# Package 'MadingleyR'

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Type Package

Title MadingleyR: General Ecosystem Model

Version 1.0.0
Author
Maintainer The package maintainer:
<b>Description</b> The MadingleyR package is wrapper around the C++ code of the Madingley model. It was intended as an accessible and transparent tool to lower barrier of using the Madingley model, thereby broadening the Madingley comminity.
LazyData true
Imports sp, raster, rgdal
Included functions:
list_output_paths
madingley_init
madingley_inputs
madingley_plot
madingley_run
plot_densities
plot_foodweb

plot\_spatialabundances11plot\_spatialbiomass12plot\_timelines13plot\_trophicpyramid14

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list\_output\_paths

MadingleyR: Get the output file paths

### **Description**

This functions returns the files names and/or paths of the files written by the MadingleyR package. It only works when an output folder is specified in the madingley\_run() function. This aids the reading of files for processing at a later stage (see example below).

# Usage

```
list_output_paths(madingley_data, full_path = TRUE)
```

### **Arguments**

```
madingley_data output generated by madingley_run()

full_path whether the function should return the full path (full_path = TRUE) or the names of the files only (full_path = FALSE)
```

```
# Set output folder
out_dir = 'path/to/outputs'
# Set spatial window
spatial\_window = c(31, 35, -5, -1)
# Call the madingley_init function (default run)
m_data = madingley_init(spatial_window = spatial_window)
# Run the model
m_data2 = madingley_run(out_dir = out_dir,
                        madingley_data = m_data,
                        years = 4)
# Get the full paths of the output files
output_paths = list_output_paths(madingley_data = m_data2)
# Get names of the output files
output_names = list_output_paths(madingley_data = m_data2, full_path = F)
# Use output_paths to load a file written
cat(output_paths$cohort_properties[1])
BinnedCohortStatistics1 = read.csv(output_paths$cohort_properties[1])
head(BinnedCohortStatistics1)
```

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madingley\_init

MadingleyR: Initialise the Madingley model

#### **Description**

This function generates the cohort and stock data.frames (initialisation parameters) needed to run madingley\_run().

### Usage

```
# Simple default init
mdata = madingley_init(spatial_window)
# All init parameters manually
mdata = madingley_init(spatial_window, cohort_def, stock_def, spatial_input, max_cohorts)
```

#### **Arguments**

spatial\_window a vector holding coordinates that specify the spatial model domain, specified in

the following order: minimum longitude, maximum longitude, minimum latitude, maximum latitude. For example spatial\_window = c(31,35,-5,-1)

cohort\_def a data.frame holding the cohort definitions used for initialising the model. The

template or default data.frame can be loaded using: madingley\_inputs('cohort definition'). If the cohort\_def is not specified, madingley\_init() loads the cohort definitions internally using the madingley\_inputs('cohort definition') function

call

stock\_def a data.frame holding the stock definitions used for initialising the model. The template or default data.frame can be loaded using: madingley\_inputs('stock

definition'). If the stock\_def is not specified, madingley\_init() loads the stock

definitions internally using the madingley\_inputs('stock definition') function call

spatial\_input an R list that holds all of the spatial inputs required for initialising the Madingley model. The default spatial inputs can be loaded using: madingley\_inputs('spatial inputs'). If the spatial\_input is not specified, madingley\_init() loads the inputs

internally using the madingley\_inputs("spatial inputs') function call

max\_cohort a single integer holding the maximum number of cohorts allowed per spatial grid cell. The Madingley model mergers similar cohorts if they are highly similar and

the maximum number of cohorts is exceeded. This is done for computational reasons (keeping the number of interactions at a reasonable level). The Madingley model merges cohorts if they are highly similar and the maximum number of cohorts is exceeded. Lowering this value will increase model performance at expenses of resolution of cohort diversity. 1000 are normally used, but 500 may be sufficient for preliminary trials. Please note, that setting this value to e.g. 500 in madingley\_init() makes sure that max 500 cohorts per grid cell are created. If in a consecutive madingley\_run() max\_cohort is set to e.g. 1000 the maximum number of cohorts (500 cohorts/grid cell) will increase during the simulation run until the new maximum (1000 cohorts/grid cell) is reached. For consistency it may be preferred to set max\_cohort to the same number in a madingley\_init()

call and the consecutive madingley\_run() simulation.

