Course 02402 Introduction to Statistics

Lecture 11: One-way Analysis of Variance, ANOVA

DTU Compute Technical University of Denmark 2800 Lyngby – Denmark

- Intro
- Model and hypothesis
- Computation decomposition and the ANOVA table
- Mypothesis test (F-test)
- Within-group variability and relation to the 2-sample t-test
- Post hoc analysis
- Model control / model validation
- A complete example from the book

Analysis of Variance

"ANalysis Of VAriance" (ANOVA) was introduced by R.A. Fisher 100 years ago as a systematic way of analysing groups and has since been a key part of statistics and its applications.

- Today: one factor (one-way ANOVA)
- Next week: two factors (two-way ANOVA)
- First factor is typically called *treatment*, second factor *block*.

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One-way ANOVA - simple example

Group A	Group B	Group C
2.8	5.5	5.8
3.6	6.3	8.3
3.4	6.1	6.9
2.3	5.7	6.1

Is there a difference (in means) between the groups A, B and C?

Analysis of variance (ANOVA) can be used for the analysis, if the observations in each group can be assumed to be normally distributed.

One-way ANOVA - simple example in R

```
# Input data
y \leftarrow c(2.8, 3.6, 3.4, 2.3,
       5.5. 6.3. 6.1. 5.7.
       5.8, 8.3, 6.9, 6.1)
## Define treatment groups
treatm <- factor(c(1, 1, 1, 1,
                   2. 2. 2. 2.
                   3, 3, 3, 3)
## Plot data by treatment groups
par(mfrow = c(1,2))
plot(y ~ as.numeric(treatm), xlab = "Treatment", ylab = "y")
boxplot(y ~ treatm, xlab = "Treatment", ylab = "y")
```

Eksempel: Cow dung and antibiotics

Decomposition of cow dung: how much organic material is left?

Control	lpha-cypermethrin	Ivermectin	Spiramycin
2.43	3.00	3.03	2.80
2.63	3.02	2.81	2.85
2.56	2.87	3.06	2.84
2.76	2.96	3.11	2.93
2.70	2.77	2.94	
2.54	2.75	3.06	

Cow dung and antibiotics – eksempel i R

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One-way ANOVA, model

• The model may be formulated as

$$Y_{ij} = \mu + \alpha_i + \varepsilon_{ij},$$

where the ε_{ij} are assumed to be independent and identically distributed (i.i.d.) with

$$\varepsilon_{ij} \sim N(0, \sigma^2)$$
.

- μ : overall mean.
- α_i : effect of group (treatment) i.
- Y_{ij} : jth measurement in group i (j runs from 1 to n_i).

One-way ANOVA, hypothesis

• We want to compare the (more than 2) means $\mu + \alpha_i$ in the model

$$Y_{ij} = \mu + \alpha_i + \varepsilon_{ij}, \quad \varepsilon_{ij} \stackrel{\text{i.i.d.}}{\sim} N(0, \sigma^2).$$

The hypothesis may be formulated as

$$H_0: \quad \alpha_i = 0 \quad \text{for all } i$$

with alternative hypothesis

$$H_1: \alpha_i \neq 0$$
 for at least one i

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One-way ANOVA, decomposition and the ANOVA table

With the model

$$Y_{ij} = \mu + \alpha_i + \varepsilon_{ij}, \quad \varepsilon_{ij} \stackrel{\text{i.i.d.}}{\sim} N(0, \sigma^2)$$

the total variation in the data can be decomposed:

$$SST = SS(Tr) + SSE$$
.

- 'One-way' refers to the fact that there is only one factor in the experiment on k levels.
- The method is called analysis of variance, because the testing is carried out by comparing certain variances.

Formulas for sums of squares

Total sum of squares ("the total variance")

$$SST = \sum_{i=1}^{k} \sum_{j=1}^{n_i} (y_{ij} - \bar{y})^2$$

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 The sum of squares for the residuals ("residual variance after model fit")

$$SSE = \sum_{i=1}^{k} \sum_{j=1}^{n_i} (y_{ij} - \bar{y}_i)^2$$

Sum of squares of treatment ("variance explained by the model")

$$SS(Tr) = \sum_{i=1}^{k} n_i (\bar{y}_i - \bar{y})^2$$

The ANOVA table

Source of	Deg. of	Sums of	Mean sum of
variation	freedom	squares	squares
Treatment	k-1	SS(Tr)	$MS(Tr) = \frac{SS(Tr)}{k-1}$
Residual	n-k	SSE	$MSE = \frac{SSE}{n-k}$
Total	n-1	SST	

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One-way ANOVA, F-test

• We have: (Theorem 8.2)

$$SST = SS(Tr) + SSE$$

and we can find the test statistic

$$F = \frac{SS(Tr)/(k-1)}{SSE/(n-k)} = \frac{MS(Tr)}{MSE}$$

where

- k is the number of levels of the factor.
- n is the total number of observations.
- Choose the significance level α , and compute the test statistic F.
- Compare the test statistic to the relevant quantile of the *F*-distribution:

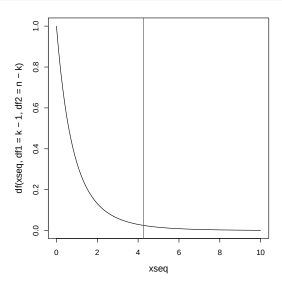
$$F \sim F_{\alpha}(k-1,n-k)$$
 (Theorem 8.6)

