## Installations:

We will be using the miaverse and miaViz R packages for this tutorial (<a href="https://microbiome.github.io/">https://microbiome.github.io/</a>)

To install the needed packages simply do:

```
BiocManager::install("mia")
BiocManager::install("microbiome/miaViz")
```

## Data:

The data can be loaded from the mia package itself, we will be using the GlobalPatterns dataset, which comes from this article: <a href="https://www.pnas.org/doi/full/10.1073/pnas.1000080107">https://www.pnas.org/doi/full/10.1073/pnas.1000080107</a>, which looked at the the microbial communities from 25 environmental samples and three known "mock communities" at an average depth of 3.1 million reads per sample in order to identify global patterns of 16S rRNA diversity at a depth of millions of sequences per sample (Caporaso, 2010).

To access the data all you need to do is:

```
library(mia)
data("GlobalPatterns", package="mia")
```

To both test that you have correctly installed the package, and have the wanted data, simply print out GlobalPatterns:

GlobalPatterns

And make sure that this is the result you get:

```
## class: TreeSummarizedExperiment
## dim: 19216 26
## metadata(0):
## assays(1): counts
## rownames(19216): 549322 522457 ... 200359 271582
## rowData names(7): Kingdom Phylum ... Genus Species
## colnames(26): CL3 CC1 ... Even2 Even3
## colData names(7): X.SampleID Primer ... SampleType Description
## reducedDimNames(0):
## mainExpName: NULL
## altExpNames(0):
## rowLinks: a LinkDataFrame (19216 rows)
## rowTree: 1 phylo tree(s) (19216 leaves)
## colLinks: NULL
## colTree: NULL
```