# R Workshop: ANOVA, ANCOVA & Factorial ANOVA

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#### Introduction to ANOVA

# **Running ANOVA**

```
library(car)
library(psych)
library(multcomp)
library(effects)
library(tidyverse)
library(sjstats)
data <- dplyr::starwars</pre>
data <- data %>%
  dplyr::select(height,mass,hair_color,species,sex) %>%
  na.omit() %>%
  filter(species == "Gungan" | species == "Human" | species == "Wookiee")(1)
height_species_aov <- aov(height ~ species, data = data)
                                                                          2
summary(height_species_aov)
                                                                          (3)
# Post Hoc Test (Tukey)
TukeyHSD(height_species_aov)
                                                                          4
# Bonferroni Adjustment vs Tukey
pairwise.t.test(data$height, data$species, p.adjust.method = "bonferroni")(5)
```

- (1) Here we are using the filter() function to filter for "Gungan", "Human" and "Wookiee"
- (2) This is the aov() function. Here we are specifying the ANOVA formula.
- 3 The summary() function will give you the output of the ANOVA
- 4 Because the species variable has more than 2 conditions, we might want to figure out which comparisons are statistically different from each other. The TukeyHSD() function allows us to perform a post hoc Tukey test
- (5) Maybe Tukey isn't your favorite correction. Another route might be to use a pairwise t test that corrects for multiple comparisons using a Bonferroni correction. You can do that with the pairwise.t.test() function.

```
Df Sum Sq Mean Sq F value Pr(>F)
species 2 6118 3059 22.5 3.86e-06 ***
Residuals 23 3127 136
```

```
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
  Tukey multiple comparisons of means
    95% family-wise confidence level
Fit: aov(formula = height ~ species, data = data)
$species
                    diff
                                lwr
                                                  p adj
                                          upr
               -30.45455 -52.022229 -8.886862 0.0048319
Human-Gungan
Wookiee-Gungan 21.00000 -8.202782 50.202782 0.1915885
Wookiee-Human
                51.45455 29.886862 73.022229 0.0000125
    Pairwise comparisons using t tests with pooled SD
data: data$height and data$species
        Gungan Human
        0.0053 -
Human
Wookiee 0.2545 1.3e-05
P value adjustment method: bonferroni
```

# **Assumptions of ANOVA**

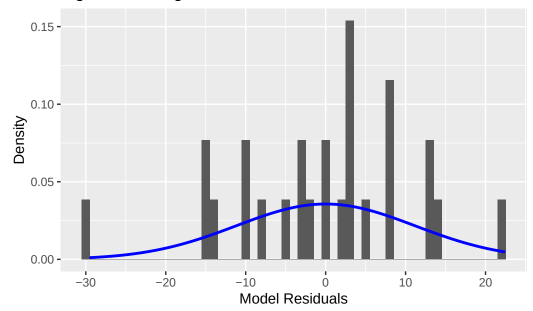
As ANOVA is just a special case of linear regression, the same assumptions exist for ANOVA that exist for linear regression. We will test each assumption below. The code will look awfully similar to the code used for linear regression previously

#### **Model Normality**

#### Graphical

```
x = "Model Residuals",
y = "Density")
density_plot
```

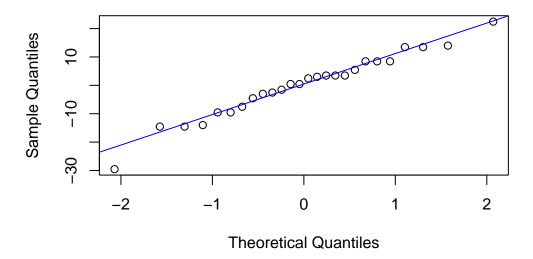
Figure 1. Histogram of Model Residual Scores



```
# OR

qqnorm(height_species_aov$residuals)
qqline(height_species_aov$residuals, col = "blue")
```

# Normal Q-Q Plot



1. When running the qqnorm() and qqline() functions at the same time, you will get a qq-plot which will graphically assess the normality of the argument given (in this case the model residuals). The col argument simply tells R which color to make the reference line.

