# Most common statistical tests in R (Part-1)

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# All about multiple comparison of means from RAW data to publication ready graphs in R.

Install and library (load) the required packages

```
• Install the packages
```

## ##

## ## filter, lag

## The following objects are masked from 'package:base':

intersect, setdiff, setequal, union

```
install.packages("ggpubr")
install.packages("ggplot2")
install.packages("dplyr")
install.packages("ggpubfigs")
or intsall ggpubfigs like this:
install.packages("devtools")
library(devtools)
devtools::install_github("JLSteenwyk/ggpubfigs")
  • Load required packages
library(ggpubr)
## Loading required package: ggplot2
library(ggplot2)
library(dplyr)
##
## Attaching package: 'dplyr'
## The following objects are masked from 'package:stats':
```

#### library(ggpubfigs)

```
##
## Attaching package: 'ggpubfigs'
## The following object is masked from 'package:ggplot2':
##
## theme_grey
```

#### One way ANOVA-test:

#### Demo data set:

We will use PlantGrowth (a builtin R data set) for this example.

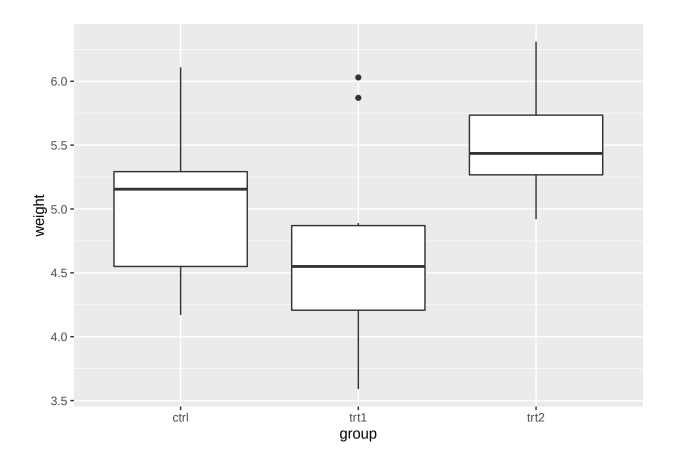
```
data("PlantGrowth")
head(PlantGrowth)
```

```
## weight group
## 1 4.17 ctrl
## 2 5.58 ctrl
## 3 5.18 ctrl
## 4 6.11 ctrl
## 5 4.50 ctrl
## 6 4.61 ctrl
```

```
View(PlantGrowth)
```

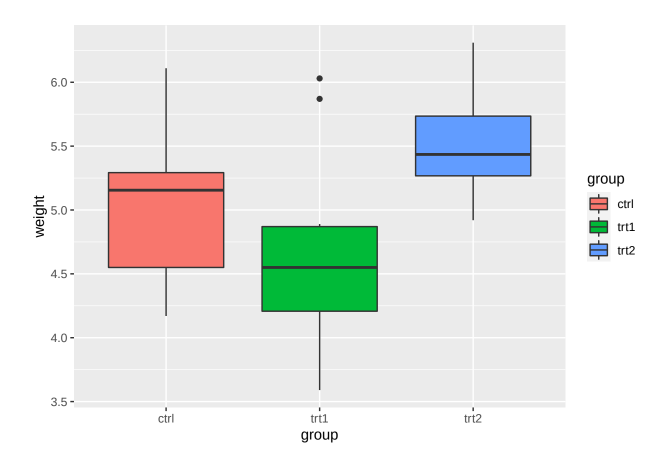
#### make a simple boxplot

```
ggplot(data = PlantGrowth, aes(x= group, y=weight))+
  geom_boxplot()
```



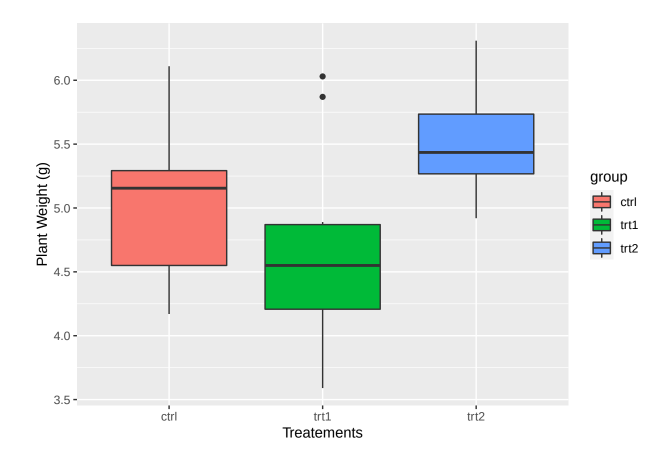
## add color (fill) argument to boxplot

```
ggplot(data = PlantGrowth, aes(x= group, y=weight, fill=group))+
  geom_boxplot()
```



