
6 4deg_20percdrier
                    casa
7 4deg_20percdrier fasy 110.708431 428.2694 0.25850184 100
8 4deg_20percdrier
                    piab 159.773037
                                      428.2694 0.37306666 100
                    pisy 55.439370
9 4deg_20percdrier
                                      428.2694 0.12944975
                                                        100
10 4deg_20percdrier
                    qupe 97.802295
                                      428.2694 0.22836629 100
```

Put the two tables together and visualize the results!



```
ggplot2::ggplot(LC, aes(fill = species, y = spec.prop, x = factor(year))) +
    ggplot2::geom_bar(position = "stack", stat = "identity") +
    ggplot2::scale_fill_manual(values = cols.all) +
    ggplot2::facet_wrap( ~ run) +
    ggplot2::ylab("Species proportion based on carbon") +
    ggplot2::theme_bw()
```



The living carbon stock doubled over 100 years due to the accumulation of biomass at the plot irrespective of the climate scenario. The comparison of the results between the two climate scenarios showed the difference in the tree species share. The living C stock of spruce was lower under the climate change reflecting its susceptibility to dry conditions, while the living C stock of Quercus pubescens, Fagus sylvatica and Pinus sylvestris were higher. Additionally, we see a significant increase in the share of the shade-tolerant beech at the expense of light-demanding chestnut.

2.2.10 Question A - Group 3

• How are biodiversity indices changing in time and across the simulated scenario(s) on Plot 3?

Run the model as it is described in the previous chapters to have at least 2 simulations completed: one with reference conditions and one scenario with some changes in the environment. Check if you have the output files in the output folder: $iLand_simulations/output/$. If you manage to simulate 2 scenarios, you can expand the code to have file3, and 3rd from each variable, or make the comparison in 2 step always comparing one scenario to the reference conditions at the time.

Open the *summerSchoolExerciseProcessBased.Rproj* project in R studio and start working there.

Read in the *tree* output table from two outputs you have for your plot:

```
path1 <- "model/iLand_simulations/output/Output_plot3.sqlite"</pre>
path2 <-
    "model/iLand simulations/output/Output_plot3 4deg_20percdrier.sqlite"
file1 <- here::here(path1)</pre>
file2 <- here::here(path2)</pre>
# Give some name for the three simulations.
name1 <- "reference"</pre>
name2 <- "4deg_20percdrier"</pre>
# Read in data using the RSQLite package
sqlite.driver <- RSQLite::dbDriver("SQLite")</pre>
db1 <-
    RSQLite::dbConnect(sqlite.driver, dbname = file1) # connect to the file1
tables.in.the.file <-
    RSQLite::dbListTables(db1) # explore the tables in the file1
print(tables.in.the.file)
[1] "carbon"
                         "carbonflow"
                                              "dynamicstand"
[4] "landscape"
                         "landscape_removed" "runinfo"
[7] "stand"
                         "tree"
# We will work with "tree" table and tree-scale data:
tree1 <- RSQLite::dbReadTable(db1, "tree")</pre>
RSQLite::dbDisconnect(db1) # disconnect to the file1
# READ IN DATA FROM THE SECOND FILE: ----
db2 <-
    RSQLite::dbConnect(sqlite.driver, dbname = file2) # connect to the file2
tree2 <- RSQLite::dbReadTable(db2, "tree")</pre>
RSQLite::dbDisconnect(db2)
```

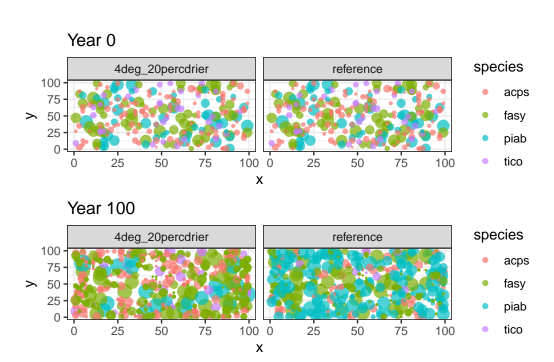
Merge the data from the two files together and make a column which tells which comes from which simulation, and study the table. The column "run" will support the plotting where we can use facets in ggplot.

head(tree)

```
height basalArea volume_m3 age
  year ru rid species id x y
                                     dbh
     0
        0
                 piab
                       1
                         7 41 80.47096 54.10505 0.5085905 11.639825 579
1
2
     0
        0
                 piab 2 57 63 77.85299 52.34484 0.4760367 10.540342 560
3
                       3 33 45 77.32411 43.94139 0.4695910 8.934730 549
     0
        0
            1
4
     0
        0
                       4 79 97 76.44166 43.43991 0.4589338 8.632307 542
            1
5
     0
        0
                 piab 5 99 35 74.65018 50.19142 0.4376748
                                                             9.292261 537
            1
6
     0
        0
            1
                 fasy 6 11 77 74.61653 42.40273 0.4372804 8.028635 530
  leafArea m2 foliageMass stemMass branchMass fineRootMass coarseRootMass
     381.4915
                 89.76272 3273.843
                                      531.3649
                                                   67.32204
                                                                   829.4560
1
2
     362.3074
                 85.24881 3017.901
                                      492.4429
                                                   63.93661
                                                                   756.3417
     387.3896
3
                 35.21723 3946.903
                                      484.7826
                                                   26.41292
                                                                   432.6764
4
     379.9038
                 34.53671 3846.260
                                      472.1521
                                                   25.90253
                                                                   421.3067
5
     339.3250
                 79.84119 2721.455
                                      447.0886
                                                   59.88089
                                                                   672.6917
                 33.14663 3642.679
                                      446.6253
     364.6130
                                                   24.85997
                                                                   398.3362
        lri lightResponse stressIndex reserve_kg treeFlags
                                                                   run
1 0.5557720
                        0
                                       157.08476
                                                          0 reference
2 0.3826686
                        0
                                     0
                                       149.18542
                                                          0 reference
3 0.4321580
                        0
                                     0
                                         61.63016
                                                          0 reference
4 0.7127665
                        0
                                     0
                                         60.43925
                                                          0 reference
5 1.0000000
                        0
                                       139.72208
                                                          0 reference
                                     0
6 0.3028383
                        0
                                         58.00661
                                                           0 reference
```

To explore a bit the simulation, plot the tree locations at the beginning and the end of the simulation! For plotting we use the colors for the species, and the size of a circles to show dbh differences. Here we divided dbh with the value 20 just for visualization, not to have too huge circles shading each others completely, but still see the trees.

```
ggplot2::ggtitle("Year 100") +
    ggplot2::facet_wrap(~ run) + ggplot2::theme_bw()
gridExtra::grid.arrange(g1, g2, ncol = 1)
```

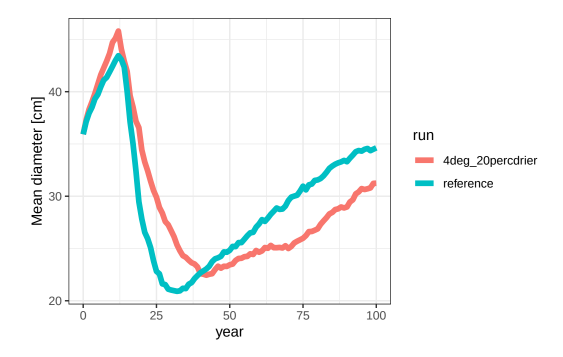


We can see that the species composition is somewhat changed and the forest become more dense. We can even identify the same trees at the same location.

Visualize some changes in time, for example the mean diameter of the trees!

```
sum.table <- tree |> dplyr::group_by(year, run) |>
    dplyr::summarise(
        N = dplyr::n(),
        MD = mean(dbh, na.rm = TRUE),
        #mean diameter
        BA = sum(basalArea)
    ) #basal area

ggplot2::ggplot(sum.table, ggplot2::aes(x = year, y = MD, color = run)) +
        ggplot2::geom_line(lwd = 2) +
        ggplot2::ylab("Mean diameter [cm]") + ggplot2::theme_bw()
```



We can see that there is an initial increase in mean dbh, then a drop after 20/25 years. This is due to the growing regeneration layer that is produced by the seeds of the existing trees. In the initial year we do not have regeneration layer in the simulations (it is possible to put there, but we do not have it now). And until the small trees grow up to higher than 4 m they are not appearing in the individual *tree* output as they are handled in cohorts.

To study biodiversity aspects, we can calculate biodiversity indicators. Let's use the adiv package for species diversity. First we need to change the database structure to have the number of trees for each species in different columns (pivot_wider) without any additional columns for the adiv package speciesdiv function. We work first only with tree data from one simulation (tree1). You can find the documentation of the adiv package here: "documents/adiv.pdf"

```
# First we need to change the
tlong1 <- tree1 |>
    dplyr::group_by(year, species) |> dplyr::summarise(N = dplyr::n()) |>
    tidyr::pivot_wider(
        id_cols = 'year',
        names_from = 'species',
        values_from = 'N',
        values_fill = 0
    ) |>
    dplyr::ungroup() |>
```

```
dplyr::select(-year)
head(tlong1)
# A tibble: 6 x 4
   acps fasy piab tico
  <int> <int> <int> <int>
           67
                  49
                        32
1
    113
2
    110
           67
                  49
                        28
3
    108
                  49
                        27
           67
4
    106
           66
                  47
                        27
5
    104
           65
                  47
                        27
6
    103
           64
                        26
                  46
```

Then we apply the "speciesdiv" function and we add back the year and run columns to have the information.

```
div1 <- data.frame(adiv::speciesdiv(tlong1)) |>
    dplyr::mutate(year = unique(tree1$year), run = name1)
head(div1)
```

```
richness GiniSimpson Simpson Shannon Margalef Menhinick McIntosh year
            0.6963785 3.293574 1.282865 0.5391300 0.2475938 0.4786064
1
2
            0.6935024 3.262668 1.274467 0.5417769 0.2509823 0.4762610
                                                                           1
3
         4
            0.6939255 3.267178 1.274173 0.5429419 0.2524778 0.4768591
                                                                          2
            0.6938000 3.265839 1.274493 0.5449263 0.2550307 0.4770630
                                                                          3
5
            0.6955241 3.284332 1.277850 0.5461435 0.2566001 0.4789300
                                                                          4
             0.6936853 3.264617 1.274063 0.5477988 0.2587385 0.4774250
6
         4
        run
1 reference
2 reference
3 reference
4 reference
5 reference
6 reference
```

We make the same steps for the other simulation results (tree2).

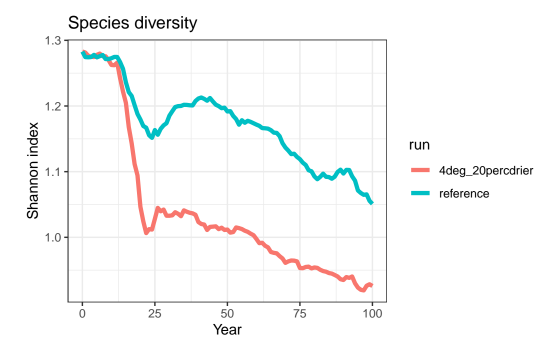
```
tlong2 <- tree2 |>
   dplyr::group_by(year, species) |> dplyr::summarise(N = n()) |>
   tidyr::pivot_wider(
```

```
id_cols = 'year',
    names_from = 'species',
    values_from = 'N',
    values_fill = 0
) |>
    dplyr::ungroup() |>
    dplyr::select(-year)

div2 <- data.frame(adiv::speciesdiv(tlong2)) |>
    dplyr::mutate(year = unique(tree2$year), run = name2)
```

We merge them all together and make a plot based on Shannon index.

```
tlong2 <- tree2 |>
    dplyr::group_by(year, species) |> dplyr::summarise(N = n()) |>
    tidyr::pivot_wider(
        id_cols = 'year',
        names_from = 'species',
        values_from = 'N',
       values_fill = 0
    ) |>
    dplyr::ungroup() |>
    dplyr::select(-year)
div2 <- data.frame(adiv::speciesdiv(tlong2)) |>
    dplyr::mutate(year = unique(tree2$year), run = name2)
div <- rbind(div1, div2)</pre>
ggplot2::ggplot(div, aes(year, Shannon , color = run)) +
    ggplot2::geom_line(lwd = 1.5) +
    ggplot2::ggtitle("Species diversity") +
    ggplot2::labs(x = "Year", y = "Shannon index") +
    ggplot2::theme_bw()
```



We see that the values of Shannon index are first stable and then tend to decrease, while the reduction under the 4deg_20percdrier is more substantial than under the reference scenario. Shannon index is calculated as follows:

$$S = -\sum \left(p \ i \ *log(p \ i \) \right)$$

Where p i is the relative abundance of the species i, calculated as the ratio of the abundance of the species i and the abundance of all species. Shannon-index is 0 when we have only one species at the stand.

We can calculate some additional biodiversity indices targeting spatial diversity using the *treespat* package developed by Francesco Chianucci (fchianucci@gmail.com). You can find a short description of the package here: "documents/treespat_package.pdf"

You can install the package using devtools:

Then, we are ready to use it! Let's calculate these two indices:

• Diameter differentiation (Gadow, 1993): Spatial size inequality defined as the mean of the ratio of smaller and larger plant sizes in the nearest neighbors of a tree. The value of the index increases with increasing average size difference between neighboring trees. 0 is implying that neighboring trees have equal size.

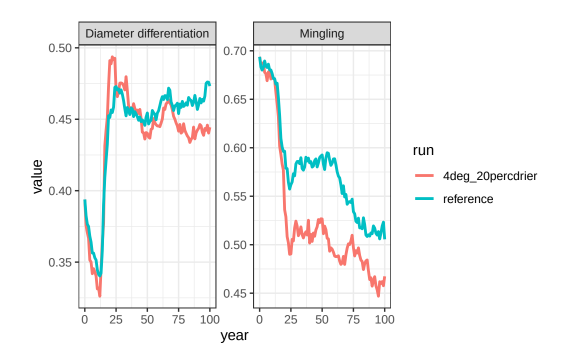
• Mingling (Aguirre et al., 2003): One very intuitive extension of taxonomic species diversity (either richness or abundance) is considering spatial mingling, namely how plants of the same (con-specific neighbors) or different (hetero-specific neighbors) species are arranged in space. The mingling index calculates the proportion of the k nearest neighbors that do not belong to the same species as the reference tree. For example, with four neighbors, the mingling attribute can assume five values, ranging from 0 (all trees are of the same species) to 1 (all trees belong to different species).

Calculate the indices based on the reference run, then based on the scenario. For each index we add extra columns with the index name, and run name for further analyses. The max.k parameter is telling how many neighboring trees we want to account for.

```
# diameter differentiation:
# mingling
diff <- data.frame(treespat::DIFF(tree1, .x = x, .y = y,.mark = dbh, xmax = 100,
                  ymax = 100, max.k = 4, shape='square', .groups=c('year')) |>
         dplyr::mutate(index="DIFF", name="Diameter differentiation", run=name1))
head(diff)
 year
            DIFF index
                                           name
                                                      run
     0 0.3938630 DIFF Diameter differentiation reference
1
2
     1 0.3825122 DIFF Diameter differentiation reference
3
     2 0.3765717 DIFF Diameter differentiation reference
     3 0.3749870 DIFF Diameter differentiation reference
5
     4 0.3681887 DIFF Diameter differentiation reference
     5 0.3639637 DIFF Diameter differentiation reference
ming <- data.frame(treespat::MING(tree1, .x = x, .y = y, .species = species,
        xmax = 100, ymax = 100, max.k = 4, shape='square', .groups=c('year')) |>
        dplyr::mutate(index="MING", name="Mingling", run=name1))
indices1<-rbind(diff |> dplyr::rename(value=DIFF),
ming |> dplyr::rename(value=MINGLING ) )
```

We calculated each index as separate variable and then merged them into indices1. Here I renamed the column names to have it all "value", for easier plotting later on. (each line has the information on which run and which index is the value) We do the same for the scenario simulation, we merge them together and make the plotting.