#### **Statistical**

(1) Another way to assess normality is to look at the skew and kurtosis of the data. Typically skew and kurtosis between -1 and 1 are acceptable. We can see this using the describe() function within the psych package.

Shapiro-Wilk normality test

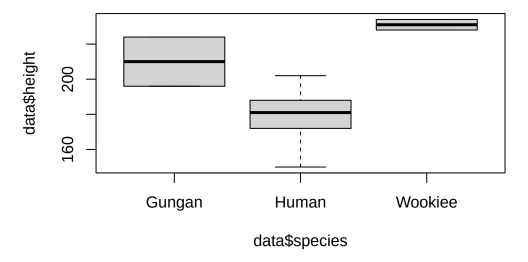
data: height\_species\_aov\$residuals
W = 0.97347, p-value = 0.7143

## Homogeneity of Variance

As with regression, homogeneity of variance is also important for ANOVA. We will graphically and statistically assess this assumption below.

## **Graphical**

(1) The boxplot() function will give us a visual representation of the DV at each level of the IV. We are looking for boxplots that are roughly the same height across each group.



#### **Statistical**

Statistically, we can assess homogeneity of variance using Levene's Test. Here we want the results to not be statistically significant.

1 The leveneTest() function from the car package takes a DV and IV as arguments.

```
Levene's Test for Homogeneity of Variance (center = median)

Df F value Pr(>F)
group 2 1.168 0.3288

23
```

#### Introduction to ANCOVA

# **Running ANCOVA**

ANCOVA refers to any ANOVA model with 2 or more parameters. These models require similar assumptions to ANOVA with a couple of additional ones. Below we will see how to run an ANCOVA and then test its assumptions.

```
data <- data %>% mutate(species = as.factor(species))
# Type I SS
ancova <- aov(height ~ species + mass, data = data)</pre>
summary(ancova)
                                                                            (1)
# Reversed Order
ancova2 <- aov(height ~ mass + species, data = data)</pre>
summary(ancova2)
                                                                            (2)
car::Anova(ancova2, type = "III")
                                                                            (3)
# Std Means
summary_data <- data %>%
  dplyr::group_by(species) %>%
  dplyr::summarise(n = n(),
                    mean = mean(height))
# Adjust means for covariate effect
adjustedMeans<- effect("species", ancova, se=TRUE)</pre>
                                                                            (4)
summary(adjustedMeans)
adjustedMeans$se
# Post Hoc Tests
posthoc <- multcomp::glht(ancova, linfct = multcomp::mcp(species = "Tukey"))(5)</pre>
summary(posthoc)
confint(posthoc)
# Effect size
sjstats::anova_stats(car::Anova(ancova,type = "III"))
                                                                            (6)
```

- 1 This displays our ANCOVA output with species and then mass.
- (2) This displays our ANCOVA output with mass and then species. We can see that 1 and 2 are different. This is because R by default does a sequential ANCOVA so order matters.

- ③ To get around this, we can use the Anova() function in the car package to specify that we want Type III sums of squares. This will then generate an ANCOVA where the order of predictors doesn't matter
- (4) When having covariates, one might wish to report adjusted means. We can do that using the effect() function from the effects package.
- (5) To run a post hoc analysis, we need to use the ghlt() function from the multcomp package. There are additional corrections you can do outside of Tukey. To investigate, please see the documentation.
- (6) To get basic ANOVA effect size statistics, we can use the anova\_stats() function from the sjstats package.

```
Df Sum Sq Mean Sq F value
                                         Pr(>F)
                 6118
                       3059.0
                                37.27 8.61e-08 ***
species
                 1322
                       1321.8
                                 16.10 0.000584 ***
mass
             1
Residuals
            22
                 1806
                         82.1
Signif. codes:
                0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
            Df Sum Sq Mean Sq F value
                                         Pr(>F)
             1
                 3737
                         3737
                                45.53 8.81e-07 ***
mass
             2
                 3702
                         1851
                                22.55 4.70e-06 ***
species
Residuals
            22
                 1806
                           82
                0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Signif. codes:
Anova Table (Type III tests)
Response: height
             Sum Sq Df F value
                                   Pr(>F)
(Intercept) 28097.6 1 342.337 6.734e-15 ***
mass
             1321.8 1 16.104 0.0005843 ***
                        22.555 4.699e-06 ***
species
             3702.4 2
Residuals
             1805.7 22
                0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Signif. codes:
 species effect
species
  Gungan
            Human Wookiee
214.4986 180.5408 215.5526
 Lower 95 Percent Confidence Limits
species
  Gungan
            Human Wookiee
```