## Change the axis labels

```
ggplot(data = PlantGrowth, aes(x= group, y=weight, fill=group))+
  geom_boxplot()+
  labs(x="Treatements", y="Plant Weight (g)")
```



#### one-way ANOVA test calculations

Two important R functions **aov()** and **summary.aov()** can be used to see the outcomes of one-way ANOVA model

```
# Analysis of variance calculation
model_aov <- aov(weight ~ group, data = PlantGrowth)
# Sumamry to view the output
summary(model_aov)

## Df Sum Sq Mean Sq F value Pr(>F)
## group 2 3.766 1.8832 4.846 0.0159 *
## Residuals 27 10.492 0.3886
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

Pr(>F) corresponding to the p-value of the test ### Interpretation of the one-way ANOVA results p-value less than 0.05 shows that model is significant, we can conclude that there are significant differences between the groups (treatments) highlighted with "\*" in the summary of the one-way ANOVA model.

#### Multiple pairwise comparison (TUKEY HSD test):

As the ANOVA test is significant, we can compute Tukey HSD (Tukey Honest Significant Differences, R function: TukeyHSD()) for performing multiple pairwise-comparison between the means of groups.

We can now compute the Tukey HSD test by using the following function of R:

```
TukeyHSD(model_aov)
```

```
## Tukey multiple comparisons of means
## 95% family-wise confidence level
##
## Fit: aov(formula = weight ~ group, data = PlantGrowth)
##
## $group
## diff lwr upr p adj
## trt1-ctrl -0.371 -1.0622161 0.3202161 0.3908711
## trt2-ctrl 0.494 -0.1972161 1.1852161 0.1979960
## trt2-trt1 0.865 0.1737839 1.5562161 0.0120064
```

- diff: difference between means of the two groups
- lwr,upr: the lower and the upper end point of the confidence interval at 95% (default)
- **p adj:** p-value after adjustment for the multiple comparisons. It can be seen from the output, that only the difference between trt2 and trt1 is significant with an adjusted p-value of 0.012, which is less than 0.05.

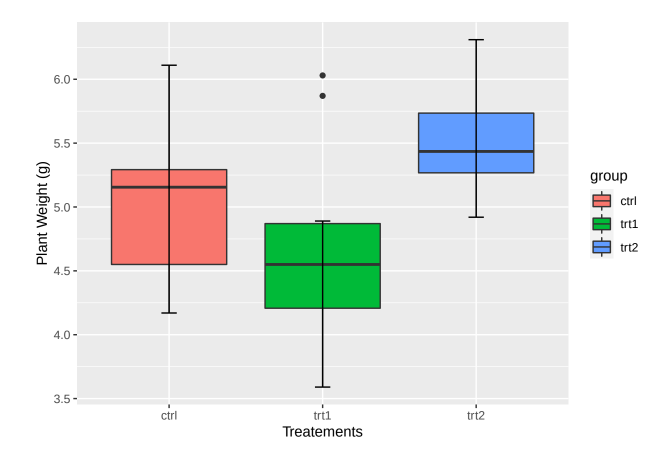
#### Visualize and specify the comparisons we want without computing tests:

Define the comparisons in an object

```
my_comparisons <- list( c("ctrl", "trt1"), c("ctrl", "trt2"), c("trt1", "trt2"))</pre>
```

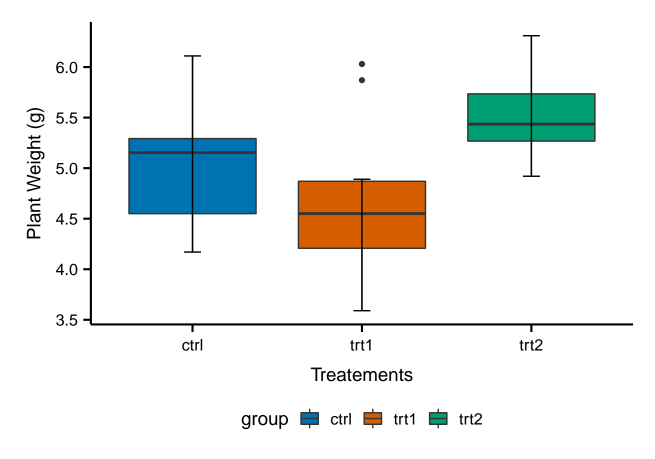
make a boxplot with errorbar caps

