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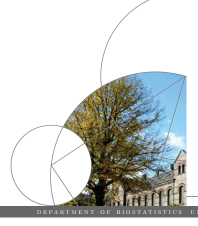


Outline

Day 4: Analysis of Variance

Paul Blanche

Section of Biostatistics, University of Copenhagen



Simple pairwise comparisons

ILO: to perform pairwise comparisons and draw rational conclusions

November 9, 2022

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Case: Irritable Bowel Syndrome Dose Response

- ▶ Data from n = 198 women.
- ► Randmomized (double-blind) to:
 - Placebo (n = 50, "dose 0")
 - Dose 1 (n = 54)
 - Dose 2 (n = 49)
 - Dose 3 (n = 45)

(Doses are blinded for confidentiality)

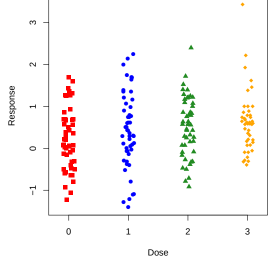


Outcome: baseline adjusted abdominal pain score at end of follow-up (12 weeks), approximately continuous variable, the larger the better.

Research questions:

- ► Does the drug work?
- ► Are there differences between doses?

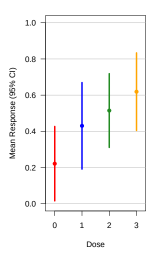
Outcome data



Dose	Mean	SD	n
0	0.22	0.72	50
1	0.43	0.88	54
2	0.51	0.71	49
3	0.62	0.71	45



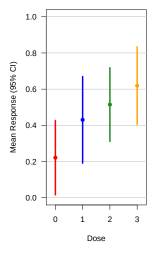
Pairwise Welch's t-test: results



p-values from pairwise t-tests:

Dose	0	1	2
1	0.19		
2	0.04	0.59	
3	0.01	0.24	0.48

Pairwise Welch's t-test: results



p-values from pairwise t-tests:

Dose	0	1	2
1	0.19		
2	0.04	0.59	
3	0.01	0.24	0.48

Have we not reported all relevant results? What is left to worry about?

Note: the y-axis has changed!



Note: the y-axis has changed!

Interpretations (1/3)

Results include:

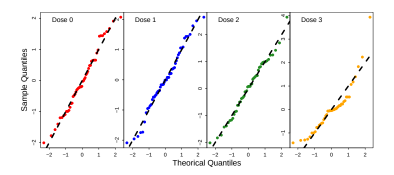
- ▶ Dose 0 not significantly different from dose 1.
- ▶ Dose 1 not significantly different from dose 2.
- ▶ But dose 0 significantly different from dose 1.

Are the results self-contradicting?

- ▶ No, this is just due to statistical uncertainty because of "small" sample sizes.
- ▶ Dose 1 may have a similar effect to either dose 0 or dose 2.

Interpretations (2/3)

What about the assumptions? Can we trust the results?



- ▶ QQplots for doses 0, 1, 2 look good but not so good for dose 3.
- ► However nothing "very" bad and decent sample size $(\geq 45 \text{ per group})$, so it seems fine.





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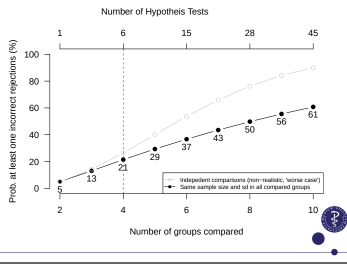
Interpretations (3/3)

Beware of the **multiple testing** issue! We have computed 6 p-values, hence the risk of making at least one false "discovery" is $\geq 5\%$.

Interpretations (3/3)

Beware of the **multiple testing** issue! We have computed 6 p-values, hence the risk of making at least one false "discovery" is > 5%.







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What statistical method should we use?

We want to control the FWER¹at 5%.

Using Bonferroni? No.

It's a conservative (i.e. sub-optimal) approach which ignores the (strong) correlation between the comparisons.

What statistical method should we use?

We want to control the FWER¹at 5%.

Using Bonferroni? No.

It's a conservative (i.e. sub-optimal) approach which ignores the (strong) correlation between the comparisons.

More modern alternative? Yes.

Use specific method and software for multiple correction that do not make any additional assumptions. The details of the method and computation are more complicated² but not the **interpretation** and **user-friendly software** exist.

¹Family-wise error rate (FWER): probability of making one or more false discoveries when performing multiple hypotheses tests (Lecture 2).