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#### **Examples**

madingley\_inputs

MadingleyR: Load the default Madingley

# **Description**

This function loads the default MadingleyR input parameters in case the model parameters need to be checked or modified depending on the simulation scenario. The function takes one string input. The available options can be printed to the R console with madingley\_inputs(), see example code below.

### Usage

```
madingley_inputs() # prints all available options
sptl_inp = madingley_inputs("spatial inputs") # loads the default spatial inputs
chrt_def = madingley_inputs("cohort definition") # loads the default cohort definitions
stck_def = madingley_inputs("stock definition") # loads the default stock definitions
mdl_prms = madingley_inputs("model parameters") # loads the default model parameters
```

# Arguments

input\_type

a string specifying the input type to load. The options are: "spatial inputs", "cohort definition", "stock definition" or "model parameters". Running the function without any string will print all available options as shown in the usage above or example below.

```
# Print all the available options to load to the R console
madingley_inputs()

# Load spatial parameters
sptl_inp = madingley_inputs(input_type = "spatial inputs")

# Load default cohort property definitions
```

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```
chrt_def = madingley_inputs(input_type = "cohort definition")
# Load default stock property definitions
stck_def = madingley_inputs(input_type = "stock definition")
# Load default model parameters
mdl_prms = madingley_inputs(input_type = "model parameters")
# Set spatial model domain
spatial\_window = c(31, 35, -5, -1)
## -----##
# Change parameters accordingly #
##----##
# Call the madingley_init function
m_data = madingley_init(cohort_def = chrt_def,
                      stock_def = stck_def,
                      spatial_inputs = sptl_inp,
                      max\_cohort = 500)
# Run the madingley model
m_data2 = madingley_run(madingley_data = m_data,
                       years = 5,
                      max\_cohort = 500,
                       cohort_def = chrt_def,
                       stock_def = stck_def,
                       spatial_inputs = sptl_inp,
                       model_parameters = mdl_prms)
```

madingley\_plot

MadingleyR: Create MadingleyR plots

# Description

Create all output plots included in the MadinleyR package. This function is intended as a tool to quickly visualize the results. All resulting plots can also be made individually by using the following functions:

```
- plot_densities()
```

- plot\_foodweb()
- plot\_spatialabundances()
- plot\_spatialbiomass()
- plot\_timelines()
- plot\_trophicpyramid()

### Usage

```
madingley_plot(madingley_data)
```

### **Arguments**

madingley\_data output generated by madingley\_run()

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### **Examples**