```
ggplot(data = PlantGrowth, aes(x= group, y=weight, fill=group))+
geom_boxplot()+
labs(x="Treatements", y="Plant Weight (g)")+
stat_boxplot(geom = 'errorbar', width=0.1)
```



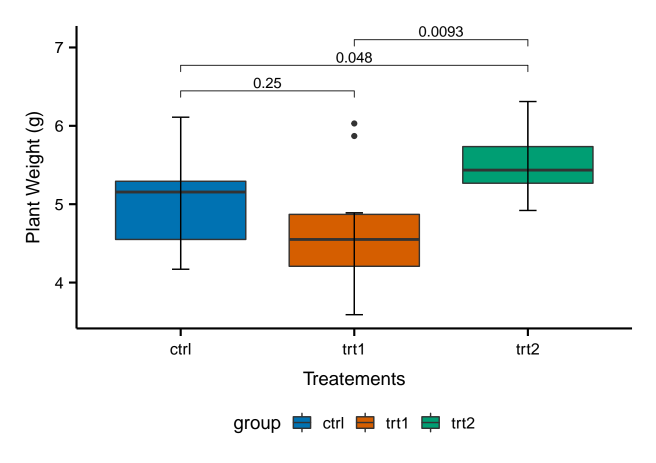
publication ready graphs can be made using ggpubfig package in the following way:

```
ggplot(data = PlantGrowth, aes(x= group, y=weight, fill=group))+
  geom_boxplot()+
  labs(x="Treatements", y="Plant Weight (g)")+
  stat_boxplot(geom = 'errorbar', width=0.1)+
scale_fill_manual(values = friendly_pal("ito_seven")) + theme_simple()
```



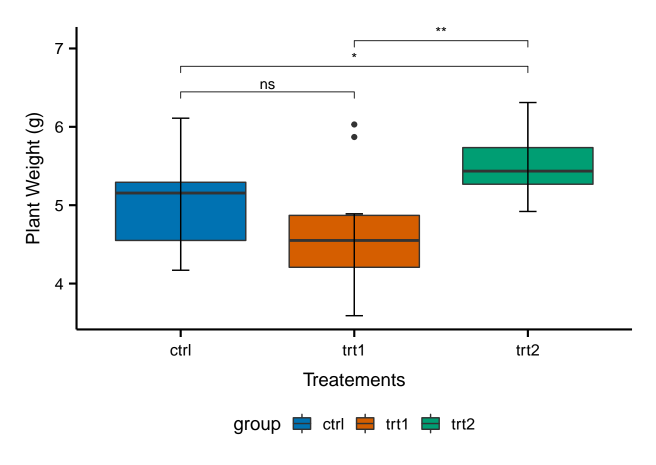
show significant differences of our comparisons on boxplot

```
ggplot(data = PlantGrowth, aes(x= group, y=weight, fill=group))+
  geom_boxplot()+
  labs(x="Treatements", y="Plant Weight (g)")+
  stat_boxplot(geom = 'errorbar', width=0.1)+
scale_fill_manual(values = friendly_pal("ito_seven")) + theme_simple()+
  stat_compare_means(method = "t.test", comparisons = my_comparisons)
```



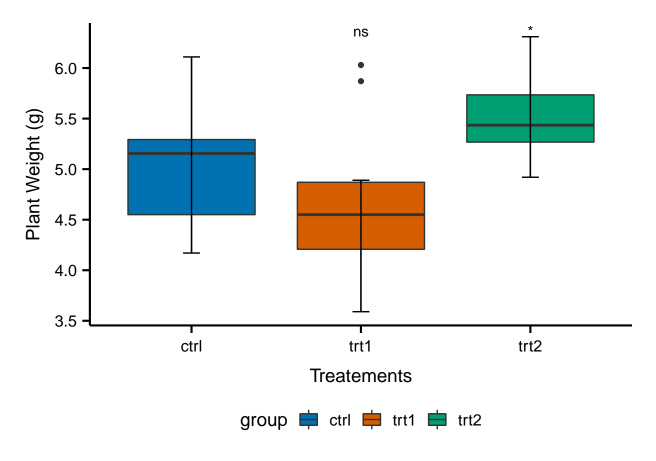
- show significant levels instead of using p-values