^{3'} ²That is why the method are still "new" and underused.

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¹Family-wise error rate (FWER): probability of making one or more false discoveries when performing multiple hypotheses tests (Lecture 2).

²That is why the method are still "new" and underused.

Recommended analysis (see R-demo for code)

Statistical methods:

Comparisons between groups were made with a heteroscedastic ANOVA model (not assuming equal variances). P-values and 95% confidence intervals were adjusted for multiple testing using the max-t test method (aka min-p method) as implemented in the multcomp-package [ref. 3] of the statistical software R [ref. 4] and described in [ref. 5].

Results (adjusted for multiple testing):

Comparison	Est. Diff	95% CI	p-value
1 - 0	0.209	[-0.202; 0.620]	0.552
2 - 0	0.293	[-0.083; 0.670]	0.185
3 - 0	0.398	[0.011; 0.784]	0.041
2 - 1	0.084	[-0.325; 0.494]	0.951
3 - 1	0.189	[-0.230; 0.607]	0.647
3 - 2	0.104	[-0.281; 0.489]	0.896

Note: p-values < 6 times the non-adjusted ones (Bonferroni).

What if we want only the comparisons to placebo?

Sometimes we only want all the comparisons to one (reference) group.

This is known as the **many-to-one** comparisons case (**Dunnett**), by contrast to the **all pairwise** comparisons case (**Tukey**).⁶

The method for this case is similar and we can use the same software.

Case results (adjusted for multiple comparisons to placebo):

Comparison	Est. Diff	95% CI	p-value
1 - 0	0.209	[-0.168; 0.586]	0.418
2 - 0	0.293	[-0.052; 0.639]	0.115
3 - 0	0.398	[0.043; 0.752]	0.023

Note: p-values \leq 3 times the non-adjusted ones (Bonferroni).



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6 Dunnett & Tukey were among the first statisticians who proposed specific, powerful, methods for these specific cas

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Pre-specification matters

The same comparison, e.g. Dose 3 versus Placebo (Dose 0), leads to the estimated mean difference 0.398, but different 95% confidence intervals (CI) and p-values (after adjusting for multiple testing) when we consider either:

- ► All-pairwise (6) comparisons: 95% Cl=[0.011; 0.784], p= 0.041.
- ightharpoonup Many-to-one (3) comparisons: 95% CI=[0.043; 0.752], p= 0.023.

Take home messages:

- ▶ The more comparisons the wider the 95% CI and the higher the p-values.
- ▶ Do not investigate more comparisons than interesting/possible (power ↓).
- ► The choice of investigating "all-pairwise" versus "many-to-one" comparisons should be done before seeing the data, i.e. pre-specified. Rigorously adjusting for multiple testing is not possible otherwise and how much we can trust the results without pre-specification is most unclear.

Digression: prespecified vs post hoc analyses

It is completely fine and often useful to performed post hoc⁷ analyses as long as:

- ▶ they are reported as such in publications⁸,
- conclusions based on them are not too strong.

"The main analyses should concentrate on the primary research questions to reduce the amount of testing of data-generated hypotheses. However, science would not proceed if analyses of questions not stated in the protocol were not allowed so, obviously, new ideas generated from the data can be pursued as long as conclusions based on such additional analyses are suitably calibrated." 9

Andersen & Skovgaard, *Regression with linear predictors* , page 473 (Springer, 2010)



³Hothorn, Bretz & Westfall (2008). Simultaneous Inference in General Parametric Models. Biometrical Journal 50(3), 346–363.

R Core Team (2022). R: A language and environment for statistical computing. R Foundation for Statistical Computing, Vien

^{10/57} Herberich, Sikorski & Hothorn. "A robust procedure for comparing multiple means under heteroscedasticity in unbalanced design

⁷A post hoc analysis is an analysis specified after the data were seen.

⁸Otherwise this is "data fishing", "data snooping" or "p-hacking" and this is considered as something in between "questionable research practice" and "scientific dishonesty and research misconduct"; see KU course "Responsible Conduct of Research".

Power and sample size calculation

When planning several comparisons, say K, with a FWER control at α , one can:

- 1. Define an adjusted type-I error $\alpha' = \alpha/K$.
- 2. Perform sample size and power calculation for each comparison as in the case of a unique comparison, using this adjusted type-I error α' as input of the formula instead of α .

Note: this is a "slightly" conservative approach 10.