```
diff <- data.frame(treespat::DIFF(tree2, .x = x, .y = y,.mark = dbh,</pre>
                                   xmax = 100, ymax = 100, max.k = 4,
                                   shape = 'square', .groups = c('year')) |>
      dplyr::mutate(index = "DIFF", name = "Diameter differentiation",
                    run = name2))
ming <- data.frame(treespat::MING(tree2, .x = x, .y = y,
                                   .species = species,
                                   xmax = 100, ymax = 100, max.k = 4,
                                   shape = 'square', .groups=c('year')) |>
                       dplyr::mutate(index = "MING", name = "Mingling",
                                      run = name2))
indices2 <- rbind(diff |> dplyr::rename(value = DIFF),
                  ming |> dplyr::rename(value = MINGLING))
indices <- rbind(indices1, indices2)</pre>
ggplot2::ggplot(indices, aes(x = year, y = value, color = run)) +
    ggplot2::geom_line(lwd = 1) +
    ggplot2::facet_wrap( ~ name, ncol = 2, scales = "free") +
    ggplot2::theme_bw()
```



The diameter differentiation increases with the increasing average difference in diameter between neighboring trees. The value 0 indicates that neighboring trees have equal diameters. The development shows that the index values decrease during the first 20/25 years, after which it steeply increases due to the occurrence of ingrowth and then the values are stabilized. The values for the reference climate scenario are slightly higher than for the climate change scenario indicating slower growth under drier and warmer conditions and/or the differences in tree species composition (proportion of tree species). The mingling index is studying the neighboring trees regarding their species. In both development there is a decrease in the index after the occurrence of regeneration. Using the climate change scenario the index shows steeper decrease.

2.2.11 Question B - Group 3

• How are the species distribution and total living biomass C content changing in time on Plot 3? Compare 0 year and 100 year status in the reference case and in the case of your scenario(s)!

Run the model as it is described in the previous chapters to have at least 2 simulations completed: one with reference conditions and one scenario with some changes in the environment. Check if you have the output files in the output folder: *iLand_simulations/output/*. If you manage to simulate 2 scenarios, you can expand the code to have file3, and 3rd from each variable, or make the comparison in 2 step always comparing one scenario to the reference conditions at the time. Open the *summerSchoolExercise-ProcessBased.Rproj* project in R studio and start working there.

Read in the *landscape* output table from two outputs you have for your plot:

```
path1 <- "model/iLand_simulations/output/Output_plot3.sqlite"
path2 <-
     "model/iLand_simulations/output/Output_plot3_4deg_2Opercdrier.sqlite"

file1 <- here::here(path1)
file2 <- here::here(path2)

# Give some name for the three simulations.
name1 <- "reference"
name2 <- "4deg_2Opercdrier"

# Read in data using the RSQLite package
sqlite.driver <- RSQLite::dbDriver("SQLite")

# We will work with "landscape" table and tree-scale data:
db1 <-</pre>
```

```
RSQLite::dbConnect(sqlite.driver, dbname = file1) # connect to the file1
landscape1 <- RSQLite::dbReadTable(db1, "landscape")
RSQLite::dbDisconnect(db1) # disconnect to the file1

db2 <-
    RSQLite::dbConnect(sqlite.driver, dbname = file2) # connect to the file2
landscape2 <- RSQLite::dbReadTable(db2, "landscape")
RSQLite::dbDisconnect(db2)</pre>
```

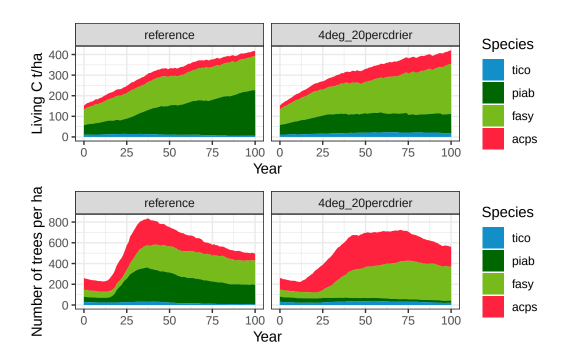
Merge the data from the three files together and make a column which tells which comes from which simulation, and study the table. The column "run" will support the plotting where we can use facets in ggplot.

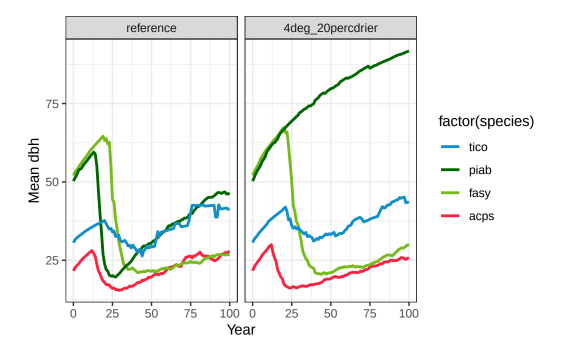
```
year area area_100m species count_ha dbh_avg_cm height_avg_m volume_m3
1
     0
          1
                     1
                          acps
                                     113
                                            21.62995
                                                          18.52183 41.57766
2
     0
          1
                                                          29.58422 218.19995
                     1
                          fasy
                                      67
                                            52.05965
3
     0
          1
                     1
                          piab
                                      49
                                            50.26190
                                                          33.79384 182.14630
4
     0
          1
                     1
                                      32
                                            30.63841
                                                          23.38127
                                                                    21.38005
                          tico
5
     1
          1
                     1
                                     110
                                            22.55262
                                                          19.19236 43.91456
                          acps
6
     1
          1
                     1
                                      67
                                            53.02260
                                                          30.03274 226.37490
                          fasy
  total_carbon_kg
                      gwl_m3 basal_area_m2
                                                NPP_kg NPPabove_kg
         19316.78
                    41.57766
                                   4.693063
                                                 0.000
                                                              0.000 0.4800021
1
2
         74974.41 218.19995
                                                 0.000
                                                              0.000 1.3766679
                                  15.193077
3
         45666.89 182.14630
                                  10.724078
                                                 0.000
                                                              0.000 0.9377907
4
                                                              0.000 0.2673509
         11816.98
                    21.38005
                                   2.379258
                                                 0.000
5
         20284.37
                    44.17968
                                   4.895663
                                              3936.955
                                                           2581.190 0.4935337
6
         79317.75 226.37490
                                  15.670265 12642.176
                                                           8302.325 1.4294626
  cohort_count_ha
                         run
1
                 0 reference
2
                 0 reference
3
                 0 reference
4
                 0 reference
5
                27 reference
6
                49 reference
```

We can see that the output is given for each year and each species in our 1ha area. Plot the living carbon (total_carbon_kg) in time, coloring by species. We give a unified

species coloring in the beginning. Plot number of trees and mean diameter. Carbon and number of trees can be plotted in an additive way, but for mean dbh we do line plot per species.

```
cols.all=c( "rops"="#e0e0e0", "acpl"="#A9A9A9", "alin"="#696969", "alvi"="#2e2e2e",
 "bepe"="#fadfad", "casa"="#7eeadf", "coav"="#20c6b6", "tipl"="#645394",
 "ulgl"="#311432", "saca"="#D8BFD8", "soar"="#DDA0DD", "soau"="#BA55D3",
 "pice"="#D27D2D", "pini"="#a81c07", "alg1"="#2ECBE9", "tico"="#128FC8",
 "potr"="#00468B", "poni"="#5BAEB7", "frex"="#fe9cb5", "cabe"="#fe6181",
 "acps"="#fe223e","lade"="#FFFE71","abal"="#FFD800", "pisy"="#A4DE02",
 "fasy"="#76BA1B", "piab"="#006600", "quro"="#FF7F00", "qupe"="#FF9900",
 "qupu"="#CC9900")
# We set here the order of the run categories for the table, to have the first
#run first and then the second. (left-right of the plots)
landscape$run <- factor(landscape$run, levels = c(name1, name2))</pre>
# Plot the living carbon content, to have tonnes/ha, we divide total carbon kg by 1000.
g1 <-
    ggplot2::ggplot(landscape, aes(year, total_carbon_kg / 1000 ,
                                   fill = factor(species))) +
    ggplot2::geom_area() +
    ggplot2::scale_fill_manual(values = cols.all,
                               guide = guide_legend(reverse = TRUE)) +
    ggplot2::facet_wrap( ~ run, nrow = 1) +
    ggplot2::labs(x = "Year", y = "Living C t/ha", fill = "Species") +
    ggplot2::theme_bw()
g2 <-
    ggplot2::ggplot(landscape, aes(year, count ha , fill = factor(species))) +
    ggplot2::geom_area() +
    ggplot2::scale_fill_manual(values = cols.all,
                               guide = guide_legend(reverse = TRUE)) +
    ggplot2::facet_wrap( ~ run, nrow = 1) +
    ggplot2::labs(x = "Year", y = "Number of trees per ha",
                  fill = "Species") +
    ggplot2::theme_bw()
gridExtra::grid.arrange(g1, g2, ncol = 1)
```





Note that we do not have management intervention, and in the model trees shorter than 4m are not included in these outputs. However as they grow taller than 4m, model start to handle them as individual trees and including in these outputs which we are looking now.

Beech profited from drier and warmer conditions at the expense of spruce. Acer pseudo-platanus and Tilia cordata also slightly increased their proportion in the living C stock and number of trees. The increase of the number of trees indicates the success of the regeneration under reference conditions, however this increase is much slower under the climate change scenario.

Mean dbh of individual species reflected the development of the number of trees. The increase in the number of trees caused the reduction of mean dbh, e.g. of beech under both scenarios between 20 and 35 simulation years. The species-specific graphs for mean dbh indicates that some species do not have successful regeneration under the climate change scenario. Spruce dbh decreased only under reference scenario, when we observed the increase of the number of trees, while under climate change scenario the mean dbh of spruce was growing indicating the growth of remaining trees of the species at the plot.

Calculate the stored C amount in the initial year (year==0) and the last year (year==100)!

```
livingCO <- data.frame(
   landscape |> dplyr::filter(year == 0) |>
```

```
dplyr::group_by(run) |>
        dplyr::summarize(sum.livingC = sum(total_carbon_kg / 1000))
print(livingCO)
               run sum.livingC
                      151.7751
         reference
2 4deg_20percdrier
                      151.7751
livingC100 <- data.frame(</pre>
    landscape |> dplyr::filter(year == 100) |>
        dplyr::group_by(run) |>
        dplyr::summarize(sum.livingC = sum(total_carbon_kg / 1000))
print(livingC100)
               run sum.livingC
         reference
                      416.3909
2 4deg_20percdrier
                      420.1472
```

The initial conditions are same for the runs, but they are ending up at different C levels.

Calculate the stored C amount PER SPECIES in the initial year (year==0) and the last year (year==100)

```
species.livingC0 <-
    data.frame(
        landscape |> dplyr::filter(year == 0) |>
              dplyr::group_by(run, species) |>
              dplyr::summarize(livingC = sum(total_carbon_kg / 1000))
    )
print(species.livingC0)
```

```
run species livingC
1
        reference
                    acps 19.31678
2
        reference
                    fasy 74.97441
3
        reference piab 45.66689
        reference tico 11.81698
5 4deg_20percdrier acps 19.31678
6 4deg_20percdrier
                   fasy 74.97441
7 4deg_20percdrier
                    piab 45.66689
8 4deg_20percdrier
                   tico 11.81698
```

```
species.livingC100 <-
    data.frame(
    landscape |> dplyr::filter(year == 100) |>
        dplyr::group_by(run, species) |>
        dplyr::summarize(livingC = sum(total_carbon_kg / 1000))
)
print(species.livingC100)
```

```
run species
                             livingC
1
                     acps 25.042637
        reference
2
        reference
                     fasy 164.555979
3
        reference
                     piab 221.267535
        reference
                     tico
                            5.524713
5 4deg_20percdrier
                     acps 65.167214
6 4deg_20percdrier
                     fasy 243.374438
7 4deg_20percdrier
                     piab 92.649456
8 4deg_20percdrier
                     tico 18.956110
```

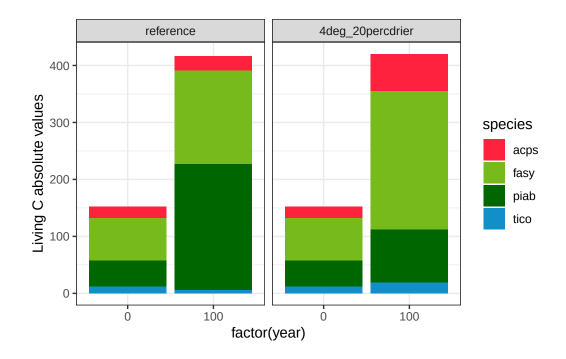
Calculate the species proportions based on the stored C amount in the initial year (year==0) and the last year (year==100) For this we need the total C amount and the species-specific C amount.

```
run species livingC sum.livingC spec.prop year
                                    151.7751 0.1272724
1
        reference
                     acps 19.31678
2
        reference
                     fasy 74.97441
                                     151.7751 0.4939837
                                                           0
3
        reference piab 45.66689
                                     151.7751 0.3008854
                                                           0
        reference
                    tico 11.81698
                                     151.7751 0.0778585
                                                           0
5 4deg 20percdrier
                   acps 19.31678
                                     151.7751 0.1272724
                                                           0
6 4deg_20percdrier
                   fasy 74.97441
                                     151.7751 0.4939837
                                                           0
7 4deg_20percdrier
                                     151.7751 0.3008854
                   piab 45.66689
                                                           0
8 4deg_20percdrier
                     tico 11.81698
                                     151.7751 0.0778585
                                                           0
```

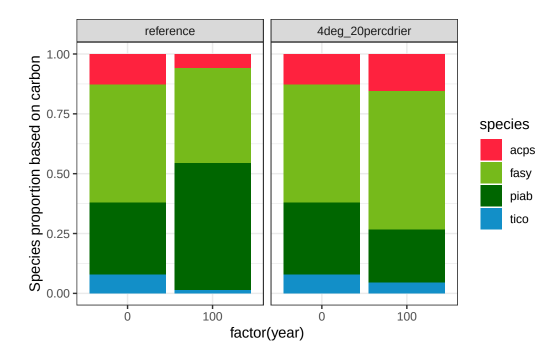
```
LC100 <-
    dplyr::left_join(species.livingC100, livingC100, by = "run")
LC100 <-</pre>
```

```
livingC sum.livingC spec.prop year
             run species
1
       reference acps 25.042637
                                    416.3909 0.06014214 100
       reference fasy 164.555979
                                    416.3909 0.39519594
        reference piab 221.267535
                                    416.3909 0.53139383
3
                                                        100
4
        reference tico
                          5.524713
                                    416.3909 0.01326809
                                                        100
5 4deg_20percdrier acps 65.167214
                                     420.1472 0.15510567
                                                        100
6 4deg_20percdrier fasy 243.374438
                                    420.1472 0.57925991
                                                        100
7 4deg_20percdrier piab 92.649456
                                     420.1472 0.22051665
                                                        100
8 4deg_20percdrier
                   tico 18.956110
                                     420.1472 0.04511778 100
```

Put the two tables together and visualize the results!



```
ggplot2::ggplot(LC, aes(fill = species, y = spec.prop, x = factor(year))) +
    ggplot2::geom_bar(position = "stack", stat = "identity") +
    ggplot2::scale_fill_manual(values = cols.all) +
    ggplot2::facet_wrap( ~ run) +
    ggplot2::ylab("Species proportion based on carbon") +
    ggplot2::theme_bw()
```



The living C stock more than doubled over 100 years due to the accumulation of biomass in trees and no management interventions. Moreover, the proportion of individual tree species changed depending on the driving climatic conditions. Under reference conditions the proportion of spruce has grown, under climate change the proportion of beech.

2.2.12 Question A - Group 4

• How are biodiversity indices changing in time and across the simulated scenario(s) on Plot 4?

Run the model as it is described in the previous chapters to have at least 2 simulations completed: one with reference conditions and one scenario with some changes in the environment. Check if you have the output files in the output folder: <code>iLand_simulations/output/</code>. If you manage to simulate 2 scenarios, you can expand the code to have file3, and 3rd from each variable, or make the comparison in 2 step always comparing one scenario to the reference conditions at the time.

Open the *summerSchoolExerciseProcessBased.Rproj* project in R studio and start working there.

Read in the *tree* output table from two outputs you have for your plot:

```
path1 <- "model/iLand_simulations/output/Output_plot4.sqlite"</pre>
path2 <-
    "model/iLand simulations/output/Output_plot4 4deg_20percdrier.sqlite"
file1 <- here::here(path1)</pre>
file2 <- here::here(path2)</pre>
# Give some name for the three simulations.
name1 <- "reference"</pre>
name2 <- "4deg_20percdrier"</pre>
# Read in data using the RSQLite package
sqlite.driver <- RSQLite::dbDriver("SQLite")</pre>
db1 <-
    RSQLite::dbConnect(sqlite.driver, dbname = file1) # connect to the file1
tables.in.the.file <-
    RSQLite::dbListTables(db1) # explore the tables in the file1
print(tables.in.the.file)
[1] "carbon"
                         "carbonflow"
                                              "dynamicstand"
[4] "landscape"
                         "landscape_removed" "runinfo"
[7] "stand"
                         "tree"
# We will work with "tree" table and tree-scale data:
tree1 <- RSQLite::dbReadTable(db1, "tree")</pre>
RSQLite::dbDisconnect(db1) # disconnect to the file1
# READ IN DATA FROM THE SECOND FILE: ----
db2 <-
    RSQLite::dbConnect(sqlite.driver, dbname = file2) # connect to the file2
tree2 <- RSQLite::dbReadTable(db2, "tree")</pre>
RSQLite::dbDisconnect(db2)
```

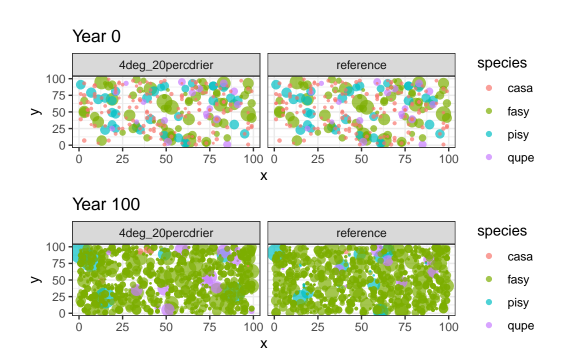
Merge the data from the two files together and make a column which tells which comes from which simulation, and study the table. The column "run" will support the plotting where we can use facets in ggplot.

head(tree)

```
year ru rid species id x y
                                     dbh
                                           height basalArea volume_m3 age
     0
        0
                 fasy
                       1 49 65 98.48459 55.96636 0.7617745
                                                              18.46041 699
1
2
     0
        0
                       2 51 15 94.27415 53.57367 0.6980316
                                                              16.19252 669
3
                       3 97 47 94.07941 53.46301 0.6951508
     0
        0
            1
                                                              16.09238 668
4
     0
        0
                 fasy 4 63
                             7 91.67390 52.09602 0.6600568 14.88928 651
            1
5
     0
        0
                 fasy 5 53 57 90.70000 51.54258 0.6461070
                                                             14.41977 644
            1
6
     0
        0
            1
                 fasy 6 13 95 89.37351 50.78876 0.6273466 13.79631 634
  leafArea m2 foliageMass stemMass branchMass fineRootMass coarseRootMass
     584.4382
                 53.13074 6802.456
                                      845.6093
                                                   39.84806
                                                                   758.3801
1
2
     542.6002
                 49.32729 6165.430
                                      764.7609
                                                   36.99547
                                                                   685.2726
3
     540.6961
                 49.15419 6136.807
                                      761.1324
                                                   36.86565
                                                                   681.9930
4
     517.4045
                 47.03678 5789.331
                                      717.1134
                                                   35.27758
                                                                   642.2182
5
     508.0949
                 46.19045 5651.843
                                      699.7122
                                                   34.64284
                                                                   626.5004
                 45.04793 5467.530
                                      676.3990
     495.5272
                                                   33.78595
                                                                   605.4482
        lri lightResponse stressIndex reserve_kg treeFlags
                                                                   run
1 0.6396535
                         0
                                     0
                                         92.97880
                                                           0 reference
2 0.4253728
                        0
                                     0
                                         86.32275
                                                           0 reference
3 1.0000000
                        0
                                     0
                                         86.01984
                                                           0 reference
4 0.4545080
                         0
                                     0
                                         82.31436
                                                           0 reference
5 0.6055760
                         0
                                         80.83328
                                                           0 reference
                                     0
6 1.0000000
                         0
                                         78.83388
                                                           0 reference
```

To explore a bit the simulation, plot the tree locations at the beginning and the end of the simulation! For plotting we use the colors for the species, and the size of a circles to show dbh differences. Here we divided dbh with the value 20 just for visualization, not to have too huge circles shading each others completely, but still see the trees.