#### 201.0113 176.5022 200.0533

Upper 95 Percent Confidence Limits species
Gungan Human Wookiee
227.9859 184.5794 231.0520
[1] 6.503427 1.947367 7.473624

Simultaneous Tests for General Linear Hypotheses

Multiple Comparisons of Means: Tukey Contrasts

Fit: aov(formula = height ~ species + mass, data = data)

Linear Hypotheses:

Estimate Std. Error t value Pr(>|t|) 6.748 -5.033 Human - Gungan == 0 -33.958 <0.001 \*\*\* Wookiee - Gungan == 0 1.054 10.333 0.102 0.994 Wookiee - Human == 0 35.012 7.846 4.462 <0.001 \*\*\* Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1 (Adjusted p values reported -- single-step method)

Simultaneous Confidence Intervals

Multiple Comparisons of Means: Tukey Contrasts

Fit: aov(formula = height ~ species + mass, data = data)

Quantile = 2.4824 95% family-wise confidence level

Linear Hypotheses:

Estimate lwr upr
Human - Gungan == 0 -33.9578 -50.7079 -17.2076
Wookiee - Gungan == 0 1.0541 -24.5973 26.7054
Wookiee - Human == 0 35.0118 15.5356 54.4880

term | sumsq | meansq | df | statistic | p.value | etasq | partial.etasq | omegasq

species	3702.445   1851.223   2   22.555   < .001   0.542	0.672   0.512
mass	1321.785   1321.785   1   16.104   0.001   0.194	0.423   0.179
Residuals	1805 669   82 076   22	1

# **Assumptions of ANCOVA**

ANCOVA requires the same assumptions as ANOVA with two additional assumptions: 1) That the predictor and covariate are independent 2) The regression slopes are homogeneous

We will test each of the traditional parametric tests as well as the two additional assumptions below.

#### **Predictor x Covariate Indepenence**

#### **Statistical**

```
predictor_assumption <- aov(mass ~ species, data = data)
summary(predictor_assumption)</pre>
```

(1) To assess this statistically, one needs to run an ANOVA looking at each predictor with each additional parameter. We do not want these to be statistically significant.

```
Df Sum Sq Mean Sq F value Pr(>F)
species 2 3390 1695.1 4.694 0.0195 *
Residuals 23 8306 361.1
---
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```



Here the statistical test for this assumption is statistically significant (which is bad). It actually means we can't do this (unless we have random assignment)

## Homogeneity of Regression Slopes

```
regression_slope_assumption <- aov(height ~ species*mass, data = data)
car::Anova(regression_slope_assumption, type = "III")
</pre>
```

1 To test this assumption, one just needs to test the interaction effects within the model. The code for this is shown here.

Anova Table (Type III tests)

```
Response: height
```

```
Sum Sq Df F value Pr(>F)

(Intercept) 149.72 1 1.9153 0.18163

species 155.01 2 0.9914 0.38856

mass 392.00 1 5.0144 0.03666 *

species:mass 242.18 2 1.5490 0.23689

Residuals 1563.48 20
---

Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

```
• Tip
```

For the above example, the interaction is NOT significant so assumption is met

# Residual Normality (DV ~ IV)

Residual normality is an assumption shared with standard regression. We can assess it the same way we did previously when looking at simple ANOVA. The same is true for the homogeneity of variance assumption.

```
height_mass_resid <- scale(residuals(aov(height ~ mass, data = data))) ①
psych::describe(height_mass_resid) ②
shapiro.test(height_mass_resid) ③</pre>
```

