BIOL432 Assignment 4 - Caroline Tang (20115082)

Project info

Date: Jan 31, 2022

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Data science

Load libraries

```
##
## Attaching package: 'dplyr'
## The following objects are masked from 'package:stats':
##
## filter, lag
## The following objects are masked from 'package:base':
##
## intersect, setdiff, setequal, union
Load in measurements data from week 1 as MData
```

Create a new dataset that is a copy of MData but with Sp instead of Organism, formatted as (G. species)

```
(short <- MData %>% mutate(Sp = gsub("(\w)\w+\s+(\w+)", "\1. \2", MData$Organism)) %>% select(Sp, Limb.length, Limb.width, Observer))
```

```
Sp Limb.length Limb.width Observer
##
## 1
            P. lotor
                               1
                                        1.0 Olivier R
        P. pygmaeus
                               2
                                        1.0
                                                Han S
## 3 P. cancrivorus
                               3
                                        1.0
                                               Luke S
            P. lotor
                               4
                                        1.0 Olivier R
         P. pygmaeus
                               5
                                        1.0 Olivier R
## 5
## 6 P. cancrivorus
                               6
                                        1.0
                                                Han S
                               7
                                               Luke S
## 7
            P. lotor
                                        1.0
## 8
        P. pygmaeus
                                        1.0
                                               Luke S
## 9 P. cancrivorus
                                        1.0
                                                Han S
```

MData <- read.csv("./InputData/A4_Caroline_Tang.csv")</pre>

```
H. sapiens
## 10
                            101
                                       1.0 Olivier R
## 11
        P. pygmaeus
                              1
                                       0.1
                                               Han S
        P. pygmaeus
                                              Luke S
## 12
                              2
                                       0.1
## 13
           P. lotor
                              3
                                       0.1
                                               Han S
                                       0.1 Olivier R
## 14
        P. pygmaeus
                              4
## 15 P. cancrivorus
                              5
                                              Luke S
                                       0.1
## 16
           P. lotor
                              6
                                       0.1
                                              Han S
                              7
                                       0.1
                                              Luke S
## 17
        P. pygmaeus
## 18 P. cancrivorus
                                       0.1
                                               Han S
                              8
## 19
           P. lotor
                              9
                                              Luke S
                                       0.1
```

Save new dataset as MeasShort.csv

write.csv(short, "./Output/MeasShort.csv")