```
ggplot2::ggtitle("Year 100") +
    ggplot2::facet_wrap(~ run) + ggplot2::theme_bw()
gridExtra::grid.arrange(g1, g2, ncol = 1)
```

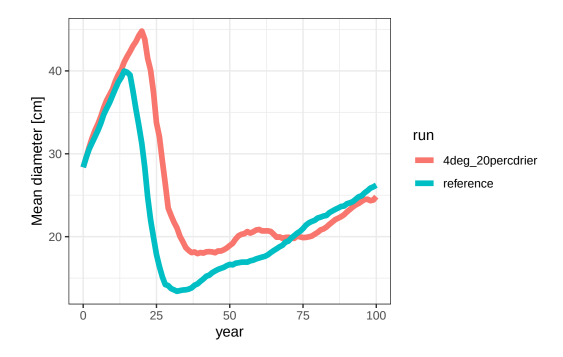


We can see that the species composition is somewhat changed and the forest become more dense. We can even identify the same trees at the same location.

Visualize some changes in time, for example the mean diameter of the trees!

```
sum.table <- tree |> dplyr::group_by(year, run) |>
    dplyr::summarise(
        N = dplyr::n(),
        MD = mean(dbh, na.rm = TRUE), #mean diameter
        BA = sum(basalArea)#basal area
)

ggplot2::ggplot(sum.table, ggplot2::aes(x = year, y = MD, color = run)) +
    ggplot2::geom_line(lwd = 2) +
    ggplot2::ylab("Mean diameter [cm]") + ggplot2::theme_bw()
```



We can see that there is an initial increase in mean dbh, then a drop after 20/25 years. This is due to the growing regeneration layer that is produced by the seeds of the existing trees. In the initial year we do not have regeneration layer in the simulations (it is possible to put there, but we do not have it now). And until the small trees grow up to higher than 4 m they are not appearing in the individual *tree* output as they are handled in cohorts.

To study biodiversity aspects, we can calculate biodiversity indicators. Let's use the adiv package for species diversity. First we need to change the database structure to have the number of trees for each species in different columns (pivot_wider) without any additional columns for the adiv package speciesdiv function. We work first only with tree data from one simulation (tree1). You can find the documentation of the adiv package here: "documents/adiv.pdf"

```
# First we need to change the
tlong1 <- tree1 |>
    dplyr::group_by(year, species) |> dplyr::summarise(N = dplyr::n()) |>
    tidyr::pivot_wider(
        id_cols = 'year',
        names_from = 'species',
        values_from = 'N',
        values_fill = 0
    ) |>
    dplyr::ungroup() |>
```

```
dplyr::select(-year)
head(tlong1)
# A tibble: 6 x 4
   casa fasy pisy qupe
  <int> <int> <int> <int>
           70
                  33
1
    152
                        18
2
    147
           70
                  32
                        18
3
    143
                  32
                        18
           69
4
    139
           68
                  30
                        18
5
    139
           68
                  30
                        18
6
    137
            68
                  30
                        18
```

Then we apply the "speciesdiv" function and we add back the year and run columns to have the information.

```
div1 <- data.frame(adiv::speciesdiv(tlong1)) |>
     dplyr::mutate(year = unique(tree1$year), run = name1)
head(div1)
```

```
richness GiniSimpson Simpson Shannon Margalef Menhinick McIntosh year
             0.6052946 2.533535 1.109706 0.5348097 0.2420910 0.3956925
1
2
             0.6092385\ 2.559105\ 1.115644\ 0.5369369\ 0.2447960\ 0.3993293
                                                                           1
3
         4
             0.6131053 2.584683 1.122656 0.5387598 0.2471208 0.4028815
                                                                           2
             0.6129335 2.583535 1.122122 0.5413928 0.2504897 0.4030962
                                                                           3
5
             0.6129335 2.583535 1.122122 0.5413928 0.2504897 0.4030962
                                                                           4
             0.6154135 2.600195 1.126170 0.5421632 0.2514778 0.4053326
6
         4
        run
1 reference
2 reference
3 reference
4 reference
5 reference
6 reference
```

We make the same steps for the other simulation results (tree2).

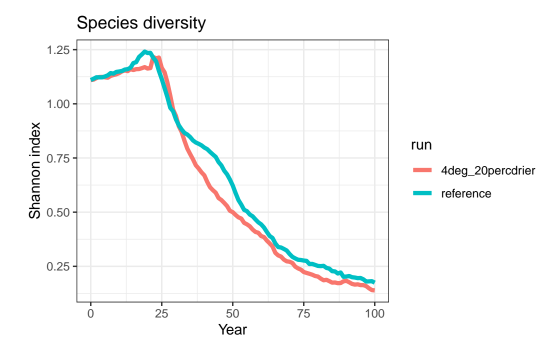
```
tlong2 <- tree2 |>
   dplyr::group_by(year, species) |> dplyr::summarise(N = n()) |>
   tidyr::pivot_wider(
```

```
id_cols = 'year',
    names_from = 'species',
    values_from = 'N',
    values_fill = 0
) |>
    dplyr::ungroup() |>
    dplyr::select(-year)

div2 <- data.frame(adiv::speciesdiv(tlong2)) |>
    dplyr::mutate(year = unique(tree2$year), run = name2)
```

We merge them all together and make a plot based on Shannon index.

```
tlong2 <- tree2 |>
    dplyr::group_by(year, species) |> dplyr::summarise(N = n()) |>
    tidyr::pivot_wider(
        id_cols = 'year',
        names_from = 'species',
        values_from = 'N',
       values_fill = 0
    ) |>
    dplyr::ungroup() |>
    dplyr::select(-year)
div2 <- data.frame(adiv::speciesdiv(tlong2)) |>
    dplyr::mutate(year = unique(tree2$year), run = name2)
div <- rbind(div1, div2)</pre>
ggplot2::ggplot(div, aes(year, Shannon , color = run)) +
    ggplot2::geom_line(lwd = 1.5) +
    ggplot2::ggtitle("Species diversity") +
    ggplot2::labs(x = "Year", y = "Shannon index") +
    ggplot2::theme_bw()
```



We see that the values of Shannon index are first stable and then tend to decrease, while the reduction under the 4deg_20percdrier is more substantial than under the reference scenario. Shannon index is calculated as follows:

$$S = -\sum \left(p \ i \ *log(p \ i \) \right)$$

Where p i is the relative abundance of the species i, calculated as the ratio of the abundance of the species i and the abundance of all species. Shannon-index is 0 when we have only one species at the stand.

We can calculate some additional biodiversity indices targeting spatial diversity using the *treespat* package developed by Francesco Chianucci (fchianucci@gmail.com). You can find a short description of the package here: "documents/treespat_package.pdf"

You can install the package using devtools:

devtools::install_gitlab('fchianucci/treespat')

Then, we are ready to use it! Let's calculate these two indices:

• Diameter differentiation (Gadow, 1993): Spatial size inequality defined as the mean of the ratio of smaller and larger plant sizes in the nearest neighbors of a tree. The value of the index increases with increasing average size difference between neighboring trees. 0 is implying that neighboring trees have equal size.

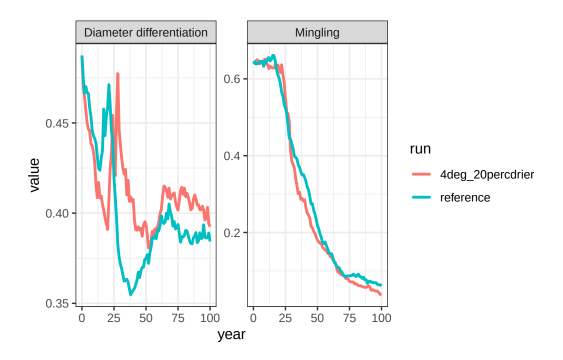
• Mingling (Aguirre et al., 2003): One very intuitive extension of taxonomic species diversity (either richness or abundance) is considering spatial mingling, namely how plants of the same (con-specific neighbors) or different (hetero-specific neighbors) species are arranged in space. The mingling index calculates the proportion of the k nearest neighbors that do not belong to the same species as the reference tree. For example, with four neighbors, the mingling attribute can assume five values, ranging from 0 (all trees are of the same species) to 1 (all trees belong to different species).

Calculate the indices based on the reference run, then based on the scenario. For each index we add extra columns with the index name, and run name for further analyses. The max.k parameter is telling how many neighboring trees we want to account for.

```
# diameter differentiation:
# mingling
diff <- data.frame(treespat::DIFF(tree1, .x = x, .y = y,.mark = dbh, xmax = 100,
                 ymax = 100, max.k = 4, shape='square', .groups=c('year')) |>
         dplyr::mutate(index="DIFF", name="Diameter differentiation", run=name1))
head(diff)
 year
            DIFF index
                                           name
                                                      run
     0 0.4874231 DIFF Diameter differentiation reference
1
2
     1 0.4752327 DIFF Diameter differentiation reference
     2 0.4665189 DIFF Diameter differentiation reference
     3 0.4701394 DIFF Diameter differentiation reference
5
     4 0.4668349 DIFF Diameter differentiation reference
     5 0.4663594 DIFF Diameter differentiation reference
ming <- data.frame(treespat::MING(tree1, .x = x, .y = y, .species = species,
        xmax = 100, ymax = 100, max.k = 4, shape='square', .groups=c('year')) |>
        dplyr::mutate(index="MING", name="Mingling", run=name1))
indices1<-rbind(diff |> dplyr::rename(value=DIFF),
ming |> dplyr::rename(value=MINGLING ) )
```

We calculated each index as separate variable and then merged them into indices1. Here I renamed the column names to have it all "value", for easier plotting later on. (each line has the information on which run and which index is the value) We do the same for the scenario simulation, we merge them together and make the plotting.

```
diff <- data.frame(treespat::DIFF(tree2, .x = x, .y = y,.mark = dbh,</pre>
      xmax = 100, ymax = 100, max.k = 4, shape = 'square',
      .groups = c('year')) |>
      dplyr::mutate(index = "DIFF", name = "Diameter differentiation",
                    run = name2))
ming <- data.frame(treespat::MING(tree2, .x = x, .y = y, .species = species,
       xmax = 100, ymax = 100, max.k = 4, shape = 'square',
       .groups = c('year')) |>
        dplyr::mutate(index = "MING", name = "Mingling", run = name2))
indices2 <- rbind(diff |> dplyr::rename(value = DIFF),
                  ming |> dplyr::rename(value = MINGLING))
indices <- rbind(indices1, indices2)</pre>
ggplot2::ggplot(indices, aes(x = year, y = value, color = run)) +
    ggplot2::geom_line(lwd = 1) +
    ggplot2::facet_wrap( ~ name, ncol = 2, scales = "free") +
    ggplot2::theme_bw()
```



The diameter differentiation increases with the increasing average difference in diameter between neighboring trees. The value 0 indicates that neighboring trees have equal

diameters. The development shows that the index values decrease during the first 20/25 years, after which it steeply increases due to the occurrence of ingrowth and then the values are stabilized. The values for the reference climate scenario are slightly higher than for the climate change scenario indicating slower growth under drier and warmer conditions and/or the differences in tree species composition (proportion of tree species). The mingling index is studying the neighboring trees regarding their species. In both development there is a decrease in the index after the occurrence of regeneration. Using the climate change scenario the index shows steeper decrease.

2.2.13 Question B - Group 4

• How are the species distribution and total living biomass C content changing in time on Plot 4? Compare 0 year and 100 year status in the reference case and in the case of your scenario(s)!

Run the model as it is described in the previous chapters to have at least 2 simulations completed: one with reference conditions and one scenario with some changes in the environment. Check if you have the output files in the output folder: <code>iLand_simulations/output/</code>. If you manage to simulate 2 scenarios, you can expand the code to have file3, and 3rd from each variable, or make the comparison in 2 step always comparing one scenario to the reference conditions at the time. Open the <code>summerSchoolExercise-ProcessBased.Rproj</code> project in R studio and start working there.

Read in the *landscape* output table from two outputs you have for your plot:

```
RSQLite::dbDisconnect(db1) # disconnect to the file1

db2 <-
    RSQLite::dbConnect(sqlite.driver, dbname = file2) # connect to the file2
landscape2 <- RSQLite::dbReadTable(db2, "landscape")
RSQLite::dbDisconnect(db2)</pre>
```

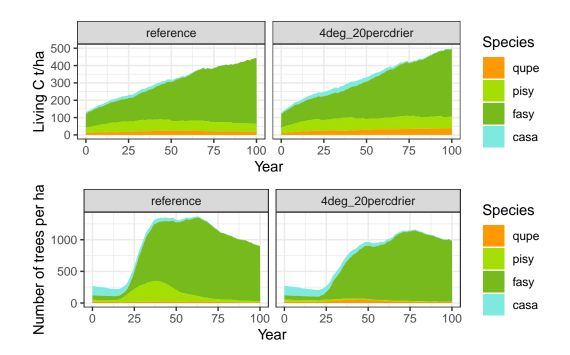
Merge the data from the three files together and make a column which tells which comes from which simulation, and study the table. The column "run" will support the plotting where we can use facets in ggplot.

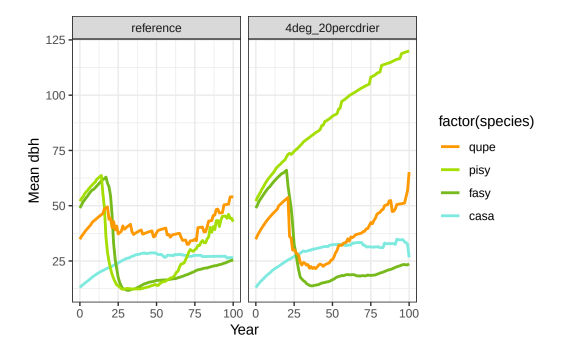
```
year area area_100m species count_ha dbh_avg_cm height_avg_m
                                                                     volume_m3
     0
          1
                     1
                                      152
                                            12.96761
                                                          10.77155
                                                                      9.465222
1
                           casa
2
     0
          1
                                            48.97035
                                                          27.82864 260.513423
                     1
                           fasy
                                       70
3
     0
          1
                     1
                                       33
                                            52.05737
                                                          29.57999
                                                                     91.870356
                           pisy
4
     0
          1
                                            34.83232
                     1
                           qupe
                                       18
                                                          17.83572
                                                                     15.683124
5
     1
           1
                     1
                           casa
                                      147
                                            13.72713
                                                          11.31040
                                                                     10.643544
     1
          1
                                       70
                                            50.20535
                                                          28.46379 268.968996
6
                           fasy
  total_carbon_kg
                       gwl_m3 basal_area_m2
                                                 NPP_kg NPPabove_kg
                                                                            LAI
         8041.268
                     9.465222
                                    2.099428
                                                  0.000
                                                                 0.00 0.3009711
1
2
        80391.775 260.513423
                                   15.847768
                                                  0.000
                                                                 0.00 1.3974759
3
        30694.307
                    91.870356
                                    7.063300
                                                  0.000
                                                                 0.00 0.8112550
4
        10327.265
                    15.683124
                                    1.791511
                                                   0.000
                                                                 0.00 0.1830485
5
         8550.914
                    10.987314
                                    2.266547
                                               2834.498
                                                             1765.12 0.3118849
6
        84739.791 268.968996
                                   16.383716 13024.190
                                                             8553.20 1.4546398
  cohort_count_ha
                          run
1
                 0 reference
2
                 0 reference
3
                 0 reference
4
                 0 reference
5
                 0 reference
               229 reference
```

We can see that the output is given for each year and each species in our 1ha area. Plot the living carbon (total_carbon_kg) in time, coloring by species. We give a unified species coloring in the beginning. Plot number of trees and mean diameter. Carbon and

number of trees can be plotted in an additive way, but for mean dbh we do line plot per species.

```
cols.all=c( "rops"="#e0e0e0", "acpl"="#A9A9A9", "alin"="#696969", "alvi"="#2e2e2e",
 "bepe"="#fadfad","casa"="#7eeadf","coav"="#20c6b6","tip1"="#645394",
 "ulgl"="#311432","saca"="#D8BFD8","soar"="#DDA0DD","soau"="#BA55D3",
 "pice"="#D27D2D", "pini"="#a81c07", "alg1"="#2ECBE9", "tico"="#128FC8",
 "potr"="#00468B", "poni"="#5BAEB7", "frex"="#fe9cb5", "cabe"="#fe6181",
 "acps"="#fe223e","lade"="#FFFE71","abal"="#FFD800", "pisy"="#A4DE02",
 "fasy"="#76BA1B", "piab"="#006600", "quro"="#FF7F00", "qupe"="#FF9900",
 "qupu"="#CC9900")
# We set here the order of the run categories for the table, to have the first
#run first and then the second. (left-right of the plots)
landscape$run <- factor(landscape$run, levels = c(name1, name2))</pre>
# Plot the living carbon content, to have tonnes/ha, we divide total_carbon kg by 1000.
g1 <-
    ggplot2::ggplot(landscape, aes(year, total_carbon_kg / 1000 , fill = factor(species)
    ggplot2::geom_area() +
    ggplot2::scale_fill_manual(values = cols.all, guide = guide_legend(reverse =
                                                                            TRUE)) +
    ggplot2::facet_wrap( ~ run, nrow = 1) +
    ggplot2::labs(x = "Year", y = "Living C t/ha", fill = "Species") +
    ggplot2::theme_bw()
g2 <-
    ggplot2::ggplot(landscape, aes(year, count_ha , fill = factor(species))) +
    ggplot2::geom_area() +
    ggplot2::scale fill manual(values = cols.all, guide = guide legend(reverse =
                                                                            TRUE)) +
    ggplot2::facet_wrap( ~ run, nrow = 1) +
    ggplot2::labs(x = "Year", y = "Number of trees per ha", fill = "Species") +
    ggplot2::theme_bw()
gridExtra::grid.arrange(g1, g2, ncol = 1)
```





Note that we do not have management intervention, and in the model trees shorter than 4m are not included in these outputs. However as they grow taller than 4m, model start to handle them as individual trees and including in these outputs which we are looking now.

At the end of the simulated 100 years, the total simulated carbon stock in living biomass under climate change scenario was slightly higher than under the reference climate. The proportion of beech has increased under both simulations. The number of trees increased in both cases after circa 25 years, but the increase was steeper under reference conditions. This increase caused by the regeneration.

Mean dbh of individual species reflected the development of the number of trees. The increase in the number of trees caused the reduction of mean dbh, e.g. of beech under both scenarios between 20 and 35 simulation years. The species-specific graphs for mean dbh indicates that some species do not have successful regeneration under the climate change scenario. Pine dbh decreased only under reference scenario, when we observed the increase of the number of trees, while under climate change scenario the mean dbh of pine was growing indicating the growth of remaining trees of the species at the plot. The small proportion of chestnut almost decrease to 0 under both scenarios, its dbh development does not show any successful regeneration processes.

Calculate the stored C amount in the initial year (year==0) and the last year (year==100)!