- (1) The scale() function will scale the residuals of the ANOVA. The residuals() function pulls out the residuals within the ANOVA
- 2 Statistical summary of the residuals using the describe() function in the psych package
- (3) Statistical test (Shapiro Wilks) of the residuals using the shapiro.test() function

```
vars n mean sd median trimmed mad min max range skew kurtosis se X1 1 26 0 1 0 -0.06 0.82 -1.85 2.7 4.55 0.71 0.44 0.2
```

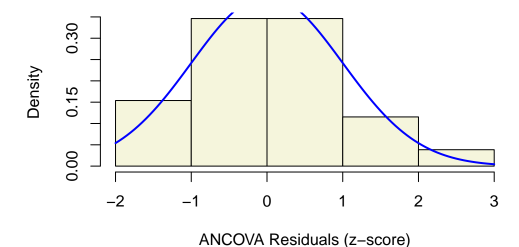
Shapiro-Wilk normality test

```
data: height_mass_resid
W = 0.95143, p-value = 0.2505
```

## **Graphical**

```
hist(height_mass_resid, col = 'beige',
    main="", xlab = "ANCOVA Residuals (z-score)",
    probability = TRUE)
curve(dnorm(x, mean = mean(height_mass_resid),
    sd = sd(height_mass_resid)),
    add = TRUE, lwd = 2, col = 'blue')
```

(1) A graphical histogram (with normal distribution overlay) of the model residuals



Homogeneity of Variance

```
# Levene's Test to Assess Equal Variance for Species
car::leveneTest(data$height ~ data$species)
①
```

(1) A statistical test (Levene's test) of the homogeneity assumption using the leveneTest() function in the car package.

```
Levene's Test for Homogeneity of Variance (center = median)

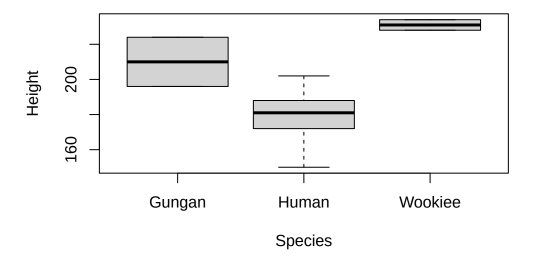
Df F value Pr(>F)
group 2 1.168 0.3288
23
```

## **Graphical**

```
# Boxplot Height by Species
boxplot(height ~ species,data=data, main="Height Variance by Species ",1)
xlab="Species", ylab="Height")
```

① A visual representation of the homogeneity of variance assumption using a boxplot

# **Height Variance by Species**



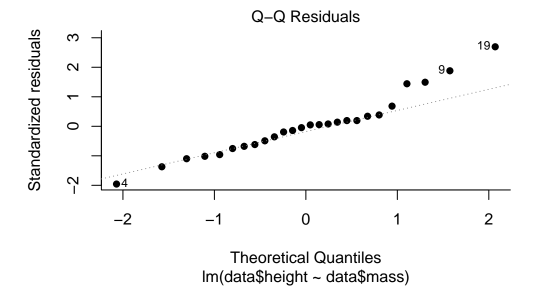
## Linearity of CV & DV

The last assumption is that the CV and DV are linearly related. We can test this using the code below graphically.

