CODING PRACTICES

Life with R



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Assign recognizable, concise names to your objects!

Let's start with the object MyData - this can be anything. Let's assume it is a raster of NDVI values during the year 1982. How do we come up with a better name?

Specificity in your naming helps you keep track of your data in your code:

Classes of objects determine how they get handled. Add them as a suffix:

The same goes for **loop indicators**! Avoid single-letter indicators and instead use something like Iter_Years - an indicator *iterating* (Iter) over years.

Also, always use <- for assigning objects!

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Use **spaces** to improve readability of your code:

```
c(1,2,rep(3,5)) # bad
c(1, 2, rep(3, 5)) # good
```

Insert **line breaks** to improve workflow. I like to break multiple lines of one function call into logical blocks. For example, in ggplot2, the first line holds data, the second contains aesthetics, the third adds a layer, and so on.

To avoid side-scrolling, use the automatic soft-wrap function in RStudio.

Be **consistent** in your coding style. Found a tidyverse solution to your problem online but don't use tidyverse syntax in the rest of your document? Adapt the solution to be in the same style as the rest of your coding!

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Don't be overly proud of your coding skills. Comment everything!

Bad comments:

- ambiguous
- sparse
- # Data Manipulation

Good comments:

- specify (what is the code doing)
- justify (why is it being done)
- # Z-Score calculation for comparability

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RMarkdown

Using Rmarkdown for your research comes with a multitude of advantages:

- Entire workflow in one program (RStudio)
- Great capabilities to present code
- Research and reports reproducible at the click of one button
- Combines R functionality and LaTEX formatting (if desired)
- Consistent formatting
- Clear presentation of code
- Dynamic documents (you can generate various output document types)
- Applicable for **almost all document types** you may desire as an output (e.g. manuscripts, presentations, posters, etc.)

Header

The **Head** is used as an information statement at the top of your code document that informs the user of the contents, author, and (sometimes) date of the last edit on said document:

Personally, I don't use the edit argument in my heads. Instead, I rely on *version* control

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Preamble

The **Preamble** is where you set up the most important parameters/guidelines for your coding script. Personally, I *highly* recommend to make your first line in the preamble read rm(list=ls()).

This is also where you load **packages**. I recommend doing so as follows:

```
install.load.package <- function(x) {
   if (!require(x, character.only = TRUE))
        install.packages(x, repos = "http://cran.us.r-project.org")
   require(x, character.only = TRUE)
}</pre>
```

This functions *checks* whether the package is already installed. If it is, it will be *loaded*. If it is not, it will be *installed* from CRAN and subsequently loaded. You use it for multiple packages like this:

```
package_vec <- c("ggplot2", "foreach", "doParallel")
sapply(package_vec, install.load.package)</pre>
```

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They do so by

- collapsing code
- adding a table of contents

```
| Separation | Sep
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```
6 - #### Libraries #### 8 | 1brary(tigyerse) 9 | 1bbrary(tigyerse) 9 | 1bbrary(fuzzysin) 11 | 6 | 1brary(vegan) 11 | 6 | 1brary(vegan) 12 | 1brary(vegan) 13 | 1brary(vegan) 14 | 1brary(vegan) 15 | 1brary(vegan) 16 | 1brary(vegan) 17 | 1brary(vegan) 17 | 1brary(vegan) 18 | 1brary
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Reproducibility

Reproducibility is paramount in science.

Working directories

It is common to set a base directory at the beginning of your code and refer to sub-directories from there. We can *do better*:

- getwd() in a project will yield the directory of the project file
- the package here offers more functions for this
- set paths to files or direcotires using the file.path() function to avoid cross-platform differences in folder path

Randomness

Random processes (e.g. sample(), rnorm(), etc.) in R are always pseudo-random (their calculation is based on your computer time). We can make sure the result of a random process is always the same between researchers by using:

-set.seed()

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Functions & Sourcing

Big projects lend themselves well to a multi-document workflow:

Functions

User-defined functions are a great way of *soft-coding* your analyses to repeat them for different input parameters and I highly recommend doing so all the time. Make sure your functions are:

- internally consistent
- well-documented
- easy to understand

Sourcing

Splitting your code into multiple documents is a great idea to ensure your project remains structured and easy to handle. You can use the source() function to combine those documents. Keep in mind that you should:

- use sensible file names
- sourced functions still need to be called

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Progress Bar

A progress bar is a great way of keeping you updated on how your code is progressing so far. It is especially useful when your code involves *loops*:

```
Data vec <- 1:100 # a vector on integers from 1 to 100
# creating progress bar
ProgBar <- txtProgressBar(min = 0, max = length(Data_vec),</pre>
                           stvle = 3)
# looping over contents of Data vec
for(Iter_ProgBar in 1:length(Data_vec)) {
  setTxtProgressBar (ProgBar, Iter ProgBar) # updating ProgBar
} # end of Data vec loop
```

Estimators

Estimators are a great way to know when to come back to your computer or server and check up on your data, code, and results. They are most useful in *loop* based approaches as they need a baseline for the estimation:

Yes, I did work on this presentation past midnight.

Parallel Processing

Got a big data approach where you carry out the same kind of analysis for many different species or data sets? It's taking forever to complete? **Parallel processing** has your back! By default, $\mathbb R$ only uses one of the cores in your computer or server for processing code. Use this, to make use of all the computational power:

```
library(doParallel) # for registering clusters
library(foreach) # for parallel processing
Cores <- detectCores() # identify the number of cores in your machine
cl <- makeCluster(Cores) # create virtual cluster
registerDoParallel(cl) # register cluster of cores
# parallel processing
foreach(Iter_Par = 1:length(Data_vec)) %dopar% {
    # your function here
} # end of parallel processing
stopCluster(cl) # stop cluster</pre>
```