 $\overline{^{_{14/57}}10}$ This calculation assumes that the Bonferroni correction will be used

Outline

Analysis of Variance (ANOVA): one-way

ILO: to describe the model, its parameters and assumptions

ILO: to explain why all assumptions are not all equally important

ILO: to interpret standard results



What is ANOVA about?

ANOVA stands for "ANalysis Of VAriance", but this is a method to compare means (via the comparisons of variances.).

Useful for answering research questions such as:

- ▶ Is this continuous outcome associated with this categorical variable?
- ▶ Is the mean outcome the same for all levels of this categorical outcome?

Examples:

- ▶ Outcome: weight, blood pressure, concentration, pain score ...
- ► Categorical variable: BMI group, age group, dose level ...

This is a very commonly used, well-known and "old" method.



ANOVA model (one-way)

The j-th observation from the i-th group is described as:

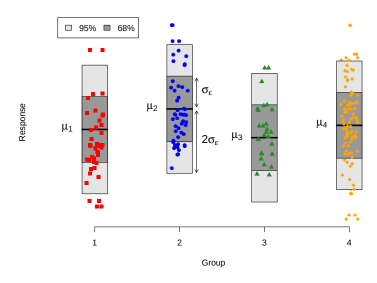
$$Y_{ij} = \mu_i + \varepsilon_{ij}$$

- \blacktriangleright μ_i is the (population) mean for group i.
- $ightharpoonup arepsilon_{ii}$'s are individual 'error' terms ("random/unexplained deviation from the mean") assumed normally distributed with zero mean and the same variance σ_{ε}^2 regardless of group.

Model assumptions (1-2 important, 3-4 not always):

- 1. Observations from different groups are independent.
- 2. Individual observations within each group are independent.
- 3. 'Error' terms are normally distributed.
- 4. The variance of 'error' terms is the same for all groups (homogeneit

Visual interpretation $(Y_{ij} = \mu_i + \varepsilon_{ij})$, hypothetical data)



 σ_{ε} tells us how **vertically spread** are the points above and below each group mean μ_i , for each group.

ANOVA (one-way): why and how?

Why using a (traditional) ANOVA?

To test the global null hypothesis " H_0 : The mean of all (K) groups are all equal", that is

$$H_0: \mu_1 = \mu_2 = \cdots = \mu_K.$$

How does it work?

By using a F-test which compares the between-group variability to the within-group variability. If the between-group variability is large enough relative to the within-group variability, then we reject H_0 .

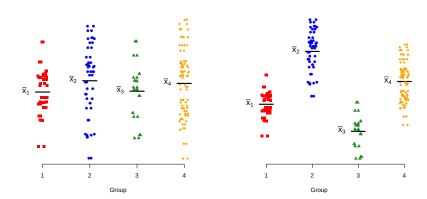
- ► Hence the name ANOVA: we analyze variances
- ► Computation possible by hand, hence the method became popular during the pre-computer age.

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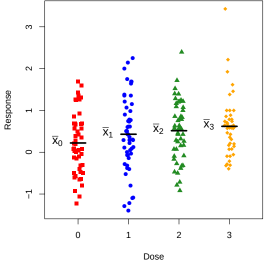
ANOVA: intuition of the F-test (hypothetical data)



- ▶ Left: the between-group variance (i.e. the variance of sample means \bar{x}_i) is small relative to the within-group variability: do not reject H_0 .
- ▶ right: the between-group variance is large relative to the within-group variability: reject H_0 .

Case: (traditional) ANOVA

- $\bar{x}_0 = 0.22$
- $\bar{x}_1 = 0.43$
- $\bar{x}_2 = 0.51$
- $\bar{x}_3 = 0.62$
- $\hat{\sigma}_{\varepsilon} = 0.76$
- ▶ p-value=0.07
- ▶ Do not reject!

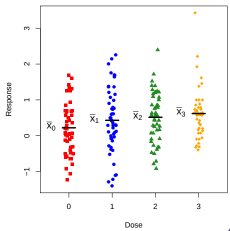


This is **not consistent** with the results from the all-pairwise comparisons...

Case: (traditional) ANOVA (without assuming homogeneity)

Let's use a more flexible model, which does not require assumption 4 (homogeneity).

- $\bar{x}_1 = 0.22, \, \hat{\sigma}_1 = 0.72$
- $\bar{x}_2 = 0.43, \ \hat{\sigma}_2 = 0.88$
- $\bar{x}_3 = 0.51, \ \hat{\sigma}_3 = 0.71$
- $\bar{x}_4 = 0.62$, $\hat{\sigma}_4 = 0.71$
- ▶ p-value=0.055
- ▶ Do not reject!