## **Graphical**

```
plot(lm(data$height ~ data$mass,data = data),
    pch = 16, bty = 'l',2)
```

(1) Visual representation of the CV to DV linearity assumption



#### Introduction to Factorial ANOVA

## **Running Factorial ANOVA**

```
library(tidyverse)
factorial_data <- starwars %>%
                                                                          1
  select(mass,homeworld,species) %>%
                                                                          (2)
  mutate(homeworld = as.factor(homeworld),
                                                                          (3)
         species = as.factor(species)) %>%
                                                                          4
  filter(homeworld == "Tatooine" | homeworld == "Naboo") %>%
                                                                          (5)
  filter(species == "Human" | species == "Droid") %>% na.omit()
                                                                          (6)
# Modeling As Factorial ANOVA
aov_factorial <- aov(mass ~ species*homeworld, data = factorial_data)</pre>
                                                                          (7)
Anova(aov_factorial, type = "III")
                                                                          (8)
# Modeling As A Regression (For SS Analyses)
reg_fanova <- lm(mass ~ homeworld*species,data = factorial_data)</pre>
                                                                          9
summary(reg_fanova)# <10>
```

- (1) Start with the starwars data set
- (2) Use the select() function to pull out mass, homeworld and specices variables
- (3) Call the mutate() function to format the homeworld variable using the as.factor() function.
- (4) Format the species variables as a factor using the as.factor() function.
- (5) Use the filter() function to filter for observations with homeworld = "Tatooine" OR "Naboo"
- (6) Use the filter() function to filter for observations with species = "Human" OR "Droid"
- 7 Create an ANOVA object using the aov() function
- (8) Use the Anova() function from the car() package to specify Type III Sums of Squares
- (9) Create an Regression object using the lm() function
- (1)0 Show output of the regression object using the summary() function.

#### Anova Table (Type III tests)

#### Response: mass

	Sum Sq	${\tt Df}$	F value	Pr(>F)
(Intercept)	1024.0	1	1.6199	0.2388
species	990.1	1	1.5662	0.2461
homeworld	308.2	1	0.4875	0.5048
species:homeworld	19.0	1	0.0301	0.8666
Rociduale	5057 2	2		

Residuals 5057.2 8

#### Call:

lm(formula = mass ~ homeworld \* species, data = factorial\_data)

#### Residuals:

```
Min 1Q Median 3Q Max -23.33 -19.50 -6.00 17.88 40.00
```

#### Coefficients:

	Estimate	Std. Error	t value	Pr(> t )
(Intercept)	32.000	25.143	1.273	0.239
${\tt homeworldTatooine}$	21.500	30.793	0.698	0.505
speciesHuman	36.333	29.032	1.251	0.246
homeworldTatooine:speciesHuman	6.167	35.557	0.173	0.867

Residual standard error: 25.14 on 8 degrees of freedom Multiple R-squared: 0.5219, Adjusted R-squared: 0.3426

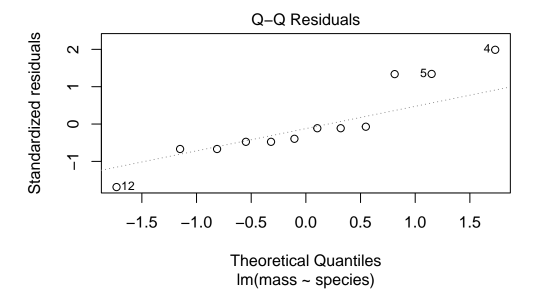
F-statistic: 2.91 on 3 and 8 DF, p-value: 0.1009

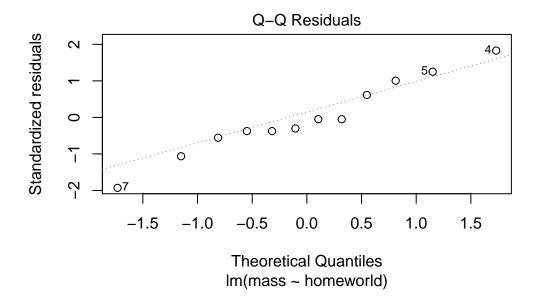
# **Assumptions of Factorial ANOVA**

# Linearity of IV to DV (Regression)

# **Graphical**

② Plot the residuals of the model using the plot() and lm() functions for mass on homeworld