```
livingCO <- data.frame(</pre>
    landscape |> dplyr::filter(year == 0) |>
        dplyr::group_by(run) |>
        dplyr::summarize(sum.livingC = sum(total_carbon_kg / 1000))
)
print(livingCO)
               run sum.livingC
                      129.4546
1
         reference
2 4deg_20percdrier
                      129.4546
livingC100 <- data.frame(</pre>
    landscape |> dplyr::filter(year == 100) |>
        dplyr::group_by(run) |>
        dplyr::summarize(sum.livingC = sum(total_carbon_kg / 1000))
print(livingC100)
               run sum.livingC
                      441.6930
         reference
2 4deg_20percdrier
                      499.1578
```

The initial conditions are same for the runs, but they are ending up at different C levels.

Calculate the stored C amount PER SPECIES in the initial year (year==0) and the last year (year==100)

```
species.livingC0 <- data.frame(
    landscape |> dplyr::filter(year == 0) |>
        dplyr::group_by(run, species) |>
        dplyr::summarize(livingC = sum(total_carbon_kg / 1000))
)
print(species.livingC0)
```

```
run species livingC
1 reference casa 8.041268
2 reference fasy 80.391775
3 reference pisy 30.694307
4 reference qupe 10.327265
5 4deg_20percdrier casa 8.041268
```

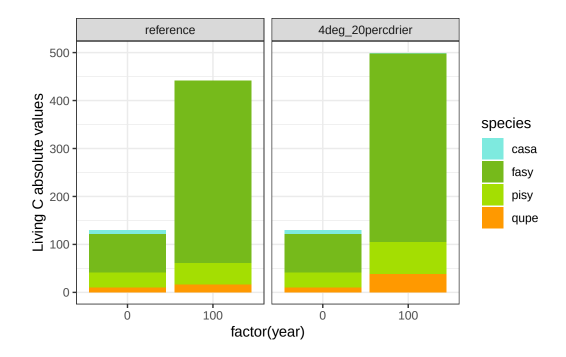
```
run species
                               livingC
1
                             0.2319933
         reference
                      casa
2
         reference
                      fasy 380.6997702
3
         reference
                      pisy
                            44.2202121
4
         reference
                      qupe
                           16.5410152
5 4deg_20percdrier
                      casa
                             1.5847763
6 4deg_20percdrier
                      fasy 392.4993454
7 4deg_20percdrier
                      pisy
                            66.8578057
8 4deg_20percdrier
                      qupe
                            38.2158285
```

Calculate the species proportions based on the stored C amount in the initial year (year==0) and the last year (year==100) For this we need the total C amount and the species-specific C amount.

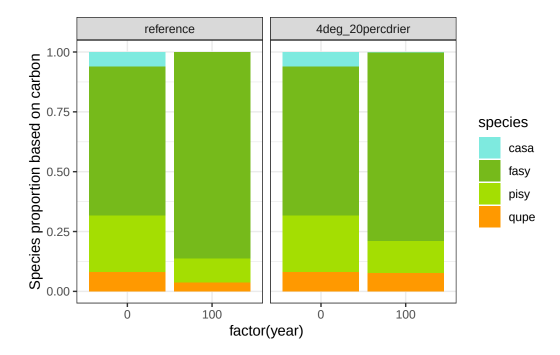
```
livingC sum.livingC spec.prop year
               run species
1
         reference
                      casa 8.041268
                                         129.4546 0.06211650
                                                                0
2
         reference
                      fasy 80.391775
                                         129.4546 0.62100355
                                                                0
3
         reference
                      pisy 30.694307
                                         129.4546 0.23710477
                                                                0
         reference
                      qupe 10.327265
                                         129.4546 0.07977518
                                                                0
5 4deg_20percdrier
                                                                0
                      casa 8.041268
                                         129.4546 0.06211650
                                        129.4546 0.62100355
6 4deg_20percdrier
                      fasy 80.391775
                                                                0
7 4deg_20percdrier
                      pisy 30.694307
                                        129.4546 0.23710477
                                                                0
8 4deg_20percdrier
                      qupe 10.327265
                                        129.4546 0.07977518
```

```
run species
                             livingC sum.livingC
                                                   spec.prop year
                                       441.6930 0.0005252366
1
        reference
                    casa
                           0.2319933
                                                             100
2
        reference
                    fasy 380.6997702
                                       441.6930 0.8619103723
                                                             100
3
        reference pisy 44.2202121
                                       441.6930 0.1001152679
                                                             100
4
                                       441.6930 0.0374491232
                                                             100
        reference qupe 16.5410152
5 4deg_20percdrier
                    casa 1.5847763
                                     499.1578 0.0031749006
                                                             100
6 4deg_20percdrier
                    fasy 392.4993454
                                       499.1578 0.7863232430
                                                             100
7 4deg_20percdrier
                    pisy 66.8578057
                                       499.1578 0.1339412339
                                                             100
8 4deg_20percdrier
                                       499.1578 0.0765606224
                                                             100
                    qupe 38.2158285
```

Put the two tables together and visualize the results!



```
ggplot2::ggplot(LC, aes(fill = species, y = spec.prop, x = factor(year))) +
    ggplot2::geom_bar(position = "stack", stat = "identity") +
    ggplot2::scale_fill_manual(values = cols.all) +
    ggplot2::facet_wrap( ~ run) +
    ggplot2::ylab("Species proportion based on carbon") +
    ggplot2::theme_bw()
```



The living C stock tripled over 100 years due to the accumulation of biomass in trees and no management interventions and reached higher level under the climate change scenario. Both cases the proportion of beech increased.

In the empirical modelling we will cover two examples of creating two models based on observed data. We will use two approaches Generalized Linear Models (GLMs) and Boosted Regression trees (BRTs). Please take all the results with a grain of salt, we are making very generalized statements from a limited data set, and we are not getting into the details of how each of the models should be assessed; the goal here is that you learn how observed data can be used to create models that help you to understand the data and relationships better.

3.1 Data

We will work with one dataset derived from the inventory developed to assess forest structure and deadwood properties of six representative forest areas in the Czech Republic (Hošek, n.d.). This dataset collects observations of the presence /absence of certain species of biodiversity importance across 99 plots.

The data was collected from square sampling plots (2500 m2 each) in six forested Czech Republic regions. These regions are representative of the main bio-regions and elevation range of forests, considering their importance in territorial representation, forestry, and ecology. The inventory sampled for biodiversity variables such as the presence/absence of different species of birds, Tracheophyta, bryophytes, fungi, lichens, and beetles.

The dataset has been generously provided by Jeňýk Hofmeister at the Czech University of Life Sciences Prague (jenyk.hofmeister@email.cz) to be used in the context of this exercise. It should not be used for other purposes or be further distributed. If you want to use this data or learn more about it, please get in touch with Jeňýk Hofmeisterhere: jenyk.hofmeister@email.cz.

The data you can download for this exercise is an aggregated summary of the original data, some aspects have been modified from the observed data, so the results you will obtain will only partially match the observed reality. This data is very similar in structure and observed variables to the one you collected in this summer school. The idea is to move from data collection, analysis to this part, where you use the data to create a model.

3.1.1 Data description

These are the variables available in the data

- longitud: longitude of the plot location
- latitude: latitude of the plot location
- forestManagementType: forest management type applied in this plot
- forestStructure: current forest structure in the plot
- slope: slope of the plot
- A.pseudoplatanus: proportion of this tree species in the plot in volume
- F.sylvatica: proportion of this tree species in the plot in volume
- L.decidua: proportion of this tree species in the plot in volume
- Q.robur: proportion of this tree species in the plot in volume
- S.aucuparia: proportion of this tree species in the plot in volume
- B.pendula: proportion of this tree species in the plot in volume
- P.abies: proportion of this tree species in the plot in volume
- P.sylvestris: proportion of this tree species in the plot in volume
- F.excelsior: proportion of this tree species in the plot in volume
- A.alba: proportion of this tree species in the plot in volume
- A.platanoides: proportion of this tree species in the plot in volume
- T.cordata: proportion of this tree species in the plot in volume
- S.racemosa: proportion of this tree species in the plot in volume
- U.glabra: proportion of this tree species in the plot in volume
- S.nigra: proportion of this tree species in the plot in volume
- P.alba: proportion of this tree species in the plot in volume
- U.minor: proportion of this tree species in the plot in volume
- S.caprea: proportion of this tree species in the plot in volume
- C.betulus: proportion of this tree species in the plot in volume
- P.nigra: proportion of this tree species in the plot in volume
- Q.petraea: proportion of this tree species in the plot in volume
- S.torminalis: proportion of this tree species in the plot in volume
- A.campestre: proportion of this tree species in the plot in volume
- P.strobus: proportion of this tree species in the plot in volume
- Q.rubra: proportion of this tree species in the plot in volume
- volAllha: total volume in the plot
- GiniDBH: Gini index calculated to assess the forest structural diversity in diameter sizes, higher values indicate more structural heterogeneity; lower values indicate more homogeneous stands
- ShannonIndexTreeSpp: The Shannon index is way to measure the diversity of tree species in the plot
- Tracheophyta_rich: Species richness across Tracheophyta species
- Birds_rich: Species richness across Birds species
- Bryophytes_rich: Species richness across Bryophytes species
- Fungi_rich: Species richness across Fungi species

- Lichens_rich: Species richness across Lichens species
- Beetles_rich: Species richness across Beetles species
- dendrocoposMajor: presence / absence of Dendrocopos major
- certhia: presence / absence of Certhia familiaris
- bryophitaNumObs: number of observed Bryophytes species
- birdNumObs: number of observed Bird species
- PlotID: ID number for the plot

3.2 Species description

3.2.1 Species 1 - Bryophytes

Bryophytes constitute an important and permanent component of the forest flora and diversity. They colonize various substrates, which are unsuitable for vascular plants, because of low light intensity or low nutrient level, such as deadwood, bark, rocks, and open soil. They provide shelter habitats, food, and nest material for many animals.

In forests, different ecological guilds of Bryophytes can be distinguished by the substrate on which they are growing, including terricolous, lignicolous, corticolous and saxicolous species that occur on soil, deadwood, bark of living trees and shrubs, or rocks, respectively. As diversity and quality of these substrates is affected by forest management, Bryophytes are suitable indicators for the effect of management on forest conditions. Especially typical woodland Bryophytes, which are strictly depending on forest conditions. It is interesting to better understand the relation of forest management effects on Bryophytes and some studies have already demonstrated their sensitivity to management practices.

3.2.2 Species 2 - Great spotted woodpecker

The great spotted woodpecker (*Dendrocopos major*) is a medium-sized woodpecker with pied black and white plumage and a red patch on the lower belly. It is found in a wide variety of woodlands, broadleaf, coniferous or mixed forests. The great spotted woodpecker spends much of its time climbing trees. It a quite generalist bird species.

3.2.3 Species 3 - Eurasian treecreeper

The Eurasian treecreeper or common treecreeper (*Certhia familiaris*) is a small passerine bird. It prefers mature trees, and in most of Europe, it tends to be found mainly in coniferous forest, especially spruce and fir.

3.3 Models

3.3.1 Generalized Linear Models (GLMs)

We propose to use a generalized linear model (GLM) to understand the abundance of different Bryophytes species in the 99 plots. Count data often conform to a Poisson distribution; in this case, we have a count of the number of species recorded at each plot.

Fitting a Poisson GLM in R is similar to analyzing covariance (or linear model), except that we now need to use the glm function. To run a GLM, we need to provide one extra piece of information beyond that required for a linear model: the family of models we want to use. In this case, we want a Poisson family, family=poisson.

3.3.2 Boosted regression trees (BRTs)

Boosted regression trees (BRT) are a combination of two powerful statistical techniques: boosting and regression trees. Boosting is a machine learning technique similar to model averaging, where the results of several competing models are merged. Unlike model averaging, however, boosting uses a forward, stage-wise procedure, where tree models are fitted iteratively to a subset of the training data. Subsets of the training data used at each iteration of the model fit are randomly selected without replacement, where the proportion of the training data used is determined by the modeler, this is defined with the "bag fraction" parameter. This procedure, known as stochastic gradient boosting, introduces an element of stochasticity that improves model accuracy and reduces overfitting (Elith, Leathwick, and Hastie 2008).

The BRT model calibration is defined by four parameters:

- the learning rate (or shrinkage parameter): The learning rate determines the contribution of each new tree to the growing model, and it is always substantially lower than 1, higher values being related to faster learning.
- the bag fraction: The bag fraction provides in-formation on which fraction of the entire data should be drawn randomly to fit the new tree. This parameter includes a random probabilistic component, making each run model different, and is aimed at improving model accuracy, speed of model creation, and the reduction of overfitting (Friedman, Hastie, and Tibshirani, n.d.).
- the tree complexity: Tree complexity controls the number of fitted interactions among variables, and determines the number of splits in each tree; for example, a value of 1 will present only one split, meaning that the model does not consider interactions; a value of 2 will result in two splits, and two interactions.

• The number of trees required for optimal prediction: The optimal number of trees is selected based on the three previous parameters. The values fitted by the final model are computed as the sum of all of the predictions of the trees, multiplied by their respective learning rates.

3.4 The exercise

3.4.1 The project folder

All the project is available in a github repository. You can download the project with this document, code, the .rproj and the correct folder structure in here [2].

[2] https://github.com/oldiya/summerSchoolExercise

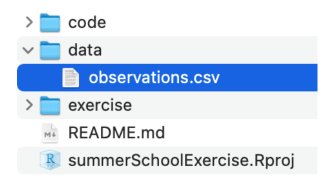
In this project you will find a folder called **code** where the codes for each question are stored. A folder called **data** is where you have to store the data that you have downloaded. A folder called **exercise** contains this document and all the required files to build it.

3.4.2 Download the data

You can download the data in here [1]

[1] https://polybox.ethz.ch/index.php/s/qERYKjzmFTr81Sq

Our recommendation is that you follow this folder structure and you put the data in the folder data of the project and the .rproj into the main folder. If you decide to organize things in a different way then you will have to change the path to the code provided to load the data in the section "Explore the data"



3.4.3 Explore the data

You can load the data in R by doing:

```
observations <- read.csv(here::here("data/observations.csv"))
```

Once you have obtained the data, your next step is to explore it. It is important to carefully analyze the structure of the data and understand its meaning. Each variable must be examined closely, along with the data structure. Remember that each row in this dataset represents one plot, identified by its PlotID, and each column represents one variable.

You can explore each of the variables by doing:

```
str(observations)
```

```
'data.frame':
               99 obs. of 45 variables:
                      : int 1 2 3 4 5 6 7 8 9 10 ...
$ X
$ longitud
                      : num 13.5 13.5 13.5 13.6 13.6 ...
$ latitude
                      : num 49.5 49.5 49.5 49.5 ...
$ forestManagementType: chr "simple clearcutting" "simple clearcutting" "simple clearcutting"
                     : chr "even-aged" "even-aged" "even-aged" "even-aged" ...
$ forestStructure
$ slope
                      : num
                            16.85 6.51 4.91 5.55 14.55 ...
$ A.pseudoplatanus
                      : int
                            0 0 0 0 0 4 1 7 16 0 ...
$ F.sylvatica
                            61 0 0 100 0 82 85 91 75 0 ...
                      : int
$ L.decidua
                      : int
                            38 0 0 0 4 6 0 0 0 0 ...
                      : int 0100000000...
$ Q.robur
$ S.aucuparia
                      : int
                            0 0 0 0 0 0 0 0 0 0 ...
$ B.pendula
                            0 0 1 0 28 0 0 0 0 0 ...
                      : int
$ P.abies
                            0 99 93 0 30 0 0 0 8 99 ...
                      : int
$ P.sylvestris
                            0 0 7 0 1 0 0 0 0 0 ...
                      : int
$ F.excelsior
                      : int 00003800000...
$ A.alba
                      : int 0000040001...
                            0 0 0 0 0 2 2 2 0 0 ...
$ A.platanoides
                      : int
$ T.cordata
                            0 0 0 0 0 2 12 0 0 0 ...
                      : int
$ S.racemosa
                      : int
                            0 0 0 0 0 0 0 0 0 0 ...
$ U.glabra
                      : int
                            0 0 0 0 0 0 0 0 1 0 ...
$ S.nigra
                      : int 0000000000...
$ P.alba
                            0 0 0 0 0 0 0 0 0 0 ...
                      : int
                            0 0 0 0 0 0 0 0 0 0 ...
                      : int
$ U.minor
$ S.caprea
                            0 0 0 0 0 0 0 0 0 0 ...
                      : int
$ C.betulus
                            0 0 0 0 0 0 0 0 0 0 ...
                      : int
$ P.nigra
                      : int 0000000000...
```

```
$ Q.petraea
                             0 0 0 0 0 0 0 0 0 0 ...
                      : int
$ S.torminalis
                             0 0 0 0 0 0 0 0 0 0 ...
                      : int
$ A.campestre
                      : int
                             0 0 0 0 0 0 0 0 0 0 ...
$ P.strobus
                      : int
                             0 0 0 0 0 0 0 0 0 0 ...
$ Q.rubra
                      : int
                             0 0 0 0 0 0 0 0 0 0 ...
$ volAllha
                      : num 592 377 438 615 194 ...
$ GiniDBH
                      : num 0.448 0.416 0.12 0.133 0.378 ...
$ ShannonIndexTreeSpp : num    0.68    0.07    0.28    0    1.28    0.78    0.54    0.39    0.75    0.06    ...
$ Tracheophyta_rich : num 0.218 0.233 0.209 0.189 0.354 ...
$ Birds rich
                      : num 0.358 0.46 0.485 0.307 0.383 ...
$ Bryophytes_rich
                      : num 0.0876 0.0876 NA 0.2629 0.3505 ...
$ Fungi_rich
                             0.199 0.133 0.114 0.218 0.262 ...
                      : num
$ Lichens_rich
                      : num 0.0659 0.1976 0.1318 0.1537 0.1537 ...
$ Beetles_rich
                      : num
                             0.16 0.16 0.234 0.258 0.221 ...
$ dendrocoposMajor
                      : int 1 1 1 0 1 1 1 1 1 1 ...
$ certhia
                      : int 000001010...
$ bryophitaNumObs
                      : int 1 1 0 3 4 5 7 4 7 1 ...
$ birdNumObs
                      : int 14 18 19 12 15 14 12 14 18 18 ...
$ PlotID
                      : int 1 2 3 4 5 6 7 8 9 10 ...
```

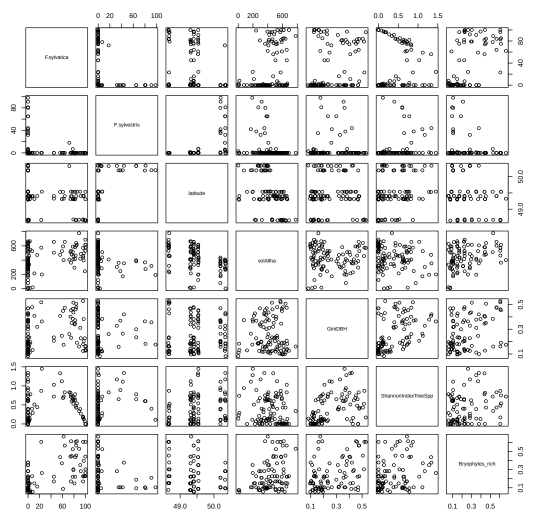
You have a description of each of the variables in the section Section 3.1.1.

You could do further analysis by exploring the data correlations and you can even plot them to explore their values and ranges better. During this process you should start thinking:

- What am I trying to understand with this model? (your question)
- What variables are important to answer my question?

You could for example explore the behavior of the variables by plotting a scatterplot:

Scatterplot Matrix

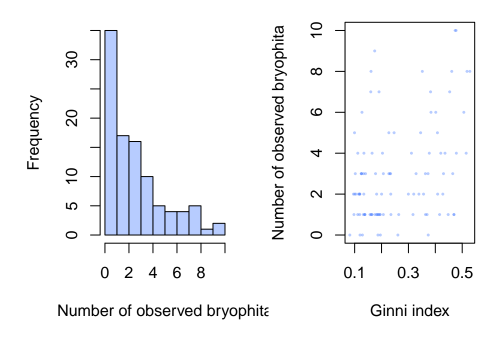


You can also create individual scatterplots or histograms for a variable of interest, for example:

```
#Divide the screen in 1 line and 3 columns
par(mfrow = c(1, 2), oma = c(0, 2, 0, 0))

#Make the margin around each graph a bit smaller
par(mar = c(4, 4, 2, 2))
# Histogram and Scatterplot
hist(
   observations$bryophitaNumObs,
```

```
main = "",
breaks = 10,
col = rgb(0.3, 0.5, 1, 0.4) ,
xlab = "Number of observed bryophita"
)
plot(
    y = observations$bryophitaNumObs,
    x = observations$GiniDBH,
    main = "" ,
    pch = 20,
    cex = 0.4,
    col = rgb(0.3, 0.5, 1, 0.4),
    xlab = "Ginni index",
    ylab = "Number of observed bryophita"
)
```



3.4.4 Question C - Group 1

• Does a more diverse forest in structure and composition have more Bryophytes species?

3.4.4.1 Fitting a Poisson GLM in R

bryoModel1 <- glm(bryophitaNumObs ~ GiniDBH,</pre>

During your data exploration, you should have selected your response variable, bryophitaNumObs. In this case, since we are trying to understand forest structure and composition, you should also choose explanatory variables for that, such as GiniDBH, which indicates the forest structural diversity in diameter sizes; higher values indicate more structural heterogeneity, and lower values indicate more homogeneous stands, or the ShannonIndexTreeSpp which assesses the diversity of tree species in the plot.

We could start by only looking at how the forest structural diversity affects the number of Bryophyte species that we have in a plot. You can create a GLM model with bryophitaNumObs as the response variable and GiniDBH as an explanatory variable. You will use the function glm in R and the family = poisson. You can see how to create and see this model here:

```
family = poisson,
                 data = observations)
summary(bryoModel1)
Call:
glm(formula = bryophitaNumObs ~ GiniDBH, family = poisson, data = observations)
Coefficients:
           Estimate Std. Error z value Pr(>|z|)
             0.4627
                        0.1388
                                 3.333 0.000859 ***
(Intercept)
GiniDBH
             2.2797
                        0.4221
                                 5.401 6.64e-08 ***
Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for poisson family taken to be 1)
   Null deviance: 188.58 on 98 degrees of freedom
Residual deviance: 159.92 on 97 degrees of freedom
AIC: 424.2
Number of Fisher Scoring iterations: 5
```

Here you can see that the coefficient table produced by a GLM is very similar to a linear model. The intercept tells us the estimated value of the response variable when the continuous explanatory variables (here, just the Gini index) has a value of 0. We

then also have coefficients describing the slope of the relationship with our continuous explanatory variables. We can see here that the number of Bryophytes species appears to show a positive relationship with the Gini index, which means that increasing structural diversity in tree diameter sizes has a positive relationship with the number of Bryophytes species in the plot.

We are also interested in understanding the relationship of the number of Bryophytes species and the tree species diversity; we can now try to add this variable into the model and see if it helps us to understand things. You can do that by adding the variable ShannonIndexTreeSpp to the model:

bryoModel2 <- glm(bryophitaNumObs ~ GiniDBH + ShannonIndexTreeSpp,</pre>

```
family = poisson,
                  data = observations)
summary(bryoModel2)
Call:
glm(formula = bryophitaNumObs ~ GiniDBH + ShannonIndexTreeSpp,
   family = poisson, data = observations)
Coefficients:
                   Estimate Std. Error z value Pr(>|z|)
(Intercept)
                     0.45277
                               0.14060
                                          3.220 0.00128 **
GiniDBH
                                0.46413
                                          4.684 2.81e-06 ***
                     2.17411
ShannonIndexTreeSpp 0.09209
                                0.16202
                                          0.568 0.56977
               0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Signif. codes:
(Dispersion parameter for poisson family taken to be 1)
   Null deviance: 188.58
                          on 98 degrees of freedom
Residual deviance: 159.60
                          on 96 degrees of freedom
AIC: 425.88
Number of Fisher Scoring iterations: 5
```

Here we can see that the variable Shannon index also has a positive relationship with the number of Bryophytes species in the plot, but its effect is much smaller. We can also observe that this variable is not significant. This does not mean it is wrong to add this variable because we want to understand its effect, but keeping this variable in the model depends on what you're trying to do and what "reality" is. Adding variables that

are not significant will not help your model (particularly your estimates) but also might not matter much (e.g., predictions). However, removing actual variables can create a useless model even if they don't meet significance.

Variable selection is a long and complicated topic. Some general rules of thumb include: (1) Include the variable if it is of interest, (2) Include the variable if you have some prior knowledge that it should be relevant. This can be misleading because it's a confirmation bias, but in most cases, this makes sense. (3) If you want a model that can generalize many cases, you should favor fewer variables.

3.4.4.2 Explanatory Power of the model

When we ran linear models, we used the coefficient of determination, or R^2 to assess how much of the variability in our response variable is explained by a given model. R^2 is based on the sums of squares of our model, and so cannot be calculated for GLMs. Instead, we can calculate the deviance explained by our model:

```
# Extract the null and residual deviance from the model
dev.null <- bryoModel1$null.deviance
dev.resid <- bryoModel1$deviance

# Calculate the deviance explained by the model
dev.explained <- (dev.null - dev.resid) / dev.null

# Round to 3 decimal places
dev.explained <- round(dev.explained, 3)

dev.explained</pre>
```

[1] 0.152

Variability in forest structure (Gini index) explain 15% of the variation in Bryophytes species richness in this study. That is an ok explanatory power for a very simple model of a complex ecological system (many factors determine the species richness for Bryophytes and we are attempting to explain everything with just one variable).

3.4.4.3 Model Assumptions

For Poisson GLMs, there is one further assumption that we have not encountered before. If data follow a Poisson distribution, then the mean of the distribution is equal to the variance. Accordingly, a Poisson distribution is represented by just one parameter λ , which describes both the mean and the variance of the distribution.

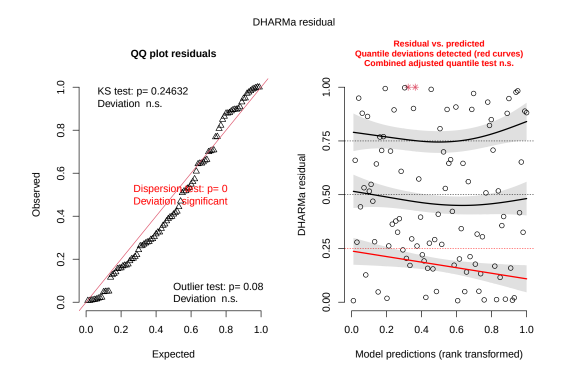
Count data in ecology are often **overdispersed**, where the variance is greater than the mean. This violates the assumption of a Poisson GLM, and means that any statistics that we calculate from the model may be unreliable.

We can get look whether a model is over-dispersed by inspecting the model summary as you did in Section 3.4.4.1. As a rule of thumb, if the response variable conforms to a true Poisson distribution, we expect the residual deviance to be approximately equal to the residual degrees of freedom. If the deviance is much greater than the degrees of freedom, this indicates over-dispersion. This is the case in our models (see outputs from running the code in Section 3.4.4.1).

To check the model assumptions in a GLM is not as straight forward as with a linear model. This is because classical residuals are not expected to behave in the same way for GLMs. We can use the DHARMa package in R for working with GLMs, which uses a simulation-based approach to compare the residuals from the actual model with the expectation if the model is behaving normally:

```
# Simulate residuals
simResids <- DHARMa::simulateResiduals(bryoModel1)

# Generate plots to compare the model residuals to expectations
plot(simResids)</pre>
```



These plots show us that this model is not behaving as we would expect in terms of homogeneity of variance and distribution of residuals. A follow up to this would be to try alternatives to deal with over-dispersed count data in GLMs such as fit a quasi-Poisson GLM or a negative binomial GLM. Unfortunately we do not have time to continuou in this exercise.

3.4.5 Question D - Group 1

• Is the number of Briophites species affected by forest management type and the forest structural diversity?

3.4.5.1 Fitting a Poisson GLM in R

Fitting a Poisson GLM in R is very similar to fitting an analysis of covariance (or linear model), except that now we need to use the glm function. To run a GLM, we need to provide one extra piece of information beyond that needed for a linear model: the family of model we want to use. In this case, we want a Poisson family.

We could start by only looking at how the forest structural diversity affects the number Bryophytes species that we have in a plot. You can do this by creating a GLM model which has bryophitaNumObs as response variable and GiniDBH as explanatory variable. For that you will use the function glm in R and use the family = poisson. You can see how to create and see this model here:

```
Null deviance: 188.58 on 98 degrees of freedom Residual deviance: 159.92 on 97 degrees of freedom
```

AIC: 424.2

Number of Fisher Scoring iterations: 5

Here you can see that the coefficient table produced by a GLM is very similar to a linear model. The intercept tells us the estimated value of the response variable when the continuous explanatory variables (here just Gini index) have a value of 0. We then also have coefficients describing the slope of the relationship with our continuous explanatory variables. We can see here that the bryophita numbers appears to show a positive relationship with Gini index, which means that increasing structural diversity in trees diameter sizes has a positive relationship with the number of bryophita species in the plot.

We are also interested in understanding the relationship of the number of observed Bryophytes species and forest management, we can now try to add this variable into the model.

Call:

```
glm(formula = bryophitaNumObs ~ forestManagementType, family = poisson,
    data = observations)
```

Coefficients:

```
(Intercept) Estimate Std. Error z value Pr(>|z|) (Intercept) 0.9651 0.1543 6.254 3.99e-10 forestManagementTypesimple clearcutting -0.1335 0.1750 -0.763 0.445 forestManagementTypeunmanaged 0.7633 0.1821 4.192 2.76e-05
```

(Dispersion parameter for poisson family taken to be 1)

```
Null deviance: 188.58 on 98 degrees of freedom
Residual deviance: 141.54 on 96 degrees of freedom
```

AIC: 407.82

Number of Fisher Scoring iterations: 5

The intercept here tells us the estimated value of the response variable when the reference groups in our grouping (categorical) variables (here for retention clear-cutting). We then also have coefficients describing the slope of the relationship with our continuous explanatory variables, and coefficients giving the estimated difference in the response variable for non-reference groupings. We can see here that number of bryophites species appears to show a negative relationship with simple clearcutting, and appears to have a positive relationship with unmanaged and retention clear-cutting.

The type simple clearcutting appears to be non significant, which only tells us about the pairwise differences between the levels. To test whether the categorical predictor, as a whole, is significant is equivalent to testing whether there is any heterogeneity in the means of the levels of the predictor. When there are no other predictors in the model, this is a classical ANOVA problem.

3.4.5.2 Explanatory Power of the model

When we run linear models, we use the coefficient of determination, or R^2 to assess how much of the variability in our response variable is explained by a given model. R^2 is based on the sums of squares of our model, and so it cannot be calculated for GLMs. Instead, we can calculate the the deviance explained by our model:

```
# Extract the null and residual deviance from the model
dev.null <- bryoModel1$null.deviance
dev.resid <- bryoModel1$deviance

# Calculate the deviance explained by the model
dev.explained <- (dev.null - dev.resid) / dev.null

# Round to 3 decimal places
dev.explained <- round(dev.explained, 3)

dev.explained</pre>
```

[1] 0.152

Variability in forest structure (Gini index) explains 15% of the variation in bryophita species richness in this study system. That is an ok explanatory power for a very simple model of a complex ecological system (many factors determine the species richness of bryophitas and we are attempting to explain everything with one variable).

3.4.5.3 Model Assumptions

For Poisson GLMs, there is one further assumption that we have not encountered before. If data follow a Poisson distribution, then the mean of the distribution is equal to the variance. Accordingly, a Poisson distribution is represented by just one parameter λ , which describes both the mean and the variance of the distribution.

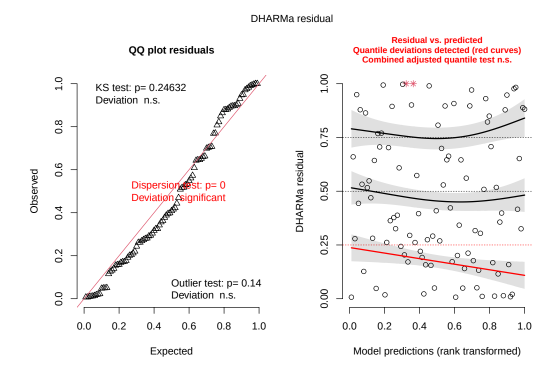
Count data in ecology are often *overdispersed*, where the variance is greater than the mean. This violates the assumption of a Poisson GLM, and means that any statistics that we calculate from the model may be unreliable.

We can get look whether a model is over-dispersed by inspecting the model summary as you did in Section 3.4.5.1. As a rule of thumb, if the response variable conforms to a true Poisson distribution, we expect the residual deviance to be approximately equal to the residual degrees of freedom. If the deviance is much greater than the degrees of freedom, this indicates over-dispersion. This is the case in our models (see outputs from running the code in Section 3.4.5.1).

To check the model assumptions in a GLM is not as straight-forward as with a linear model. This is because classical residuals are not expected to behave in the same way for GLMs. We can use the DHARMa package in R for working with GLMs, which uses a simulation-based approach to compare the residuals from the actual model with the expectation if the model is behaving normally.

```
# Simulate residuals
simResids <- DHARMa::simulateResiduals(bryoModel1)

# Generate plots to compare the model residuals to expectations
plot(simResids)</pre>
```



These plots show us that this model is not behaving as we would expect in terms of homogeneity of variance and distribution of residuals. A follow up to this would be to try alternatives to deal with over-dispersed count data in GLMs such as fit a quasi-Poisson GLM or a negative binomial GLM. Unfortunately we do not have to continue in this exercise.

3.4.6 Question C - Group 2

• Does a more diverse forest in structure and composition have more bird species?

3.4.6.1 Fitting a Poisson GLM in R

During your data exploration you should have selected your response variable: birdNumObs. In this case since we are trying to understand forest structure and composition you should also select explanatory variables for that such as GiniDBH which indicates the forest structural diversity in diameter sizes, higher values indicate more structural heterogeneity and lower values indicate more homogeneous stands, or the ShannonIndexTreeSpp which assess the diversity of tree species in the plot.

We could start by only looking at how the forest structural diversity affects the number bird species that we have in a plot. You can do this by creating a GLM model which has birdNumObs as response variable and GiniDBH as explanatory variable. For that you will use the function glmin R and use the family = poisson. You can see how to create and see this model here:

```
birdModel1 <- glm(birdNumObs ~ GiniDBH,
                  family = poisson,
                  data = observations)
summary(birdModel1)
Call:
glm(formula = birdNumObs ~ GiniDBH, family = poisson, data = observations)
Coefficients:
            Estimate Std. Error z value Pr(>|z|)
             2.63827
                       0.05636 46.809
                                          <2e-16 ***
(Intercept)
GiniDBH
             0.42725
                        0.19030
                                  2.245
                                          0.0248 *
                0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Signif. codes:
(Dispersion parameter for poisson family taken to be 1)
   Null deviance: 59.277
                           on 98 degrees of freedom
Residual deviance: 54.280 on 97 degrees of freedom
AIC: 511.57
Number of Fisher Scoring iterations: 4
```

Here you can see that the coefficient table produced by a GLM is very similar to a linear model. The intercept tells us the estimated value of the response variable when the continuous explanatory variables (here just Gini index) has a value of 0. We then also have coefficients describing the slope of the relationship with our continuous explanatory variables. We can see here that the number of bird species appears to show a positive relationship with Gini index, which means that increasing structural diversity in trees diameter sizes has a positive relationship with the number of bird species in the plot.

We are also interested in understanding the relationship of the number of bird species and the trees species diversity, we can now try to add this variable into the model and see if it help us to understand things. You can do that by adding the variable ShannonIndexTreeSpp into the model

```
Call:
glm(formula = birdNumObs ~ GiniDBH + ShannonIndexTreeSpp, family = poisson,
   data = observations)
Coefficients:
                    Estimate Std. Error z value Pr(>|z|)
(Intercept)
                     2.63499
                                0.05665 46.513
                                                  <2e-16 ***
GiniDBH
                     0.33323
                                0.21661
                                          1.538
                                                   0.124
                                0.07444
                                          0.930
                                                   0.352
ShannonIndexTreeSpp 0.06923
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for poisson family taken to be 1)
   Null deviance: 59.277
                          on 98 degrees of freedom
Residual deviance: 53.419
                          on 96 degrees of freedom
AIC: 512.71
```

Number of Fisher Scoring iterations: 4

Here we can see that the variable Shannon index also has a positive relationship with the number of bird species in the plot but its effect is much smaller. We can also observe that this variable is not significant and that including this variable also made Gini index variable not significant. This does not mean that is wrong to add this variable, because we want to understand its effect, but keeping this variable in the model or not depends on what you're trying to do, and what "reality" is. Adding variables that are not needed will not help your model (particularly your estimates), but also might not matter much (e.g. predictions). However, removing variables that are real, even if they don't meet significance, can create a useless model.

Variable selection is a long and complicated topic. some general rules of thumb include: (1) Include the variable if it is of interest, (2) Include the variable if you have some prior knowledge that it should be relevant. This can be misleading, because it's a confirmation bias, but in most cases this makes sense. (3) If you want a model that can generalize to many cases, you should favor fewer variables.

3.4.6.2 Explanatory Power of the model

When we ran linear models, we used the coefficient of determination, or R^2 to assess how much of the variability in our response variable is explained by a given model. R^2 is based on the sums of squares of our model, and so cannot be calculated for GLMs. Instead, we can calculate the deviance explained by our model:

```
# Extract the null and residual deviance from the model
dev.null <- birdModel1$null.deviance
dev.resid <- birdModel1$deviance

# Calculate the deviance explained by the model
dev.explained <- (dev.null - dev.resid) / dev.null

# Round to 3 decimal places
dev.explained <- round(dev.explained, 3)

dev.explained</pre>
```

[1] 0.084

Variability in forest structure (Gini index) explain 8% of the variation in bird species richness in this study. That is an ok explanatory power for a very simple model of a complex ecological system (many factors determine the species richness for birds and we are attempting to explain everything with just one variable).

3.4.6.3 Model Assumptions

For Poisson GLMs, there is one further assumption that we have not encountered before. If data follow a Poisson distribution, then the mean of the distribution is equal to the variance. Accordingly, a Poisson distribution is represented by just one parameter λ , which describes both the mean and the variance of the distribution.

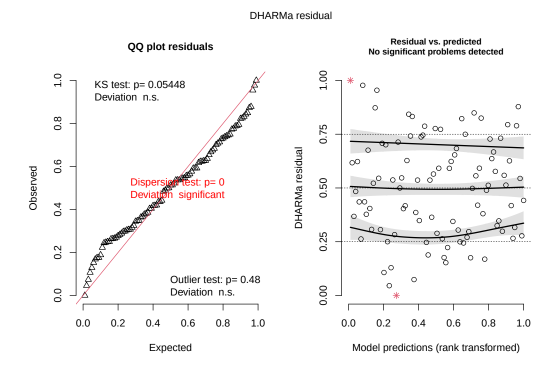
Count data in ecology are often **overdispersed**, where the variance is greater than the mean. This violates the assumption of a Poisson GLM, and means that any statistics that we calculate from the model may be unreliable.

We can get look whether a model is over-dispersed by inspecting the model summary as you did in Section 3.4.6.1. As a rule of thumb, if the response variable conforms to a true Poisson distribution, we expect the residual deviance to be approximately equal to the residual degrees of freedom. If the deviance is much greater than the degrees of freedom, this indicates over-dispersion. This is the case in our models (see outputs from running the code in Section 3.4.6.1).

To check the model assumptions in a GLM is not as straight-forward as with a linear model. This is because classical residuals are not expected to behave in the same way for GLMs. We can use the DHARMa package in R for working with GLMs, which uses a simulation-based approach to compare the residuals from the actual model with the expectation if the model is behaving normally:

```
# Simulate residuals
simResids <- DHARMa::simulateResiduals(birdModel1)

# Generate plots to compare the model residuals to expectations
plot(simResids)</pre>
```



These plots show us that this model is not behaving as we would expect in terms of homogeneity of variance and distribution of residuals. A follow up to this would be to try alternatives to deal with over-dispersed count data in GLMs such as fit a quasi-Poisson GLM or a negative binomial GLM. Unfortunately we do not have time to continuou in this exercise.

3.4.7 Question D - Group 2

• Is the number of bird species affected by forest management type and the forest structural diversity?

3.4.7.1 Fitting a Poisson GLM in R

Fitting a Poisson GLM in R is very similar to fitting an analysis of covariance (or linear model), except that now we need to use the glm function. To run a GLM, we need to provide one extra piece of information beyond that needed for a linear model: the family of model we want to use. In this case, we want a Poisson family.

We could start by only looking at how the forest structural diversity affects the number bird species that we have in a plot. You can do this by creating a GLM model which has birdNumObs as response variable and GiniDBH as explanatory variable. For that you will use the function glm in R and use the family = poisson. You can see how to create and see this model here:

```
birdModel1 <- glm(birdNumObs ~ GiniDBH,</pre>
                  family = poisson,
                  data = observations)
summary(birdModel1)
Call:
glm(formula = birdNumObs ~ GiniDBH, family = poisson, data = observations)
Coefficients:
            Estimate Std. Error z value Pr(>|z|)
(Intercept) 2.63827
                        0.05636 46.809
                                          <2e-16 ***
GiniDBH
             0.42725
                        0.19030
                                  2.245
                                          0.0248 *
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for poisson family taken to be 1)
    Null deviance: 59.277 on 98 degrees of freedom
Residual deviance: 54.280
                          on 97
                                  degrees of freedom
AIC: 511.57
Number of Fisher Scoring iterations: 4
```

Here you can see that the coefficient table produced by a GLM is very similar to a linear model. The intercept tells us the estimated value of the response variable when the continuous explanatory variables (here just Gini index) have a value of 0. We then also have coefficients describing the slope of the relationship with our continuous explanatory variables. We can see here that the bird numbers appears to show a positive relationship with Gini index, which means that increasing structural diversity in trees diameter sizes has a positive relationship with the number of bird species in the plot.

We are also interested in understanding the relationship of the number of observed bird species and forest management, we can now try to add this variable into the model.

birdModel2 <- glm(birdNumObs ~ forestManagementType,</pre>

```
family = poisson,
                  data = observations)
summary(birdModel2)
Call:
glm(formula = birdNumObs ~ forestManagementType, family = poisson,
    data = observations)
Coefficients:
                                        Estimate Std. Error z value Pr(>|z|)
(Intercept)
                                         2.83688
                                                    0.06052 46.873
                                                                      <2e-16
forestManagementTypesimple clearcutting -0.11744
                                                    0.06850 -1.714
                                                                      0.0865
forestManagementTypeunmanaged
                                        -0.06429
                                                    0.08338 -0.771
                                                                      0.4407
(Intercept)
forestManagementTypesimple clearcutting .
forestManagementTypeunmanaged
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for poisson family taken to be 1)
    Null deviance: 59.277
                          on 98 degrees of freedom
Residual deviance: 56.205 on 96 degrees of freedom
AIC: 515.49
Number of Fisher Scoring iterations: 4
```

The intercept here tells us the estimated value of the response variable when the reference groups in our grouping (categorical) variables (here for retention clear-cutting). We

then also have coefficients describing the slope of the relationship with our continuous explanatory variables, and coefficients giving the estimated difference in the response variable for non-reference groupings. We can see here that number of bird species appears to show a negative relationship with simple clearcutting, unmanaged and a positive relationship with retention clear-cutting.

The type simple clearcutting and unmanaged appear to be no significant, which only tell us about the pairwise differences between the levels. To test whether the categorical predictor, as a whole, is significant is equivalent to testing whether there is any heterogeneity in the means of the levels of the predictor. When there are no other predictors in the model, this is a classical ANOVA problem.

3.4.7.2 Explanatory Power of the model

When we ran linear models, we used the coefficient of determination, or \mathbb{R}^2 to assess how much of the variability in our response variable is explained by a given model. \mathbb{R}^2 is based on the sums of squares of our model, and so cannot be calculated for GLMs. Instead, we can calculate the the deviance explained by our model:

```
# Extract the null and residual deviance from the model
dev.null <- birdModel1$null.deviance
dev.resid <- birdModel1$deviance

# Calculate the deviance explained by the model
dev.explained <- (dev.null - dev.resid) / dev.null

# Round to 3 decimal places
dev.explained <- round(dev.explained, 3)

dev.explained</pre>
```

[1] 0.084

Variability in forest structure (Gini index) explain 15% of the variation in bird species richness in this study system. That is an ok explanatory power for a very simple model of a complex ecological system (many factors determine the species richness and we are attempting to explain everything with one variable).

3.4.7.3 Model Assumptions

For Poisson GLMs, there is one further assumption that we have not encountered before. If data follow a Poisson distribution, then the mean of the distribution is equal to the

variance. Accordingly, a Poisson distribution is represented by just one parameter λ , which describes both the mean and the variance of the distribution.

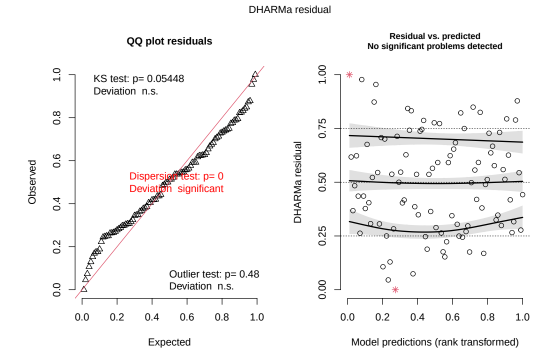
Count data in ecology are often *overdispersed*, where the variance is greater than the mean. This violates the assumption of a Poisson GLM, and means that any statistics that we calculate from the model may be unreliable.

We can get look whether a model is over-dispersed by inspecting the model summary as you did in Section 3.4.7.1. As a rule of thumb, if the response variable conforms to a true Poisson distribution, we expect the residual deviance to be approximately equal to the residual degrees of freedom. If the deviance is much greater than the degrees of freedom, this indicates over-dispersion. This is the case in our models (see outputs from running the code in Section 3.4.7.1).

To check the model assumptions in a GLM is not as straight forward as with a linear model. This is because classical residuals are not expected to behave in the same way for GLMs. We can use the DHARMa package in R for working with GLMs, which uses a simulation-based approach to compare the residuals from the actual model with the expectation if the model is behaving normally.

```
# Simulate residuals
simResids <- DHARMa::simulateResiduals(birdModel1)

# Generate plots to compare the model residuals to expectations
plot(simResids)</pre>
```



These plots show us that this model is not behaving as we would expect in terms of homogeneity of variance and distribution of residuals. A follow up to this would be to try alternatives to deal with over-dispersed count data in GLMs such as fit a quasi-Poisson GLM or a negative binomial GLM. Unfortunately we do not have to continuou in this exercise.

3.4.8 Question C - Group 3

• Is the presence of the Great spotted woodpecker affected by forest density?

3.4.8.1 Fitting a BRT in R

In this case we want to assess the occurrence of certain species across the plots. In other words, we want to assess what is the probability of a certain species with biodiversity interest to be present in a plot based on the variables that describe the forest of that plot. In this case the response variable is dendrocoposMajor that represents if the Great spotted woodpecker has been observed in this plot or not.

Then you need to select some variables of interest, after you have explored the data you can decide which variables you want to use to fit this model. We are proposing to select the following variables:

- latitude as proxy for plot location or/and climate
- forestManagementType to assess if different management types have different impact in the presence / absence of Great spotted woodpecker.
- volAllha that is the total volume in the plot, as a proxy of how dense the plot is. Higher volumes will mean that the forest is more dense.
- GiniDBH showing how homogeneous the plot is in trees diameters. A value closer to 1 indicate a more structural heterogeneity, lower values indicate more homogeneous plots.

You can create a vector **selVar** in which you add the names of the selected variables. Then you only take those variables from the data that you will use to create the model.

```
# Select variables from the dataset for the model
selVar <- c(
    "dendrocoposMajor",
    "latitude",
    "forestManagementType",
    "volAllha",
    "GiniDBH",
    "ShannonIndexTreeSpp"
)

# Filter the dataset to the selected variables
modelDataSel <- observations[, colnames(observations) %in% selVar]</pre>
```

Unfortunately the amount of that we have in this dataset it is not enough to fit a BRT model for these variables. We are going to do an obviously wrong thing for the shake of being able to demonstrate how to fit a BRT model. In the next code you are going to repeat the same dataset multiple times:

Now it is important to assess if the variables have the right categories. Variables should be type numeric or factor.

```
summary(modelDataSel)
```

```
latitude forestManagementType volAllha GiniDBH
Min.:48.65 Length:495 Min.: 1.681 Min.:0.08209
1st Qu::49.31 Class:character 1st Qu::319.920 1st Qu::0.13684
```

```
Median :49.40
                Mode :character
                                       Median :434.729
                                                          Median: 0.20358
       :49.49
                                              :425.694
                                                                 :0.25683
Mean
                                       Mean
                                                          Mean
                                                          3rd Qu.:0.37368
3rd Qu.:50.19
                                       3rd Qu.:558.355
Max.
       :50.34
                                       Max.
                                              :777.882
                                                          Max.
                                                                 :0.52852
ShannonIndexTreeSpp dendrocoposMajor
Min.
       :0.000
                     Min.
                            :0.0000
1st Qu.:0.060
                     1st Qu.:0.0000
Median :0.280
                     Median :1.0000
Mean
       :0.392
                     Mean
                            :0.7071
3rd Qu.:0.680
                     3rd Qu.:1.0000
Max.
       :1.450
                     Max.
                            :1.0000
```

```
# Two variables are character, we assign to factor instead:
modelDataSel$forestManagementType <-
    as.factor(modelDataSel$forestManagementType)</pre>
```

In the next step you can see how you can run the model with the selected variables and model parameters. You have a description of the models parameters in the Section 3.3.2 . In this example we are going to use the default parameters for the calibration, where learning rate = 0.01 and tree complexity = 1 and cross-validation = 10-fold. However, the bag fraction is changed from the default value, 0.75, to 0.5. As a family we used the Bernoulli family, because we are predicting presence/absence per plot. These data have 495 plots, comprising 350 presence records for the Great spotted woodpecker. You can check these numbers by doing:

```
table(modelDataSel$dendrocoposMajor)
```

```
0 1
145 350
```

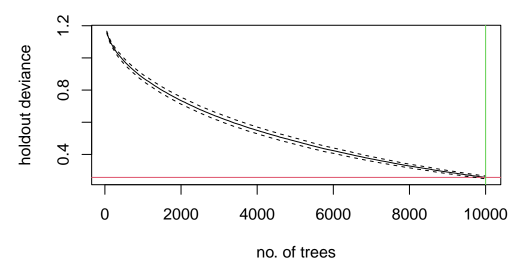
As a first guess you could decide there are enough data to model interactions of reasonable complexity, and a lr of about 0.01 could be a reasonable starting point. You can use the model creation function that steps forward and identifies the optimal number of trees (nt) by doing this:

```
family <- "bernoulli"
tc = 1  # tree complexity
lr = 0.01 # learning rate-shrinkage
bag = 0.5 # bag fraction

modelBRT <- dismo::gbm.step(</pre>
```

```
data = modelDataSel,
    #indices of predictor variables in data
    gbm.x = 1:5,
    #index of response variable in data:
    gbm.y = 6,
    family = family,
    tree.complexity = tc,
    learning.rate = lr,
    bag.fraction = bag
)
```

dendrocoposMajor, d - 1, lr - 0.01



Running a model such as that described above writes progress reports to the screen, makes a graph, and returns an object containing a number of components. The R console results reports a brief model summary all the values are also retained in the model object.

The model is built with the default 10-fold cross-validation (CV). In the plotted graph the solid black curve is the mean, and the dotted curves 1 standard error, for the changes in predictive deviance (i.e., as measured on the excluded folds of the CV). The red line shows the minimum of the mean, and the green line the number of trees at which that occurs. The final model that is returned in the model object is built on the full data set, using the number of trees identified as optimal.

Ideally, you should invest time in modifying the parameters and find the parameters that provide the models with the minimum deviance resulting from the best combination of bag, tree complexity and learning rate values. For the shake of limited timing, we will only test here the default values.

3.4.8.2 Model behaviour

You can summarize the model parameters used and the cross validation statistics from the fitted model by doing this:

```
# We make a table with the summary statistics
results <- data.frame(
    # Model parameters
    Tree.Complexity = modelBRT$gbm.call$tree.complexity,
    Learning.Rate = modelBRT$gbm.call$learning.rate,
    Bag.Fraction = modelBRT$gbm.call$bag.fraction,
    Interaction.depth = modelBRT$interaction.depth,
    Shrinkage = modelBRT$shrinkage,
    N.trees = modelBRT$n.trees,
    # Cross validation statistics
    ## mean total deviance
    Deviance = modelBRT$self.statistics$mean.resid,
    # mean residual deviance
    AUC = modelBRT$self.statistics$discrimination,
    # training data AUC score
    Corr = modelBRT$self.statistics$correlation,
    # training data correlation
    ## Cross Validation statistics
    # We calculate each statistic within each fold (at the identified optimal number
    # of trees that is calculated on the mean change in predictive deviance over all fold
    #then present here the mean and standard error of those fold-based statistics.
    devianceCV = modelBRT$cv.statistics$deviance.mean,
    # estimated cv deviance
    devianceCVse = modelBRT$cv.statistics$deviance.se,
    # estimated cv deviance se
```

```
CorrCV = modelBRT$cv.statistics$correlation.mean,
    #cv correlation
    CorrCVse = modelBRT$cv.statistics$correlation.se,
    #cv correlation se

AUCcv = modelBRT$cv.statistics$discrimination.mean,
    # cv AUC score
    AUCcvSE = modelBRT$cv.statistics$discrimination.se
) # cv AUC score se
print(t(results))
```

[,1]1.000000e+00 Tree.Complexity Learning.Rate 1.000000e-02 Bag.Fraction 5.000000e-01 Interaction.depth 1.000000e+00 Shrinkage 1.000000e-02 N.trees 1.000000e+04 Deviance 2.180946e-01 AUC 1.000000e+00 Corr 9.746990e-01 2.571289e-01 devianceCV devianceCVse 8.660155e-03 CorrCV 9.600872e-01 CorrCVse 4.284915e-03 AUCcv 1.000000e+00 AUCcvSE 0.000000e+00

3.4.8.3 Model output analysis

We can look at the relative contribution of each of the predictor variables. The measures are based on the number of times the variable is selected for splitting, weighted by the improvement of the model as a result of each split averaged across all trees. The relative contribution of each of the variables is scaled so the sum is 100%, with higher numbers indicating stronger influence in the response.