Still **not consistent** with the results from the all-pairwise comparisons. although the modeling assumptions are similar! (only the hypothesis test method is difficult.)

Power of F-test vs all-pairwise comparisons

We can test the global null hypothesis $H_0: \mu_1 = \mu_2 = \cdots = \mu_K$ using:

- ► F-test.
- ▶ min-p test (aka max-t test): perform all-pairwise comparisons and compute the p-value for H_0 as the minimum of the multiplicity adjusted p-values of all the comparisons (hence the "min-p" name). The rational is that we can safely reject H_0 if there is at least one significant difference after adjusting for multiple testing.

Which approach is the most powerful?

- ► F-test when the means of all groups are different although there is no particularly large difference between any two groups.
- min-p test: when there exists a particularly large difference between any two groups.

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Critics of the F-test and recommendations (1/2)

- When the F-test is significant we can conclude to differences between the groups but we do not know between which groups! (frustrating....). Historically, people used to proceed in two steps: 1) test the global null $H_0: \mu_1 = \mu_2 = \cdots = \mu_K$, 2) if H_0 is rejected, proceed to make pairwise comparisons, but why not directly start with pairwise comparisons, especially because...
- ► F-test and pairwise comparisons are inconsistent: either may find a significant difference the other doesn't. When it happens it is frustrating and hard to explain.
- ▶ When the F-test is not significant it is difficult to know whether it is due to lack of effect or lack of evidence, because there are no corresponding CIs.

Critics of the F-test and recommendations (2/2)

Recommendations:

- ▶ Unless you are not interested in the pairwise comparisons or have another specific reason in mind (e.g. power), prefer the all-pairwise comparisons and min-p (aka max-t) approach to the F-test.
- ▶ If you are not interested in the pairwise comparisons but **only** in knowing whether a continuous outcome is associated with a categorical, and if you want to keep the analysis as "simple and common" as possible, then you can prefer the F-test to the more "modern" min-p approach.



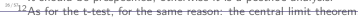


Checking the ANOVA model assumptions

But the F-test and ANOVA are still very much used... so, what should we know about checking their modeling assumptions?

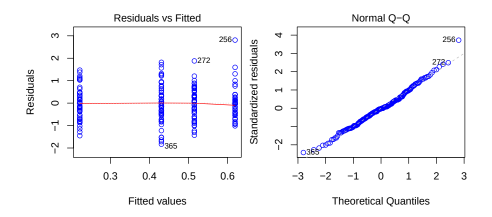
- ► Assumptions 1-2 (independence):
 - rely on the study design.
- ► Assumptions 3 (homogeneity of variances):
 - check with residual plots or compute sd in each group (best).
 - can be relaxed if needed (see R-demo for code).
 - ▶ log-transforming the data might help to obtain homogeneity of the variances. ¹¹
- ► Assumptions 4 (normality):
 - check with qqplot.
 - ▶ not needed with large sample sizes in each group. 12







Case: model checking (default) plots



Note: these are similar plots to those of the linear models. ¹³



^{27/57}13The 'default' left plot does not use "jitter"... Comparing SDs directly can be better

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ANOVA: usual software parametrization (1/4)

R code for ANOVA

fitlm <- lm(resp~dosefact, data=d)
summary(fitlm)</pre>

which returns (among other things)

Coefficients:

Estimate Std. Error t value Pr(>|t|)
(Intercept) 0.2213 0.1079 2.051 0.0416 *
dosefact1 0.2091 0.1498 1.396 0.1643
dosefact2 0.2935 0.1534 1.913 0.0572 .
dosefact3 0.3977 0.1568 2.537 0.0120 *
--Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Residual standard error: 0.7631 on 194 degrees of freedom Multiple R-squared: 0.03513,Adjusted R-squared: 0.02021 F-statistic: 2.354 on 3 and 194 DF, p-value: 0.07335

ANOVA: usual software parametrization (2/4)

- (Intercept): est. mean in the reference group: $\bar{x}_0 = 0.2213$.
- ▶ dosefact1: est. mean difference: $\bar{x}_1 \bar{x}_0 = 0.2091$.
- ▶ dosefact2: est. mean difference: $\bar{x}_2 \bar{x}_0 = 0.2935$.
- ▶ dosefact3: est. mean difference: $\bar{x}_3 \bar{x}_0 = 0.3977$.
- ► F-statistic: provides F-test p-value: 0.07335.
- ▶ Residual standard error: estimate of σ_{ε} : 0.7631