```
# Print all the available options to load to the R console
madingley_inputs( )
# Load spatial parameters
sptl_inp = madingley_inputs("spatial inputs")
# Load default cohort property definitions
chrt_def = madingley_inputs("cohort definition")
# Load default stock property definitions
stck_def = madingley_inputs("stock definition")
# Load default model parameters
mdl_prms = madingley_inputs("model parameters")
# Set spatial model domain
spatial\_window = c(31, 35, -5, -1)
# Call the madingley_init function
m_data = madingley_init(cohort_def = chrt_def,
                        stock_def = stck_def,
                        spatial_inputs = sptl_inp,
                        spatial_window = spatial_window,
                        max\_cohort = 500)
# Run the madingley model using the m_data as input
m_data2 = madingley_run(madingley_data = m_data,
                        years = 5,
                        max\_cohort = 500,
                        cohort_def = chrt_def,
                        stock_def = stck_def,
                        spatial_inputs = sptl_inp,
                        model_parameters = mdl_prms)
# Create the MadingleyR plots
madingley_plot(m_data2)
```

madingley\_run

MadingleyR: Run the Madingley model

# Description

This function runs the Madingley model using outputs generated by madingley\_init() or by a previous model run.

### Usage

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> stock\_def = stck\_def, spatial\_inputs = sptl\_inp, model\_parameters = mdl\_prms, dispersal\_off = FALSE, silenced = FALSE)

# **Arguments**

out\_dir

path to the folder used for writing the Madingley simulation outputs (csv file). If out\_dir is not specified, madingley\_run() writes the outputs to the temporary R folder. The outputs in this folder are required for the plotting functions included in MadingleyR. If the outputs need to be analysed manually or plots need to be made at a later time, please specify the output folder to make sure your files are

madingley\_data madingley output list generated by madingley\_init() or by a previous model run (output list of madingley\_run())

years

a single integer to set the simulation length (in years)

cohort\_def

a data.frame holding the cohort definitions used for initialising the model. The template or default data.frame can be loaded using: madingley\_inputs('cohort definition'). If the cohort def is not specified, madingley run() loads the cohort definitions internally using the madingley inputs('cohort definition') function

stock\_def

a data.frame holding the stock definitions used for initialising the model. The template or default data.frame can be loaded using madingley\_inputs('stock definition'). If the stock\_def is not specified, madingley\_run() loads the stock definitions internally using the madingley\_inputs('stock definition') function call

spatial\_input

an R list that holds all the spatial inputs required for initialising the Madingley model. The default spatial inputs can be loaded using madingley\_inputs('spatial inputs'). If the spatial\_input is not specified, madingley\_run() loads the inputs internally using the madingley\_inputs("spatial inputs") function call

model\_parameters

a data.frame with the default model parameters. The function call madingley\_inputs('model parameters') loads the model inputs alongside a description corresponding to each parameters. If the model\_parameters argument is not specified, madingley run() loads the default model parameters internally using the madingley\_inputs('model parameters') function call

max cohort

a single integer holding the maximum number of cohorts allowed per spatial grid cell for computational reasons (keeping the number of interactions at a reasonable level). The Madingley model merges cohorts if they are highly similar and the maximum number of cohorts is exceeded. Lowering this value will increase model performance at expenses of resolution of cohort diversity. 1000 are normally used, but 500 may be sufficient for preliminary trials.

output\_timestep

by default the model writes the outputs during all (12) months of the last simulation year (years - 1). This can be altered by using the output\_timestep argument. The output\_timestep argument is formatted as a vector containing four integers: 1) the year to output the log10 binned cohort properties 2) the year to output the full cohort properties 3) the year to output the log10 binned feeding interactions (required by the food-web plot) and 4) the year to output the full stock (autotroph) files. Please note, exporting the results will always start from the 8 madingley\_run

year set and continues until the end of the simulation. For example, when setting output\_timestep = c(5,3,5,4) and years = 5 in Madingley\_run():

- the log 10 binned cohort properties (set to 5) are never written (years = output\_timestep = 5)
- the full cohort properties (set to 3) are written during the last two years (after the 3rd year until the end of the simulation)
- the log10 binned feeding interactions (set to 5) are never written (years = output\_timestep = 5)
- the full stock files (set to 4) are written only during the last year of the simulation (after the 4th year).

Be careful outputting many large files, it can easily fill up your hard disk. Also note that outputting the feeding interactions significantly slows down the model runtime, only use this option if needed! The fact that outputting results will start from the year set and continue to the end of the simulation, can produce a lot of results when outputs are required somewhere halfway the simulation. Let's say in a 100-year simulation we need to output the model results at year = 50, we will end up with around 50 years of outputs (from year 50 to year 100). For these cases I recommend to run the model for 50 years, output the results at the final simulation year and continue the simulation for another 50 years using a new madingley\_run() instance using the previous 50 year simulation as an input (see final example below).

dispersal\_off

If TRUE turns the dispersal off between grid cells in the Madingley model. By default dispersal\_off is set to FALSE, thereby allowing dispersal between grid cells

silenced

When set to TRUE it suppresses the outputs printed to the R console. By default it is set to FALSE