```
# Variables contribution
modelBRT$contributions
```

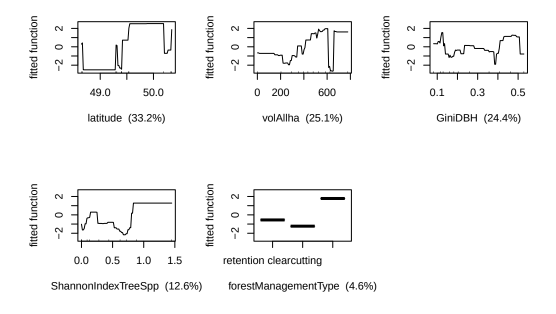
```
var rel.inf
latitude latitude 33.223565
volAllha volAllha 25.100356
GiniDBH GiniDBH 24.438618
ShannonIndexTreeSpp ShannonIndexTreeSpp 12.639937
forestManagementType forestManagementType 4.597523
```

Here we can see that the two variables with the highest influence in the response are latitude, and volAllhsa.

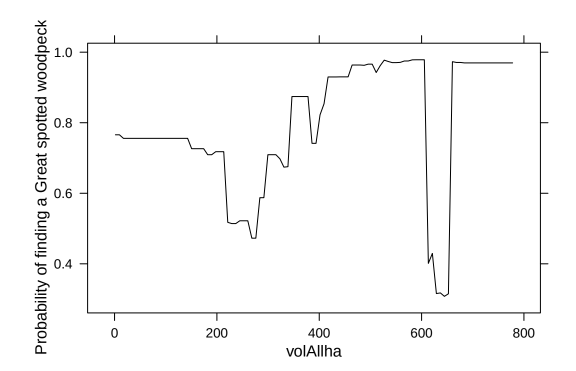
Now we can evaluate the model behavior via partial dependence plots, showing the effect of each of the variables on the response by accounting for the average effects of all other predictors in the model:

```
dismo::gbm.plot(
    modelBRT,
    n.plots = 6,
    plot.layout = c(2, 3),
    write.title = F
)
```

Warning in dismo::gbm.plot(modelBRT, n.plots = 6, plot.layout = c(2, 3), : reducing no of plotted predictors to maximum available (5)



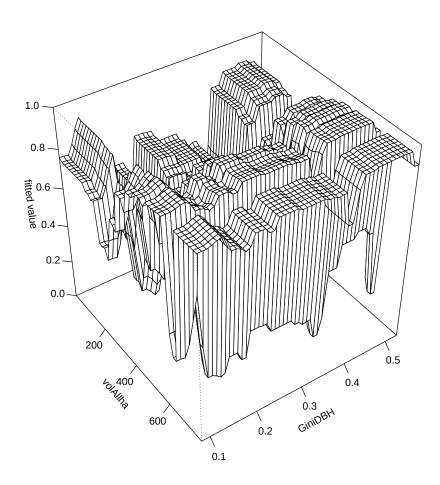
In partial dependence plots the predictions are on the scale of f(x). In this case, for the Bernoulli loss the returned value is on the log odds scale. You can see how this plot will look by plotting with the function from the package gbm and using the type "response". Since we are interested in finding out if the forest density has an impact on the presence of the Great spotted woodpecker we can have a look to the plot density variable volAllha:



It seems that there is an increase in probability of finding a Great spotted woodpecker with higher forest densities, but the trend it is not very clear. We could also analyse the interaction effects, of density and for example overall bird richness. The model predictions can be obtained for each pair of predictor variables, setting all other predictors to their means.

To plot this pairwise interactions we have to do:

```
dismo::gbm.perspec(modelBRT, 3, 4)
```



Here we can see that both increasing bird diversity and forest density provides the highest probabilities for finding the Great spotted woodpecker.

3.4.9 Question D - Group 3

• Is the presence of the Great spotted woodpecker affected by forest diversity?

3.4.9.1 Fitting a BRT in R

In this case we want to assess the occurrence of certain species across the plots. In other words, we want to assess what is the probability of a certain species with biodiversity interest to be present in a plot based on the variables that describe the forest of that plot. In this case the response variable is dendrocoposMajor that represents if the Great spotted woodpecker has been observed in this plot or not.

Then you need to select some variables of interest, after you have explored the data you can decide which variables you want to use to fit this model. We are proposing to select the following variables:

- latitude as proxy for plot location or/and climate
- forestManagementType to assess if different management types have different impact in the presence / absence of Great spotted woodpecker.
- volAllha that is the total volume in the plot, as a proxy of how dense the plot is. Higher volumes will mean that the forest is more dense.
- GiniDBH showing how homogeneous the plot is in trees diameters. A value closer to 1 will mean that indicate more structural heterogeneity, lower values indicate more homogeneous plots.

You can create a vector **selVar** in which you add the names of the selected variables. Then you only take those variables from the data that you will use to create the model.

```
# Select variables from the dataset for the model
selVar <- c(
    "dendrocoposMajor",
    "latitude",
    "forestManagementType",
    "volAllha",
    "GiniDBH",
    "ShannonIndexTreeSpp"
)

# Filter the dataset to the selected variables
modelDataSel <- observations[, colnames(observations) %in% selVar]</pre>
```

Unfortunately the amount of that we have in this dataset it is not enough to fit a BRT model for these variables. We are going to do an obviously wrong thing for the shake of being able to demonstrate how to fit a BRT model. In the next code you are going to repeat the same dataset multiple times:

Now it is important to assess if the variables have the right categories. Variables should be type numeric or factor.

summary(modelDataSel)

```
latitude
                 forestManagementType
                                          volAllha
                                                              GiniDBH
       :48.65
                 Length: 495
                                               : 1.681
                                                                  :0.08209
Min.
                                       Min.
                                                          Min.
1st Qu.:49.31
                 Class :character
                                       1st Qu.:319.920
                                                           1st Qu.:0.13684
Median :49.40
                 Mode
                      :character
                                       Median: 434.729
                                                          Median: 0.20358
       :49.49
Mean
                                       Mean
                                               :425.694
                                                          Mean
                                                                  :0.25683
3rd Qu.:50.19
                                       3rd Qu.:558.355
                                                           3rd Qu.:0.37368
Max.
       :50.34
                                       Max.
                                               :777.882
                                                          Max.
                                                                  :0.52852
ShannonIndexTreeSpp dendrocoposMajor
Min.
       :0.000
                     Min.
                             :0.0000
1st Qu.:0.060
                     1st Qu.:0.0000
Median :0.280
                     Median :1.0000
Mean
       :0.392
                     Mean
                             :0.7071
3rd Qu.:0.680
                     3rd Qu.:1.0000
Max.
       :1.450
                     Max.
                             :1.0000
```

```
# Two variables are character, we assign to factor instead:
modelDataSel$forestManagementType <-
    as.factor(modelDataSel$forestManagementType)</pre>
```

In the next step you can see how you can run the model with the selected variables and model parameters. You have a description of the models parameters in the Section 3.3.2. In this example we are going to use the default parameters for the calibration, where learning rate = 0.01 and tree complexity = 1 and cross-validation = 10-fold. However, the bag fraction is changed from the default value, 0.75, to 0.5. As a family we used the Bernoulli family, because we are predicting presence/absence per plot. These data have 495 plots, comprising 350 presence records for the Great spotted woodpecker. You can check these numbers by doing:

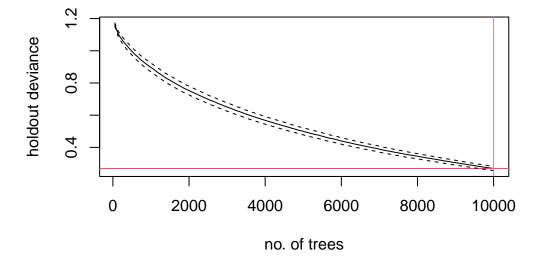
```
table(modelDataSel$dendrocoposMajor)
```

```
0 1
145 350
```

As a first guess you could decide there are enough data to model interactions of reasonable complexity, and a lr of about 0.01 could be a reasonable starting point. You can use the model creation function that steps forward and identifies the optimal number of trees (nt) by doing this:

```
family <- "bernoulli"
tc = 1
          # tree complexity
lr = 0.01 # learning rate-shrinkage
bag = 0.5 \# bag fraction
modelBRT <- dismo::gbm.step(</pre>
    data = modelDataSel,
    #indices of predictor variables in data
    gbm.x = 1:5,
    #index of response variable in data:
    gbm.y = 6,
    family = family,
    tree.complexity = tc,
    learning.rate = lr,
    bag.fraction = bag
)
```

dendrocoposMajor, d - 1, lr - 0.01



Running a model such as that described above writes progress reports to the screen, makes a graph, and returns an object containing a number of components. The R console results reports a brief model summary all the values are also retained in the model object.

The model is built with the default 10-fold cross-validation (CV). In the plotted graph the solid black curve is the mean, and the dotted curves 1 standard error, for the changes in predictive deviance (i.e., as measured on the excluded folds of the CV). The red line shows the minimum of the mean, and the green line the number of trees at which that occurs. The final model that is returned in the model object is built on the full data set, using the number of trees identified as optimal.

Ideally, you should invest time in modifying the parameters and find the parameters that provide the models with the minimum deviance resulting from the best combination of bag, tree complexity and learning rate values. For the shake of limited timing, we will only test here the default values.

3.4.9.2 Model behaviour

You can summarize the model parameters used and the cross validation statistics from the fitted model by doing this:

```
# We make a table with the summary statistics
results <- data.frame(
    # Model parameters
    Tree.Complexity = modelBRT$gbm.call$tree.complexity,
    Learning.Rate = modelBRT$gbm.call$learning.rate,
    Bag.Fraction = modelBRT$gbm.call$bag.fraction,
    Interaction.depth = modelBRT$interaction.depth,
    Shrinkage = modelBRT$shrinkage,
    N.trees = modelBRT$n.trees,
    # Cross validation statistics
    ## mean total deviance
    Deviance = modelBRT$self.statistics$mean.resid,
    # mean residual deviance
    AUC = modelBRT$self.statistics$discrimination,
    # training data AUC score
    Corr = modelBRT$self.statistics$correlation,
    # training data correlation
```

```
## Cross Validation statistics
    # We calculate each statistic within each fold (at the identified optimal number
    # of trees that is calculated on the mean change in predictive deviance over all fold
    #then present here the mean and standard error of those fold-based statistics.
    devianceCV = modelBRT$cv.statistics$deviance.mean,
    # estimated cv deviance
    devianceCVse = modelBRT$cv.statistics$deviance.se,
    # estimated cv deviance se
    CorrCV = modelBRT$cv.statistics$correlation.mean,
    #cv correlation
    CorrCVse = modelBRT$cv.statistics$correlation.se,
    #cv correlation se
    AUCcv = modelBRT$cv.statistics$discrimination.mean,
    # cv AUC score
   AUCcvSE = modelBRT$cv.statistics$discrimination.se
) # cv AUC score se
print(t(results))
```

```
[,1]
                  1.000000e+00
Tree.Complexity
Learning.Rate
                  1.000000e-02
Bag.Fraction
                  5.000000e-01
Interaction.depth 1.000000e+00
                  1.000000e-02
Shrinkage
N.trees
                  1.000000e+04
Deviance
                  2.183994e-01
AUC
                 1.000000e+00
                 9.750907e-01
Corr
devianceCV
                 2.691805e-01
devianceCVse
                 1.296261e-02
CorrCV
                  9.535493e-01
CorrCVse
                  6.295274e-03
AUCcv
                  1.000000e+00
AUCcvSE
                  0.000000e+00
```

3.4.9.3 Model output analysis

We can look at the relative contribution of each of the predictor variables. The measures are based on the number of times the variable is selected for splitting, weighted by the improvement of the model as a result of each split averaged across all trees. The relative contribution of each of the variables is scaled so the sum is 100%, with higher numbers indicating stronger influence in the response.

```
# Variables contribution
modelBRT$contributions
```

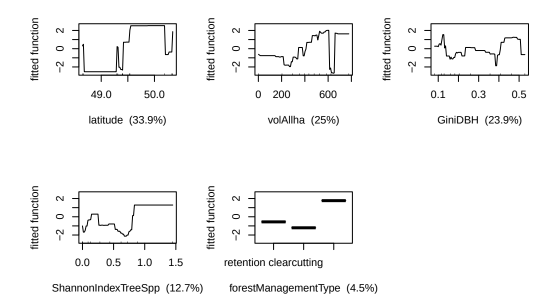
```
var rel.inf
latitude latitude 33.907767
volAllha volAllha 25.041328
GiniDBH GiniDBH 23.857644
ShannonIndexTreeSpp ShannonIndexTreeSpp 12.656084
forestManagementType forestManagementType 4.537178
```

Here we can see that the two variables with the highest influence in the response are latitude, and volAllhsa.

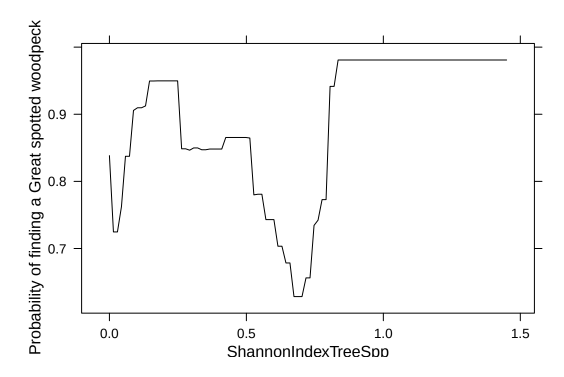
Now we can evaluate the model behavior via partial dependence plots, showing the effect of each of the variables on the response by accounting for the average effects of all other predictors in the model:

```
dismo::gbm.plot(
    modelBRT,
    n.plots = 6,
    plot.layout = c(2, 3),
    write.title = F
)
```

Warning in dismo::gbm.plot(modelBRT, n.plots = 6, plot.layout = c(2, 3), : reducing no of plotted predictors to maximum available (5)



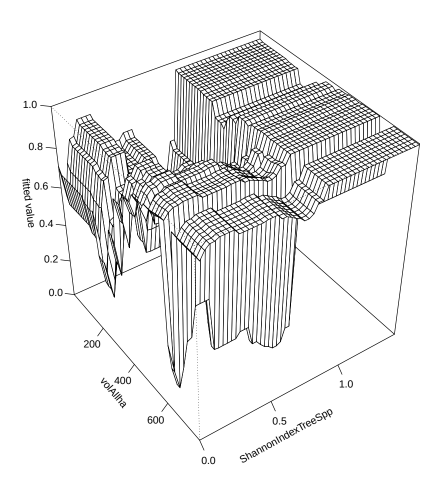
In this partial dependence plots the predictions are on the scale of f(x). In this case, for the Bernoulli loss the returned value is on the log odds scale. You can see how this plot will look by plotting with the function from the package gbm and using the type "response". Since we are interesting in finding out if the forest diversity has an impact in the presence of the Great spotted woodpecker we can have a look to the plot density variable volAllha:



It seems that there is an increase in probability of finding a Great spotted woodpecker with higher forest densities, but the trend it is not very clear. We could also analyse the interaction effects, of forest diversity and for example forest density. The model predictions can be obtained for each pair of predictor variables, setting all other predictors to their means.

To plot this pairwise interactions we have to do:

dismo::gbm.perspec(modelBRT, 3, 5)



Here we can not see a very clear combined behavior between forest diversity and forest structural diversity in respect to the probabilities for finding the Great spotted woodpecker.

3.4.10 Question C - Group 4

• Is the presence of the Eurasian treecreeper affected by forest density?

3.4.10.1 Fitting a BRT in R

In this case we want to assess the occurrence of certain species across the plots. In other words, we want to assess what is the probability of a certain species with biodiversity interest to be present in a plot based on the variables that describe the forest of that plot. In this case the response variable is **certhia** that represents if the Eurasian treecreeper has been observed in this plot or not.

Then you need to select some variables of interest, after you have explored the data you can decide which variables you want to use to fit this model. We are proposing to select the following variables:

- latitude as proxy for plot location or/and climate
- forestManagementType to assess if different management types have different impact in the presence / absence of Great spotted woodpecker.
- volAllha that is the total volume in the plot, as a proxy of how dense the plot is. Higher volumes will mean that the forest is more dense.
- GiniDBH showing how homogeneous the plot is in trees diameters. A value closer to 1 will mean that indicate more structural heterogeneity, lower values indicate more homogeneous plots.

You can create a vector **selVar** in which you add the names of the selected variables. Then you only take those variables from the data that you will use to create the model.

```
# Select variables from the dataset for the model
selVar <- c(
    "certhia",
    "latitude",
    "forestManagementType",
    "volAllha",
    "GiniDBH",
    "ShannonIndexTreeSpp"
)

# Filter the dataset to the selected variables
modelDataSel <- observations[, colnames(observations) %in% selVar]</pre>
```

Unfortunately the amount of that we have in this dataset it is not enough to fit a BRT model for these variables. We are going to do an obviously wrong thing for the shake of being able to demonstrate how to fit a BRT model. In the next code you are going to repeat the same dataset multiple times:

Now it is important to assess if the variables have the right categories. Variables should be type numeric or factor.

summary(modelDataSel)

```
GiniDBH
   latitude
                 forestManagementType
                                          volAllha
Min.
       :48.65
                 Length:495
                                       Min.
                                              : 1.681
                                                          Min.
                                                                  :0.08209
1st Qu.:49.31
                 Class : character
                                       1st Qu.:319.920
                                                          1st Qu.:0.13684
Median :49.40
                 Mode :character
                                       Median: 434.729
                                                          Median :0.20358
Mean
       :49.49
                                       Mean
                                               :425.694
                                                                  :0.25683
                                                          Mean
3rd Qu.:50.19
                                       3rd Qu.:558.355
                                                          3rd Qu.:0.37368
                                       Max.
Max.
       :50.34
                                              :777.882
                                                          Max.
                                                                  :0.52852
{\tt ShannonIndexTreeSpp}
                        certhia
Min.
       :0.000
                            :0.0000
                     Min.
1st Qu.:0.060
                     1st Qu.:0.0000
Median :0.280
                     Median :1.0000
Mean
       :0.392
                            :0.5859
                     Mean
3rd Qu.:0.680
                     3rd Qu.:1.0000
Max.
       :1.450
                            :1.0000
                     Max.
```

```
# Two variables are character, we assign to factor instead:
modelDataSel$forestManagementType <-
    as.factor(modelDataSel$forestManagementType)</pre>
```

In the next step you can see how you can run the model with the selected variables and model parameters. You have a description of the models parameters in the Section 3.3.2 . In this example we are going to use the default parameters for the calibration, where learning rate = 0.01 and tree complexity = 1 and cross-validation = 10-fold. However, the bag fraction is changed from the default value, 0.75, to 0.5. As a family we used the Bernoulli family, because we are predicting presence/absence per plot. These data have 495 plots, comprising 290 presence records for the Eurasian treecreeper. You can check these numbers by doing:

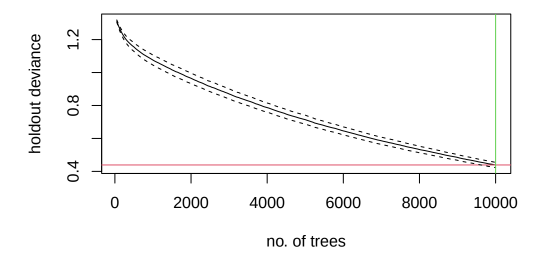
```
table(modelDataSel$certhia)
```

0 1 205 290

As a first guess you could decide there are enough data to model interactions of reasonable complexity, and a lr of about 0.01 could be a reasonable starting point. You can use the model creation function that steps forward and identifies the optimal number of trees (nt) by doing this:

```
family <- "bernoulli"
tc = 1
          # tree complexity
lr = 0.01 # learning rate-shrinkage
bag = 0.5 \# bag fraction
modelBRT <- dismo::gbm.step(</pre>
    data = modelDataSel,
    #indices of predictor variables in data
    gbm.x = 1:5,
    #index of response variable in data:
    gbm.y = 6,
    family = family,
    tree.complexity = tc,
    learning.rate = lr,
    bag.fraction = bag
)
```

certhia, d - 1, lr - 0.01



Running a model such as that described above writes progress reports to the screen, makes a graph, and returns an object containing a number of components. The R console results reports a brief model summary all the values are also retained in the model object.

The model is built with the default 10-fold cross-validation (CV). In the plotted graph the solid black curve is the mean, and the dotted curves 1 standard error, for the changes in predictive deviance (i.e., as measured on the excluded folds of the CV). The red line shows the minimum of the mean, and the green line the number of trees at which that occurs. The final model that is returned in the model object is built on the full data set, using the number of trees identified as optimal.

Ideally, you should invest time in modifying the parameters and find the parameters that provide the models with the minimum deviance resulting from the best combination of bag, tree complexity and learning rate values. For the shake of limited timing, we will only test here the default values.

3.4.10.2 Model behaviour

You can summarized the model parameters used and the cross validation statistics from the fitted model by doing this:

```
# We make a table with the summary statistics
results <- data.frame(
    # Model parameters
    Tree.Complexity = modelBRT$gbm.call$tree.complexity,
    Learning.Rate = modelBRT$gbm.call$learning.rate,
    Bag.Fraction = modelBRT$gbm.call$bag.fraction,
    Interaction.depth = modelBRT$interaction.depth,
    Shrinkage = modelBRT$shrinkage,
    N.trees = modelBRT$n.trees,
    # Cross validation statistics
    ## mean total deviance
    Deviance = modelBRT$self.statistics$mean.resid,
    # mean residual deviance
    AUC = modelBRT$self.statistics$discrimination,
    # training data AUC score
    Corr = modelBRT$self.statistics$correlation,
    # training data correlation
```

```
## Cross Validation statistics
    # We calculate each statistic within each fold (at the identified optimal number
    # of trees that is calculated on the mean change in predictive deviance over all fold
    #then present here the mean and standard error of those fold-based statistics.
    devianceCV = modelBRT$cv.statistics$deviance.mean,
    # estimated cv deviance
    devianceCVse = modelBRT$cv.statistics$deviance.se,
    # estimated cv deviance se
    CorrCV = modelBRT$cv.statistics$correlation.mean,
    #cv correlation
    CorrCVse = modelBRT$cv.statistics$correlation.se,
    #cv correlation se
    AUCcv = modelBRT$cv.statistics$discrimination.mean,
    # cv AUC score
   AUCcvSE = modelBRT$cv.statistics$discrimination.se
) # cv AUC score se
print(t(results))
```

[,1]1.000000e+00 Tree.Complexity Learning.Rate 1.000000e-02 Bag.Fraction 5.000000e-01 Interaction.depth 1.000000e+00 1.000000e-02 Shrinkage N.trees 1.000000e+04 Deviance 3.654408e-01 AUC 1.000000e+00 9.531487e-01 Corr devianceCV 4.390420e-01 devianceCVse 1.584192e-02 CorrCV 9.233031e-01 CorrCVse 6.657569e-03 AUCcv 9.996700e-01 AUCcvSE 3.300000e-04

3.4.10.3 Model output analysis

We can look at the relative contribution of each of the predictor variables. The measures are based on the number of times the variable is selected for splitting, weighted by the improvement of the model as a result of each split averaged across all trees. The relative contribution of each of the variables is scaled so the sum is 100%, with higher numbers indicating stronger influence in the response.

```
# Variables contribution
modelBRT$contributions
```

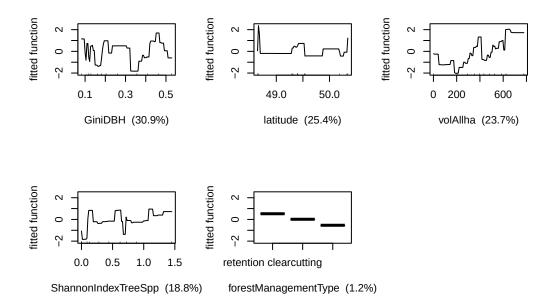
```
var rel.inf
GiniDBH GiniDBH 30.867613
latitude latitude 25.446834
volAllha volAllha 23.695681
ShannonIndexTreeSpp ShannonIndexTreeSpp 18.759858
forestManagementType forestManagementType 1.230014
```

Here we can see that the two variables with the highest influence in the response are GiniDBH and volAllhsa.

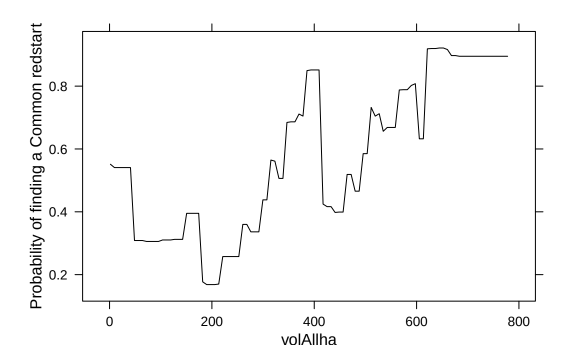
Now we can evaluate the model behavior via partial dependence plots, showing the effect of each of the variables on the response by accounting for the average effects of all other predictors in the model:

```
dismo::gbm.plot(
    modelBRT,
    n.plots = 6,
    plot.layout = c(2, 3),
    write.title = F
)
```

Warning in dismo::gbm.plot(modelBRT, n.plots = 6, plot.layout = c(2, 3), : reducing no of plotted predictors to maximum available (5)



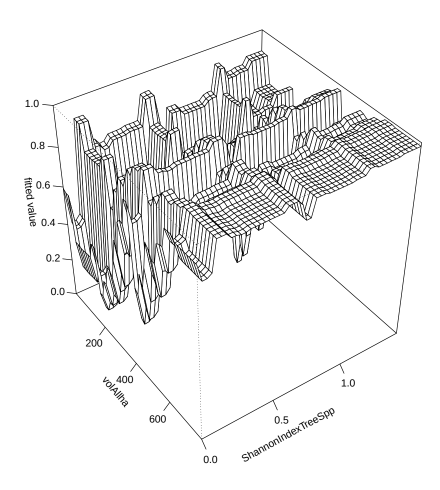
In this partial dependence plots the predictions are on the scale of f(x). In this case, for the Bernoulli loss the returned value is on the log odds scale. You can see how this plot will look by plotting with the function from the package gbm and using the type "response". Since we are interesting in finding out if the forest density has an impact in the presence of the Eurasian treecreeper affected we can have a look to the plot density variable volAllha:



It seems that there is an increase in probability of finding a Eurasian treecreeper with higher forest densities. We could also analyse the interaction effects, of forest diversity and for example forest density. The model predictions can be obtained for each pair of predictor variables, setting all other predictors to their means.

To plot this pairwise interactions we have to do:

dismo::gbm.perspec(modelBRT, 3, 5)



It seems that the highest chances to see a Eurasian tree creeper is in dense forests with highest tree diversity.

3.4.11 Question D - Group 4

• Is the presence of the Eurasian treecreeper affected by forest management?

3.4.11.1 Fitting a BRT in R

In this case we want to assess the occurrence of certain species across the plots. In other words, we want to assess what is the probability of a certain species with biodiversity interest to be present in a plot based on the variables that describe the forest of that plot. In this case the response variable is **certhia** that represents if the Eurasian treecreeper has been observed in this plot or not.

Then you need to select some variables of interest, after you have explored the data you can decide which variables you want to use to fit this model. We are proposing to select the following variables:

- latitude as proxy for plot location or/and climate
- forestManagementType to assess if different management types have different impact in the presence / absence of Great spotted woodpecker.
- volAllha that is the total volume in the plot, as a proxy of how dense the plot is. Higher volumes will mean that the forest is more dense.
- GiniDBH showing how homogeneous the plot is in trees diameters. A value closer to 1 will mean that indicate more structural heterogeneity, lower values indicate more homogeneous plots.

You can create a vector **selVar** in which you add the names of the selected variables. Then you only take those variables from the data that you will use to create the model.

```
# Select variables from the dataset for the model
selVar <- c(
    "certhia",
    "latitude",
    "forestManagementType",
    "volAllha",
    "GiniDBH",
    "ShannonIndexTreeSpp"
)

# Filter the dataset to the selected variables
modelDataSel <- observations[, colnames(observations) %in% selVar]</pre>
```

Unfortunately the amount of that we have in this dataset it is not enough to fit a BRT model for these variables. We are going to do an obviously wrong thing for the shake of being able to demonstrate how to fit a BRT model. In the next code you are going to repeat the same dataset multiple times:

Now it is important to assess if the variables have the right categories. Variables should be type numeric or factor.

summary(modelDataSel)

```
GiniDBH
   latitude
                 forestManagementType
                                          volAllha
Min.
       :48.65
                 Length:495
                                       Min.
                                              : 1.681
                                                          Min.
                                                                  :0.08209
1st Qu.:49.31
                 Class : character
                                       1st Qu.:319.920
                                                          1st Qu.:0.13684
Median :49.40
                 Mode :character
                                       Median: 434.729
                                                          Median :0.20358
Mean
       :49.49
                                       Mean
                                               :425.694
                                                          Mean
                                                                  :0.25683
3rd Qu.:50.19
                                       3rd Qu.:558.355
                                                          3rd Qu.:0.37368
                                       Max.
Max.
       :50.34
                                              :777.882
                                                          Max.
                                                                  :0.52852
{\tt ShannonIndexTreeSpp}
                        certhia
Min.
       :0.000
                            :0.0000
                     Min.
1st Qu.:0.060
                     1st Qu.:0.0000
Median :0.280
                     Median :1.0000
Mean
       :0.392
                            :0.5859
                     Mean
3rd Qu.:0.680
                     3rd Qu.:1.0000
Max.
       :1.450
                            :1.0000
                     Max.
```

```
# Two variables are character, we assign to factor instead:
modelDataSel$forestManagementType <-
    as.factor(modelDataSel$forestManagementType)</pre>
```

In the next step you can see how you can run the model with the selected variables and model parameters. You have a description of the models parameters in the Section 3.3.2 . In this example we are going to use the default parameters for the calibration, where learning rate = 0.01 and tree complexity = 1 and cross-validation = 10-fold. However, the bag fraction is changed from the default value, 0.75, to 0.5. As a family we used the Bernoulli family, because we are predicting presence/absence per plot. These data have 495 plots, comprising 290 presence records for the Eurasian treecreeper. You can check these numbers by doing:

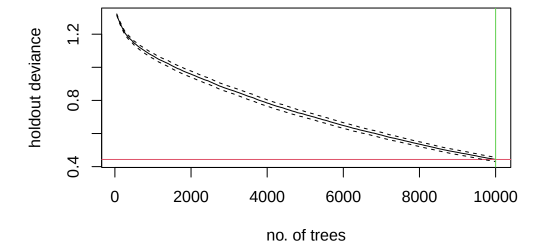
```
table(modelDataSel$certhia)
```

0 1 205 290

As a first guess you could decide there are enough data to model interactions of reasonable complexity, and a lr of about 0.01 could be a reasonable starting point. You can use the model creation function that steps forward and identifies the optimal number of trees (nt) by doing this:

```
family <- "bernoulli"
tc = 1
          # tree complexity
lr = 0.01 # learning rate-shrinkage
bag = 0.5 \# bag fraction
modelBRT <- dismo::gbm.step(</pre>
    data = modelDataSel,
    #indices of predictor variables in data
    gbm.x = 1:5,
    #index of response variable in data:
    gbm.y = 6,
    family = family,
    tree.complexity = tc,
    learning.rate = lr,
    bag.fraction = bag
)
```

certhia, d - 1, lr - 0.01



Running a model such as that described above writes progress reports to the screen, makes a graph, and returns an object containing a number of components. The R console results reports a brief model summary all the values are also retained in the model object.

The model is built with the default 10-fold cross-validation (CV). In the plotted graph the solid black curve is the mean, and the dotted curves 1 standard error, for the changes in predictive deviance (i.e., as measured on the excluded folds of the CV). The red line shows the minimum of the mean, and the green line the number of trees at which that occurs. The final model that is returned in the model object is built on the full data set, using the number of trees identified as optimal.

Ideally, you should invest time in modifying the parameters and find the parameters that provide the models with the minimum deviance resulting from the best combination of bag, tree complexity and learning rate values. For the shake of limited timing, we will only test here the default values.

3.4.11.2 Model behaviour

You can summarize the model parameters used and the cross validation statistics from the fitted model by doing this:

```
# We make a table with the summary statistics
results <- data.frame(
    # Model parameters
    Tree.Complexity = modelBRT$gbm.call$tree.complexity,
    Learning.Rate = modelBRT$gbm.call$learning.rate,
    Bag.Fraction = modelBRT$gbm.call$bag.fraction,
    Interaction.depth = modelBRT$interaction.depth,
    Shrinkage = modelBRT$shrinkage,
    N.trees = modelBRT$n.trees,
    # Cross validation statistics
    ## mean total deviance
    Deviance = modelBRT$self.statistics$mean.resid,
    # mean residual deviance
    AUC = modelBRT$self.statistics$discrimination,
    # training data AUC score
    Corr = modelBRT$self.statistics$correlation,
    # training data correlation
```

```
## Cross Validation statistics
    # We calculate each statistic within each fold (at the identified optimal number
    # of trees that is calculated on the mean change in predictive deviance over all fold
    #then present here the mean and standard error of those fold-based statistics.
    devianceCV = modelBRT$cv.statistics$deviance.mean,
    # estimated cv deviance
    devianceCVse = modelBRT$cv.statistics$deviance.se,
    # estimated cv deviance se
    CorrCV = modelBRT$cv.statistics$correlation.mean,
    #cv correlation
    CorrCVse = modelBRT$cv.statistics$correlation.se,
    #cv correlation se
    AUCcv = modelBRT$cv.statistics$discrimination.mean,
    # cv AUC score
   AUCcvSE = modelBRT$cv.statistics$discrimination.se
) # cv AUC score se
print(t(results))
```

[,1]1.000000e+00 Tree.Complexity Learning.Rate 1.000000e-02 Bag.Fraction 5.000000e-01 Interaction.depth 1.000000e+00 1.000000e-02 Shrinkage N.trees 1.000000e+04 Deviance 3.641224e-01 AUC 1.000000e+00 9.542670e-01 Corr devianceCV 4.432030e-01 1.257987e-02 devianceCVse CorrCV 9.202141e-01 CorrCVse 6.692980e-03 AUCcv 9.993500e-01 4.982748e-04 AUCcvSE

3.4.11.3 Model output analysis

We can look at the relative contribution of each of the predictor variables. The measures are based on the number of times the variable is selected for splitting, weighted by the improvement of the model as a result of each split averaged across all trees. The relative contribution of each of the variables is scaled so the sum is 100%, with higher numbers indicating stronger influence in the response.

```
# Variables contribution
modelBRT$contributions
```

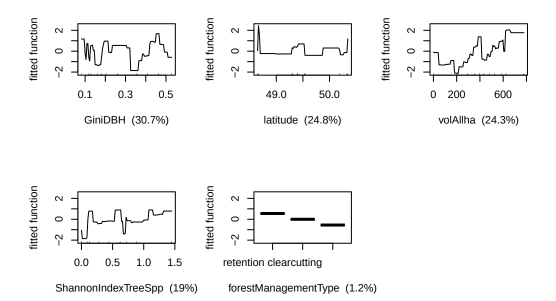
```
var rel.inf
GiniDBH GiniDBH 30.687490
latitude latitude 24.779168
volAllha volAllha 24.342973
ShannonIndexTreeSpp ShannonIndexTreeSpp 18.955284
forestManagementType forestManagementType 1.235085
```

Here we can see that the two variables with the highest influence in the response are GiniDBH and volAllhsa.

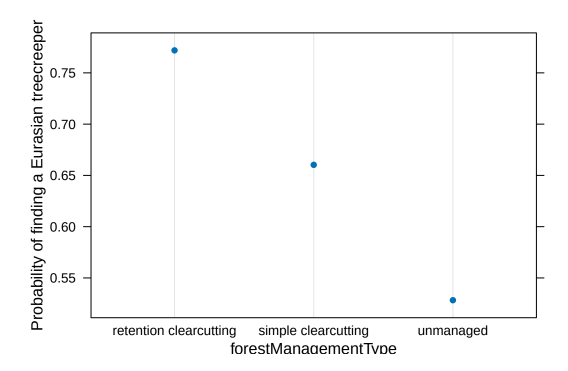
Now we can evaluate the model behavior via partial dependence plots, showing the effect of each of the variables on the response by accounting for the average effects of all other predictors in the model:

```
dismo::gbm.plot(
    modelBRT,
    n.plots = 6,
    plot.layout = c(2, 3),
    write.title = F
)
```

Warning in dismo::gbm.plot(modelBRT, n.plots = 6, plot.layout = c(2, 3), : reducing no of plotted predictors to maximum available (5)



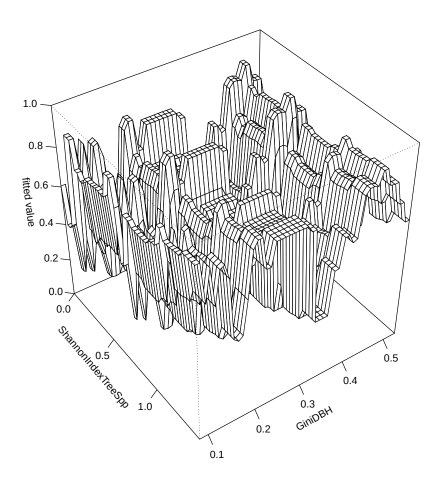
In this partial dependence plots the predictions are on the scale of f(x). In this case, for the Bernoulli loss the returned value is on the log odds scale. You can see how this plot will look by plotting with the function from the package gbm and using the type "response". Since we are interesting in finding out if the forest density has an impact in the presence of the Eurasian treecreeper affected we can have a look to the plot density variable volAllha:



It seems that there is a small difference in the probability of finding the Eurasian treecreeper different management strategies. We could also analyse the interaction effects, of forest diversity and forest structual diversity. The model predictions can be obtained for each pair of predictor variables, setting all other predictors to their means.

To plot this pairwise interactions we have to do:

dismo::gbm.perspec(modelBRT, 5, 4)



It seems that there are not clear patterns in the combined effect of forest structure diversity and forest tree species diversity.

4 References

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