Data importation

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1. endpoint

Comparisons between raw and cleaned data

ABER Data Analysis Timepoints

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Set the right working directory.

```
setwd("C:/Users/elise/Documents/Mémoire/Main/Data/Templates/ABER")
```

Data importation

Reimport the data sets extracted from the Data Preparation and Data Analysis R Markdown.

```
list.files()
```

```
## [1] "endpoint.txt" "plant_info.txt"
```

```
plant_info <- read.table("plant_info.txt", header = TRUE, sep = "\t")</pre>
endpoint <- read.table("endpoint.txt", header = TRUE, sep = "\t")</pre>
# plant info
plant info <- lapply(plant info, factor)</pre>
# endpoint
matching cols <- intersect(names(endpoint), names(plant info))</pre>
endpoint[, matching_cols] <- lapply(endpoint[, matching_cols], factor)</pre>
endpoint$Date <- date(endpoint$Date)</pre>
endpoint$Timestamp <- NA</pre>
platform <- "ABER"
# endpoint
df <- endpoint[,colSums(is.na(endpoint))<nrow(endpoint)]</pre>
genotype_index <- which(colnames(df) == "Genotype")</pre>
variables <- colnames(df[, c(3:(genotype_index - 1))]) # We remove the two first column
s that are "Unit.ID" and "Date"
print(paste(platform, ": The variables for endpoint are", paste(variables, collapse =
", "), sep = " "))
```

```
## [1] "ABER : The variables for endpoint are DW_shoot_g, FW_shoot_g, DW_root_g, FW_roo
t_g, Leaf_number"
```

```
endpoint$Plant_type <- substr(endpoint$Genotype, nchar(as.character(endpoint$Genotyp
e)), nchar(as.character(endpoint$Genotype)))</pre>
```

Get the cleaned endpoint data

```
endpoint_clean <- endpoint
# Run the function on the dataset for all the variables
endpoint_clean <- detect_replace_outliers_by_genotype(endpoint_clean)</pre>
```

Time point objects

Generation of the timePoints objects using the function "createTimePoints".

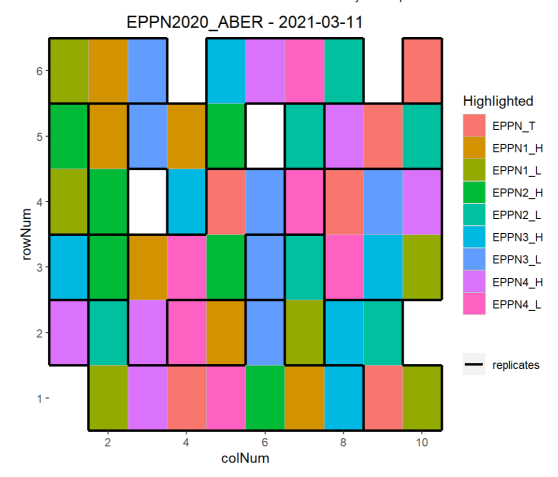
```
timePoint_endpoint <- createTimePoints(dat = endpoint,</pre>
                                        experimentName = "EPPN2020_ABER",
                                        genotype = "Genotype",
                                        timePoint = "Date",
                                        plotId = "Unit.ID",
                                        rowNum = "Row",
                                        colNum = "Column",
                                        repId = "Replication")
timePoint_endpoint_clean <- createTimePoints(dat = endpoint_clean,</pre>
                                        experimentName = "EPPN2020_ABER",
                                        genotype = "Genotype",
                                        timePoint = "Date",
                                        plotId = "Unit.ID",
                                        rowNum = "Row",
                                        colNum = "Column",
                                        repId = "Replication")
```

Gentoypic layout

Check the layout of the platforms' genotypes.

```
genotypes_list <- as.character(unique(endpoint$Genotype))

plot(timePoint_endpoint,
    plotType = "layout",
    highlight = genotypes_list,
    showGeno = FALSE)</pre>
```



1. endpoint

Comparisons between raw and cleaned data

View timePoint object.

```
summary(timePoint_endpoint)

## timePoint_endpoint contains data for experiment EPPN2020_ABER.
##
## It contains 1 time points.
## First time point: 2021-03-11
## Last time point: 2021-03-11
##
## No check genotypes are defined.

getTimePoints(timePoint_endpoint)

## timeNumber timePoint
## 1 1 2021-03-11
```

Count the number of observations per trait.

```
for (trait_name in traits) {
  print(paste("How many data observations for", trait_name))
  num_observations <- countValid(timePoint_endpoint, trait_name)
  print(num_observations)
}</pre>
```