#### Statistical

- ① Statistical summary of the model residuals using the describe() function in the psych package
- 2 Statistical test (Shapiro Wilk) of the residuals using the resid() and shapiro.test() functions

```
vars n mean sd median trimmed mad min max range skew kurtosis se X1 1 12 0 21.44 -6 -1.67 22.61 -23.33 40 63.33 0.43 -1.4 6.19
```

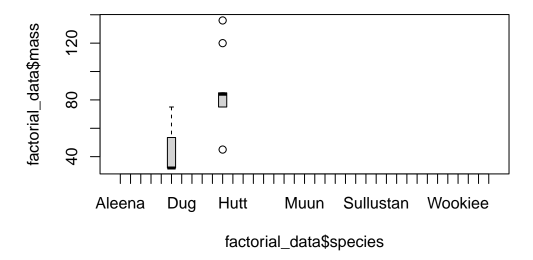
Shapiro-Wilk normality test

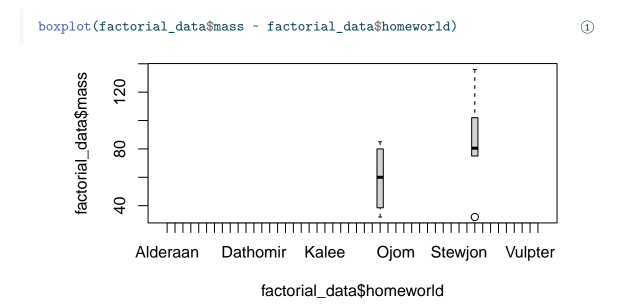
data: resid(aov\_factorial)
W = 0.90166, p-value = 0.1666

## Homogeneity of Variance (Regression)

#### **Graphical**

- (3) Fitted object of the ANCOVA
- (4) Plot of the residual x fitted data using the plot() function

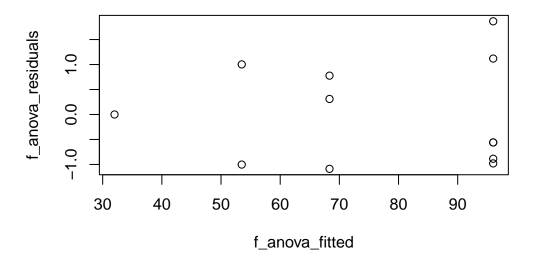




```
# Residuals
f_anova_residuals <- scale(residuals(lm(mass ~ species*homeworld, data = factorial_data)))
f_anova_fitted <- fitted(lm(mass ~ species*homeworld, data = factorial_data))

plot(f_anova_residuals ~ f_anova_fitted)

4</pre>
```



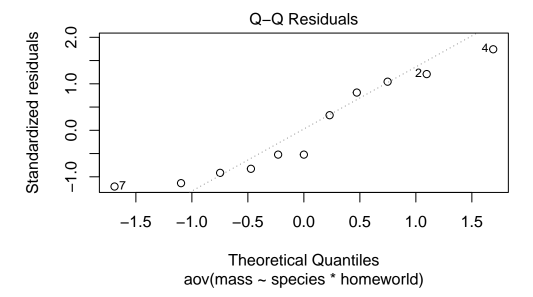
- 1 Levene test of mass x species using leveneTest() function
- 2 Levene test of mass x homeworld using leveneTest() function
- (3) Levene test of mass x species/homeworld interaction using leveneTest() function

## Normallity of Residuals

## Graphical

plot(aov\_factorial,2)

1 Plot ANOVA residuals using plot() function



#### **Statistical**

fanova\_residuals <- aov\_factorial\$residuals ①

shapiro.test(fanova\_residuals) ②

psych::describe(fanova\_residuals) ③

- (1) Create residual data set
- 2 Statistical test of residual normality using shapiro.test() function
- (3) Descriptive statistics of residuals using describe() function in the psych package

Shapiro-Wilk normality test

data: fanova\_residuals
W = 0.90166, p-value = 0.1666

vars n mean sd median trimmed mad min max range skew kurtosis se X1 1 12 0 21.44 -6 -1.67 22.61 -23.33 40 63.33 0.43 -1.4 6.19