```
# Print all the available options to load to the R console
madingley_inputs( )
# Load spatial parameters
sptl_inp = madingley_inputs("spatial inputs")
# Load default cohort property definitions
chrt_def = madingley_inputs("cohort definition")
# Load default stock property definitions
stck_def = madingley_inputs("stock definition")
# Load default model parameters
mdl_prms = madingley_inputs("model parameters")
# Set spatial model domain
spatial\_window = c(31, 35, -5, -1)
## -----##
# Change parameters accordingly
# e.g.: sptl_inp, chrt_def, stck_def, mdl_prms #
# Call the madingley_init function
m_data = madingley_init(cohort_def = chrt_def,
                       stock_def = stck_def,
```

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```
spatial_inputs = sptl_inp,
                       spatial_window = spatial_window,
                      max\_cohort = 500)
# Change parameters accordingly
# e.g.: m_data sptl_inp, chrt_def, stck_def, mdl_prms #
# Run the madingley model using the m_data as input
m_data2 = madingley_run(madingley_data = m_data,
                      years = 5,
                      max\_cohort = 500,
                       cohort_def = chrt_def,
                       stock_def = stck_def,
                       spatial_inputs = sptl_inp,
                      model_parameters = mdl_prms)
# Change parameters accordingly
# e.g.: m_data2 sptl_inp, chrt_def, stck_def, mdl_prms #
# Continue simulation run using the m_data2 as input
m_data3 = madingley_run(out_dir = "C:/MadingleyOutputs/",
                      madingley_data = m_data2,
                       years = 50,
                       max\_cohort = 500,
                       cohort_def = chrt_def,
                       stock_def = stck_def,
                       spatial_inputs = sptl_inp,
                      model_parameters = mdl_prms)
## -----##
# Export outputs for a single year halfway the simulation duration
##----##
# Call the madingley_init function
m_data1 = madingley_init(spatial_window = spatial_window, max_cohort = 500)
# Run for 50 years and export results during last simulation year
m_data2 = madingley_run(out_dir = "C:/MadingleyOutputs/", madingley_data = m_data1,
       years = 50, max_cohort = 500, output_timestep = c(49,49,49,49))
# Run for an additional 50 years and export during last simulation year
m_data3 = madingley_run(out_dir = "C:/MadingleyOutputs/", madingley_data = m_data2,
       years = 50, max_cohort = 500, output_timestep = c(49,49,49,49))
```

plot\_densities

MadingleyR: Plot abundance or biomass density against body mass

### **Description**

It plots the abundance or biomass density against (log10) body mass.

# Usage

```
plot_densities(m_data2, weighted = 'biomass' , xlim=c(-4, 5)) plot_densities(m_data2, weighted = 'abundance' , xlim=c(-4, 5))
```

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### Arguments

madingley\_data madingley output list returned by madingley\_run()

weighted select either 'biomass' or 'abundance' (pass as string)

xlim min and max log10 body mass plotted

ylim min and max density

col the colours used (length needs to be equal to the number of groups plotted)

plot If TRUE it generates the plot, if FALSE it returns the data output formatted for

plotting which can be used for further analysis

### **Examples**

```
# set model params
spatial_window = c(31,35,-5,-1) # Serengeti

# init model
m_data = madingley_init(spatial_window = spatial_window)

# run model
m_data2 = madingley_run(madingley_data = m_data, years = 3)

# make plots
plot_densities(m_data2, weighted = 'biomass' , xlim=c(-4, 5))
plot_densities(m_data2, weighted = 'abundance' , xlim=c(-4, 5))
```

plot\_foodweb

MadingleyR: Make food web plot

# Description

It plots the log10-binned body mass food web.

# Usage

```
plot_foodweb(m_data2, max_flows = 5, sample_data = 100)
```

# Arguments

madingley\_data madingley output list returned by madingley\_run()
max\_flows set the maximum number of flows per node

color TRUE (color version of the plot) or FALSE (grey scale version of the plot)

sample\_data By default the function takes all data into consideration (sample\_data = 100) but

can be computationally intensive. To speed up the visualization of the results it might be useful to only consider a random sample of the data, e.g. sample\_data = 30 will sample 30% of the data randomly and use it to make the food-web plot

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#### **Examples**

```
# set model params
spatial_window = c(31,35,-5,-1) # Serengeti

# init model
m_data = madingley_init(spatial_window = spatial_window)

# run model
m_data2 = madingley_run(madingley_data = m_data, years = 3)

# make plot
plot_foodweb(m_data2, max_flows = 5, sample_data = 100)
```

plot\_spatialabundances

MadingleyR: Create spatial abundance plot

# Description

It plots the spatial abundances of functional groups spatially.