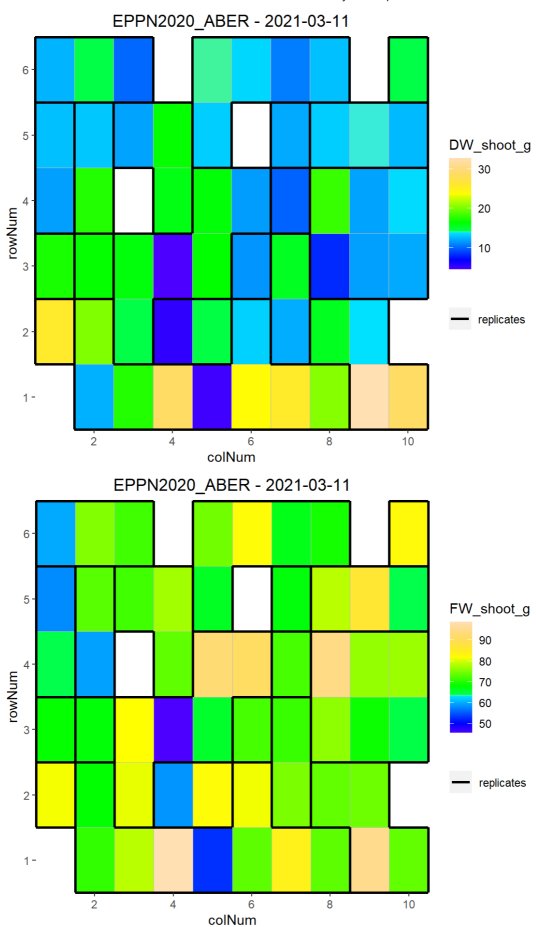
```
## [1] "How many data observations for DW_shoot_g"
## 2021-03-11
##
## [1] "How many data observations for FW_shoot_g"
## 2021-03-11
##
           54
## [1] "How many data observations for DW_root_g"
## 2021-03-11
##
           54
## [1] "How many data observations for FW_root_g"
## 2021-03-11
           54
## [1] "How many data observations for Leaf_number"
## 2021-03-11
##
           54
```

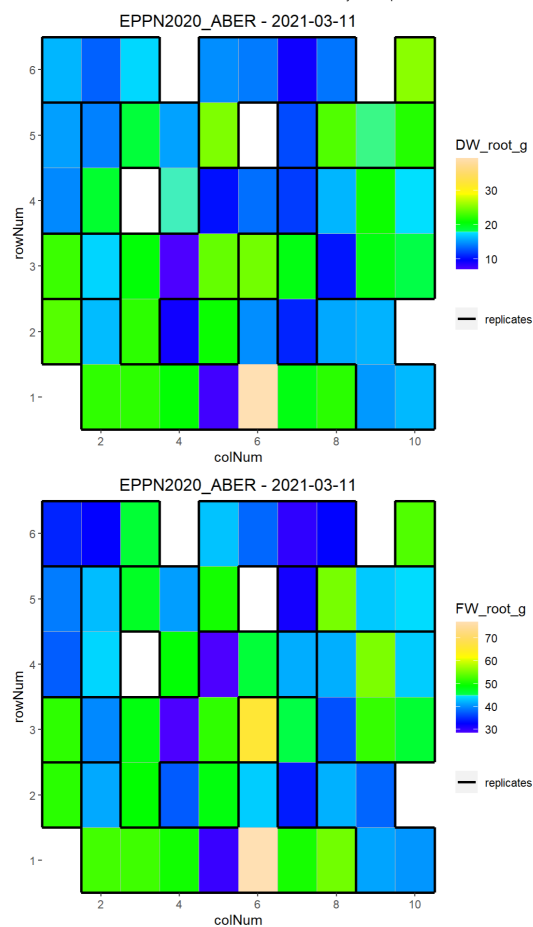
```
for (trait_name in traits) {
  print(paste("How many cleaned data observations for", trait_name))
  num_observations <- countValid(timePoint_endpoint_clean, trait_name)
  print(num_observations)
}</pre>
```

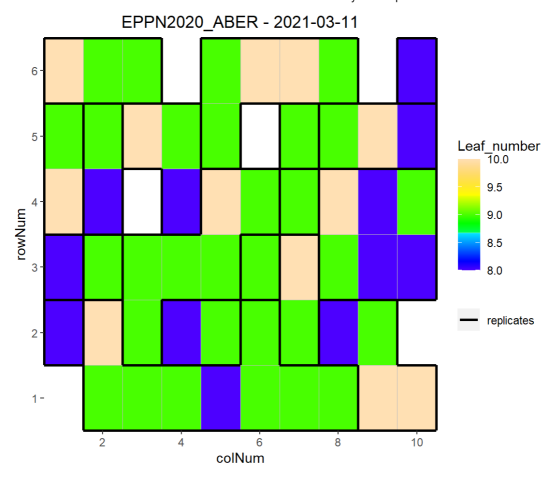
```
## [1] "How many cleaned data observations for DW_shoot_g"
## 2021-03-11
## 46
## [1] "How many cleaned data observations for FW_shoot_g"
## 2021-03-11
## 54
## [1] "How many cleaned data observations for DW_root_g"
## 2021-03-11
## 52
## [1] "How many cleaned data observations for FW_root_g"
## 2021-03-11
## 51
## [1] "How many cleaned data observations for Leaf_number"
## 2021-03-11
## 48
```

Check the heatmap of the data at harvest

```
for (trait_name in traits) {
  plot(timePoint_endpoint,
    plotType = "layout",
    timePoints = 1,
    traits = trait_name)
}
```







```
for (trait_name in traits) {
  plot(timePoint_endpoint_clean,
     plotType = "layout",
     timePoints = 1,
     traits = trait_name)
}
```