ANOVA: usual software parametrization (2/4)

- ▶ (Intercept): est. mean in the reference group: $\bar{x}_0 = 0.2213$.
- ▶ dosefact1: est. mean difference: $\bar{x}_1 \bar{x}_0 = 0.2091$.
- ▶ dosefact2: est. mean difference: $\bar{x}_2 \bar{x}_0 = 0.2935$.
- \blacktriangleright dosefact3: est. mean difference: $\bar{x}_3 \bar{x}_0 = 0.3977$.
- ► F-statistic: provides F-test p-value: 0.07335.
- ▶ Residual standard error: estimate of σ_{ε} : 0.7631
- ▶ p-values for the mean differences are not adjusted for multiple testing.

ANOVA: usual software parametrization (2/4)

- (Intercept): est. mean in the reference group: $\bar{x}_0 = 0.2213$.
- ▶ dosefact1: est. mean difference: $\bar{x}_1 \bar{x}_0 = 0.2091$.
- ▶ dosefact2: est. mean difference: $\bar{x}_2 \bar{x}_0 = 0.2935$.
- ightharpoonup dosefact3: est. mean difference: $\bar{x}_3 \bar{x}_0 = 0.3977$.
- ► F-statistic: provides F-test p-value: 0.07335.
- ▶ Residual standard error: estimate of σ_{ε} : 0.7631
- ▶ p-values for the mean differences are not adjusted for multiple testing.
- "default" summary presents only comparisons between the reference group and others (3 out of 6 possible). This is arbitrary!
- Note that if Dose 1 had been chosen as the reference group, among the 3 differences shown in the output none would be significant.



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ANOVA: usual software parametrization (3/4)

R code for ANOVA when the reference Dose is now Dose 1.

d\$dosefact <- relevel(d\$dosefact,ref="1")
fitlm <- lm(resp~dosefact, data=d)
summary(fitlm)</pre>

which returns (among other things)

Coefficients:

Estimate Std. Error t value Pr(>|t|) (Intercept) 0.4304 0.1038 4.145 5.08e-05 dosefact0 -0.2091 0.1498 0.164 dosefact2 0.0844 0.1505 0.561 0.576 1.225 dosefact3 0.1887 0.1540 0.222 Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' '1

Residual standard error: 0.7631 on 194 degrees of freedom Multiple R-squared: 0.03513, Adjusted R-squared: 0.02021 F-statistic: 2.354 on 3 and 194 DF, p-value: 0.07335

ANOVA: usual software parametrization (4/4)

The ANOVA model is actually a specific kind of linear model. ¹⁴ The mean of each group is described by the regression formula:

$$\mu_i = \alpha + \beta_1 \cdot I(\text{group}_i = \text{Dose 1}) + \beta_2 \cdot I(\text{group}_i = \text{Dose 2}) + \beta_3 \cdot I(\text{group}_i = \text{Dose 3})$$

where I() is the indicator function:

$$I(condition) = \begin{cases} 1, & \text{if condition is true} \\ 0, & \text{if condition is false} \end{cases}$$

Gro	ир	Dose 0	Dose 1	Dose 2	Dose 3
Mea	an	α	$\alpha+eta_1$	$lpha + eta_2$	$\alpha + eta_3$
Esti	mate	0.2213	0.2213 + 0.2091	0.2213 + 0.2935	0.2213 +0.3977

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^{31/57}14 more on linear model in Lecture 7; it explains the use of the 1m function in R

Outline

Simple pairwise comparisons

ILO: to perform pairwise comparisons and draw rational conclusions

Analysis of Variance (ANOVA): one-way

ILO: to describe the model, its parameters and assumptions

ILO: to explain why all assumptions are not all equally important

ILO: to interpret standard results

Analysis of Variance (ANOVA): two-way

ILO: to contrast one- and two-way ANOVA

ILO: to explain why all assumptions are not all equally important

ILO: to interpret standard results



Two-way ANOVA

What is it about?

Analysis the mean of a continuous outcome depending on two categorical variables.

Why and when is it useful?

- 1. to increase power and precision of the estimates.
- 2. to correct/adjust for differences between the groups that we primarily aim to compare (e.g. to adjust for baseline differences; to get closer to "causal" conclusions ¹⁵).
- 3. to (sometimes) better handle missing data.

Note: points 2 and 3 are closely related. ¹⁶

¹⁵More on Lecture 7.