The F-distribution and the F-test

```
# Number of groups
k < -3
# Total number of observations
n < -12
# Sequence for plot
xseq \leftarrow seq(0, 10, by = 0.1)
# Plot density of the F-distribution
plot(xseq, df(xseq, df1 = k-1, df2 = n-k), type = "l")
# Plot critical value for significance level 5%
cr \leftarrow qf(0.95, df1 = k-1, df2 = n-k)
abline(v = cr. col = "red")
```

Remember, this is "under HO" (i.e. we compute as if HO is true)

An F-distribution with a critical value



The ANOVA table

Source of	Deg. of	Sums of	Mean sum of	Test-	<i>p</i> -
variation	freedom	squares	squares	statistic F	value
treatment	k-1	SS(Tr)		$F_{\rm obs} = \frac{MS(Tr)}{MSE}$	$P(F > F_{\text{obs}})$
Residual	n-k	SSE	$MSE = \frac{SSE}{n-k}$		
Total	n-1	SST			

One-way ANOVA F-test "by hand"

```
k \leftarrow 3; n \leftarrow 12 # Number of groups k, total number of observations n
# Total variation, SST
(SST \leftarrow sum((y - mean(y))^2))
# Residual variance after model fit, SSE
y1 \leftarrow y[1:4]; y2 \leftarrow y[5:8]; y3 \leftarrow y[9:12]
(SSE \leftarrow sum((v1 - mean(v1))^2) +
         sum((y2 - mean(y2))^2) +
         sum((y3 - mean(y3))^2))
# Variance explained by the model, SS(Tr)
(SSTr <- SST - SSE)
# Test statistic
(Fobs \leftarrow (SSTr/(k-1)) / (SSE/(n-k)))
# P-va.1.v.e.
(1 - pf(Fobs, df1 = k-1, df2 = n-k))
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Within-group variability and relation to the 2-sample t-test (Theorem 8.4)

The residual sum of squares, SSE, divided by n-k, also called residual mean square, MSE = SSE/(n-k), is the average within-group variability:

$$MSE = \frac{SSE}{n-k} = \frac{(n_1 - 1)s_1^2 + \dots + (n_k - 1)s_k^2}{n-k}$$
 (1)

$$s_i^2 = \frac{1}{n_i - 1} \sum_{i=1}^{n_i} (y_{ij} - \bar{y}_i)^2$$

ONLY when k = 2: (cf. Method 3.52)

$$MSE = s_p^2 = \frac{(n_1 - 1)s_1^2 + (n_2 - 1)s_2^2}{n - 2}$$

$$F_{\rm obs} = t_{\rm obs}^2$$

where $t_{\rm obs}$ is the pooled t-test statistic from Methods 3.52 and 3.53.

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Post hoc confidence interval - Method 8.9

 A single pre-planned confidence interval for the difference between treatment i and j is found as:

$$\bar{y}_i - \bar{y}_j \pm t_{1-\alpha/2} \sqrt{\frac{SSE}{n-k} \left(\frac{1}{n_i} + \frac{1}{n_j}\right)} \tag{2}$$

where $t_{1-\alpha/2}$ is based on the t-distribution with n-k degrees of freedom.

- Note the fewer degrees of freedom as more unknowns are estimated in the computation of $MSE=SSE/(n-k)=s_p^2$ (i.e. pooled variance estimate)
- If all M = k(k-1)/2 combinations of pairwise confidence intervals are found use the formula M times, but each time with $\alpha_{\mathsf{Bonferroni}} = \alpha/M$.

Post hoc pairwise hypothesis test- Method 8.10

• A single pre-planned level lpha hypothesis test:

$$H_0: \mu_i = \mu_j, H_1: \mu_i \neq \mu_j$$

is carried out as:

$$t_{\text{obs}} = \frac{\bar{y}_i - \bar{y}_j}{\sqrt{MSE\left(\frac{1}{n_i} + \frac{1}{n_j}\right)}} \tag{3}$$

and

$$p$$
 - value = $2P(t > |t_{obs}|)$

where the *t*-distribution with n-k degrees of freedom is used.

• If all M=k(k-1)/2 combinations of pairwise hypothesis tests are carried out use the approach M times, but each time with significance level $\alpha_{\mathsf{Bonferroni}}=\alpha/M$.

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Variance homogeneity

Look at a box plot to check whether the variability seems different across the groups.

```
# Check assumption of homogeneous variance using, e.g.,
# a box plot.
plot(treatm, y)
```

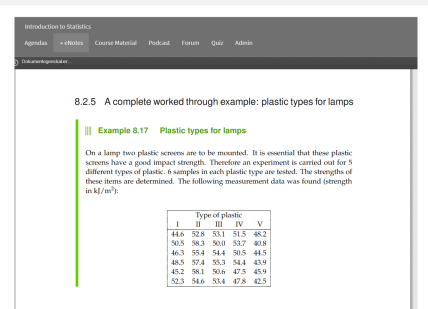
Normal assumption

Look at a normal QQ-plot of the residuals

```
# Check normality of residuals using a normal QQ-plot
fit1 <- lm(y ~ treatm)
qqnorm(fit1$residuals)
qqline(fit1$residuals)</pre>
```

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A complete example - from the book



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