Tutorial for R microeco package (v1.7.0)

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## Chapter 1

## Background

R language (?) and its packages ecosystem are wonderful tools for data analysis. In community ecology, a series of packages are available for statistical analysis, such as vegan (?), ape (?) and picante (?). However, with the development of the high-throughput sequencing techniques, the increasing data amount and complexity of studies make the data mining in microbiome a challenge. There have been some R packages created specifically for the statistics and visualization of microbiome data, such as phyloseq (?), microbiome (https://github.com/microbiome/microbiome), (http://www.github.com/umerijaz/microbiomeSeq), (https://github.com/KasperSkytte/ampvis2), MicrobiomeR(https: //github.com/vallenderlab/MicrobiomeR), theseus (?), rANOMALY (?),microbial (https://github.com/guokai8/microbial), tidyMicro plicon (https://github.com/microbiota/amplicon), **MicrobiotaProcess** (https://github.com/YuLab-SMU/MicrobiotaProcess) and so on. tion, some web tools associated with R language are also useful for microbiome data analysis, such as Shiny-phyloseq (?), MicrobiomeExplorer (?), animalcules (?) and Namco (?). Even so, researchers still lack a flexible, comprehensive and modularized R package to analyze and manage the data fast and easily. Based on this background, we created the R microeco package (?) (https://github.com/ChiLiubio/microeco). Besides, we also developed the file2meco package (https://github.com/ChiLiubio/file2meco) for the data input from some famous tools easily.