# Usage

```
plot_spatialabundances(m_data2, functional_filter = TRUE)
```

### Arguments

# Input arguments

madingley output list returned by madingley\_run()

functional\_filter

madingley\_data TRUE or FALSE

plot If set to TRUE it makes the plot, if set to FALSE the function returns a raster or

raster brick (default = TRUE)

Visual plotting arguments

box TRUE or FALSE (default = TRUE)

axes TRUE or FALSE (default = TRUE)

names plot titles: by default it uses the numbering of the functional groups following

the cohort definitions (see madingley\_inputs('cohort definition'))

cex\_legend\_title

font size of legend title (default = 1)

cex\_axis font size of legend numbering (default = 1.3)

legend\_mar the legend margin (default = 20)

 ${\tt distance\_legend\_title}$ 

the distance from the legend to the legend title (default = 4)

legend\_width the width of the legend bar (default = 2.5)

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#### **Examples**

```
# set model params
spatial_window = c(31,35,-5,-1) # Serengeti
# init model
m_data = madingley_init(spatial_window = spatial_window)
# run model
m_data2 = madingley_run(madingley_data = m_data, years = 3)
# make plot
plot_spatialabundances(m_data2, functional_filter = TRUE)
```

plot\_spatialbiomass

MadingleyR: Create spatial biomass plot

### **Description**

It plots the spatial biomass per grid cell of functional groups spatially.

### Usage

```
plot_spatialbiomass(m_data2, functional_filter = TRUE)
```

# Arguments

### Input arguments

madingley output list returned by madingley\_run()

functional\_filter

 ${\tt madingley\_data} \ \ TRUE \ or \ FALSE$ 

plot If set to TRUE it makes the plot, if set to FALSE the function returns a raster or

raster brick (default = TRUE)

Visual plotting arguments

box TRUE or FALSE (default = TRUE)

axes TRUE or FALSE (default = TRUE)

names plot titles: by default it uses the numbering of the functional groups following

the cohort definitions (see madingley\_inputs('cohort definition'))

cex\_legend\_title

font size of legend title (default = 1)

cex\_axis font size of legend numbering (default = 1.3)

 $legend_mar$  the legend margin (default = 20)

 ${\tt distance\_legend\_title}$ 

the distance from the legend to the legend title (default = 4)

legend\_width the width of the legend bar (default = 2.5)

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#### **Examples**

```
# set model params
spatial_window = c(31,35,-5,-1) # Serengeti

# init model
m_data = madingley_init(spatial_window = spatial_window)

# run model
m_data2 = madingley_run(madingley_data = m_data, years = 3)

# make plot
plot_spatialbiomass(m_data2, functional_filter = TRUE)
```

plot\_timelines

MadingleyR: Create timeline plot

### **Description**

It creates a plot the total biomass of functional groups over time.

# Usage

```
plot_timelines(m_data2, select = 'functional groups')
plot_timelines(m_data2, select = 'feeding guilds')
```

# Arguments

### Input arguments

```
madingley output list returned by madingley_run()

meddengley_data can be set to: 'functional groups' or 'feeding guilds'

plot If FALSE it returns a data.frame with the time line data formatted for plotting (default = TRUE).
```

```
# set model params
spatial_window = c(31,35,-5,-1) # Serengeti

# init model
m_data = madingley_init(spatial_window = spatial_window)

# run model
m_data2 = madingley_run(madingley_data = m_data, years = 3)

# make plots
plot_timelines(m_data2, select = 'functional groups')
plot_timelines(m_data2, select = 'feeding guilds')
```

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plot\_trophicpyramid MadingleyR: Creates trophic pyramide plot

# **Description**

The function plots a trophic pyramid from the Madingley simulation outputs.

### Usage

```
plot_trophicpyramid(m_data2)
```

### **Arguments**

```
madingley_data madingley output list returned by madingley_run()

flow_round number of decimals for rounding the quantification of flows (default = 2)

mass_round number of decimals for rounding the quantification of mass (default = 2)

col vector including the colours for each trophic level

text_col the colour of the text quantifying the mass of each trophic level
```

```
# set model params
spatial_window = c(31,35,-5,-1) # Serengeti

# init model
m_data = madingley_init(spatial_window = spatial_window)

# run model
m_data2 = madingley_run(madingley_data = m_data, years = 3)

# make plot
plot_trophicpyramid(m_data2)
```