```
ggplot(data = PlantGrowth, aes(x= group, y=weight, fill=group))+
  geom_boxplot()+
  labs(x="Treatements", y="Plant Weight (g)")+
  stat_boxplot(geom = 'errorbar', width=0.1)+
  scale_fill_manual(values = friendly_pal("ito_seven")) + theme_simple()+
  stat_compare_means(method = "t.test", comparisons = my_comparisons, label = "p.signif") # Pairwise comparisons
```



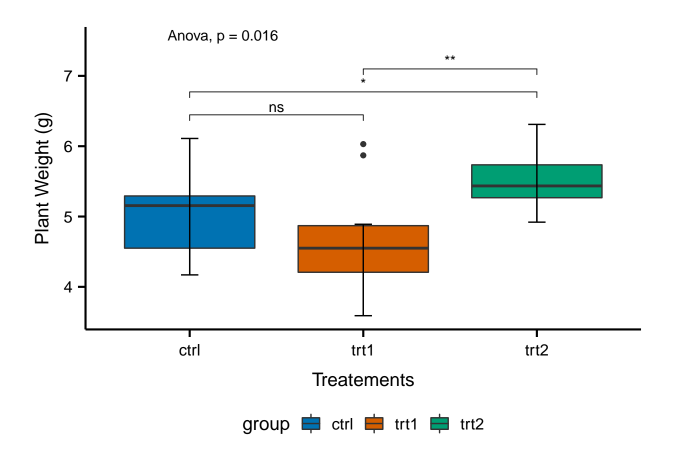
- Show significant differences based on a reference treatment, in this case our reference group treatment will be  $\operatorname{ctrl}$ 

```
ggplot(data = PlantGrowth, aes(x= group, y=weight, fill=group))+
  geom_boxplot()+
  labs(x="Treatements", y="Plant Weight (g)")+
  stat_boxplot(geom = 'errorbar', width=0.1)+
  scale_fill_manual(values = friendly_pal("ito_seven")) +theme_simple()+
  stat_compare_means(method = "t.test", ref.group = "ctrl", label = "p.signif")
```

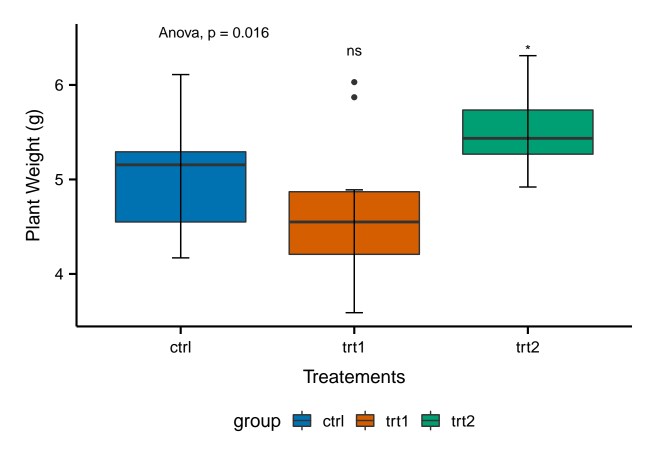


\*\*show ANOVA results

```
ggplot(data = PlantGrowth, aes(x= group, y=weight, fill=group))+
  geom_boxplot()+
  labs(x="Treatements", y="Plant Weight (g)")+
  stat_boxplot(geom = 'errorbar', width=0.1)+
  scale_fill_manual(values = friendly_pal("ito_seven")) +theme_simple()+
  stat_compare_means(method = "t.test", comparisons = my_comparisons, label = "p.signif")+ # Pairwise c
  stat_compare_means(method = "anova", label.y = 7.5)
```



```
ggplot(data = PlantGrowth, aes(x= group, y=weight, fill=group))+
geom_boxplot()+
labs(x="Treatements", y="Plant Weight (g)")+
stat_boxplot(geom = 'errorbar', width=0.1)+
scale_fill_manual(values = friendly_pal("ito_seven")) + theme_simple()+
stat_compare_means(method = "t.test", ref.group = "ctrl", label = "p.signif")+
stat_compare_means(method = "anova", label.y = 6.5)
```



For more content please subscribe our youtube channel Codanics Best regards, Dr Muhammad Aammar Tufail