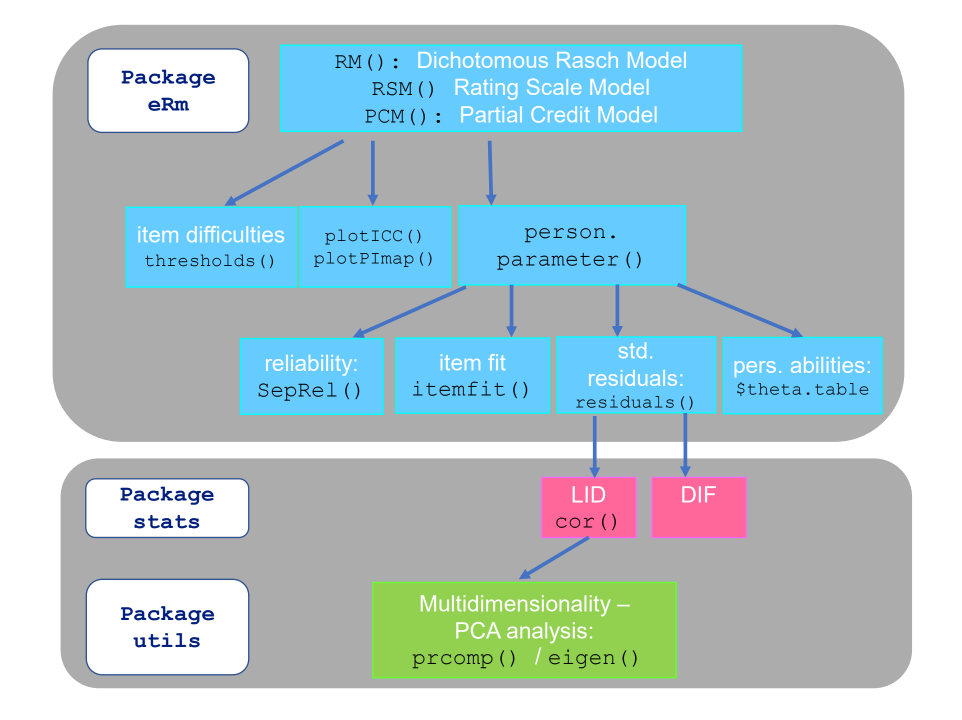
# Getting Started with WHOMDS

R-Package for Analysis of the MDS-Data



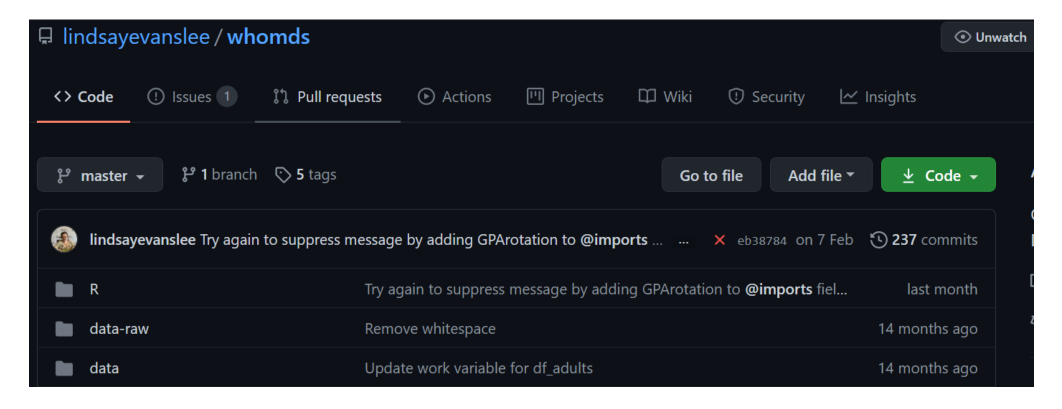
# Github: WHOMDS

- The WHOMDS is an R-package with functions to run MDS analyses with R.
- Package Maintainer: Lindsay Lee
- WHOMDS facilitates the Rasch analysis of the MDS-metrics
- R-studio should be installed.
- Make sure that you have the latest R-version
- The package is on Github, a repository that facilitate sharing and working on code, not only R.

# **WHOMDS**

#### https://github.com/lindsayevanslee/whomds

Press the green button and «download the zip» on your computer



### README.rmd file

 Run the README.rmd file using the knit button and select 'Knit to github\_document'

```
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### README.rmd file

 The README file indicates that WHOMDS can be charged directly from the Github.

#### Run in the R-Console:

```
devtools::install_github("lindsayevanslee/whomds",
build vignettes = TRUE)
```

- A few packages need to be downloaded ahead, this can take some time.
- Let's Switch to R and R-Studio and see how it works