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Case: weight gains in rats

- ▶ Data from n = 40 rats, fed on four diets.
- ► Two amounts of protein (low and high)
- ► Two sources of protein (beef and cereal)
- ► Randomized, Factorial, Balanced experiment.

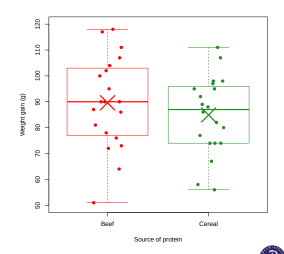


Outcome: weight gain in grams.

Research question: Does one of the two sources of proteins lead to larger weight gains (in average)?

Simple comparison: t-test

- $\bar{x}_1 = 89.6, \, \hat{\sigma}_1 = 17.7$
- $\bar{x}_2 = 84.9$, $\hat{\sigma}_2 = 15.0$
- ▶ p-value of t-test=0.37
- ► Difference in mean 4.70, 95% CI = [-5.81;15.21]



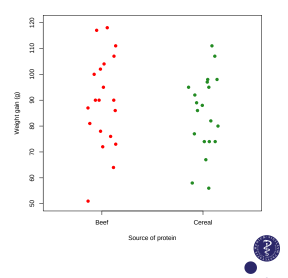


Note: crosses show the means.

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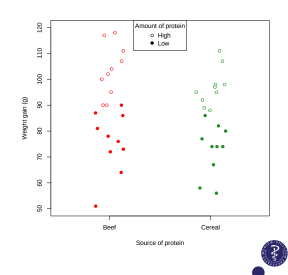
The data

Adjusting increases power & precision



Theory (maths) shows that adjusting on a variable (strongly) associated with the outcome will generally increase the power of the analysis and the precision of the estimates (i.e. smaller s.e., narrower CI).

To get the intuition, let's imagine this hypothetical situation...



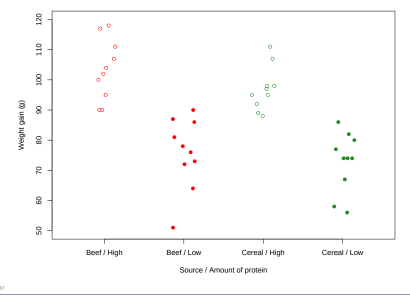
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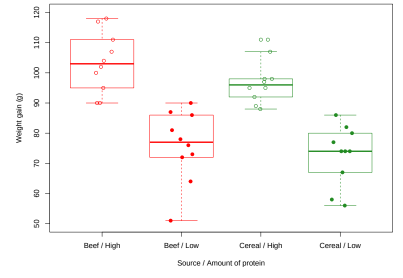
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Unfolding the four groups (hypothetical) (1/3)

Unfolding the four groups (hypothetical) (2/3)

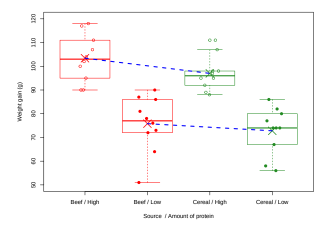




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Unfolding the four groups (hypothetical) (3/3)



- ► Some evidence in each subgroup (High and Low)
- The evidence of each subgroup is based on less subjects ($\searrow n \Rightarrow \searrow$ evidence/power) but based on data with less variability ($\searrow \sigma \Rightarrow \nearrow$ evidence/power), which somehow "balances out".
- So, overall there is more evidence (i.e. smaller s.e. and more power)

The two-way ANOVA model (without interaction)

The k-th observation from the (i, j)-th combination group (e.g. source of protein i and amount of protein j) is described as:

$$Y_{ijk} = \mu_{ij} + \varepsilon_{ijk}$$

= $\gamma_i + \eta_j + \varepsilon_{ijk}$ (assuming no interaction).

- \blacktriangleright $\mu_{ij} = \gamma_i + \eta_j$ is the mean for the (i, j)-th combination group.
- $ightharpoonup arepsilon_{ijk}$'s are individual 'error' terms ("random/unexplained deviation from the mean") assumed normally distributed with zero mean and the same variance σ_{ε}^2 regardless of group.



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Two-way ANOVA assumptions (without interaction)

Model assumptions (1-4 similar to that of the one-way ANOVA):

- 1. Observations from different groups are independent.
- 2. Individual observations within each group are independent.
- 3. 'Error' terms are normally distributed.
- 4. The variance of 'error' terms is the same for all groups (homogeneity).
- 5. There is no interaction (\rightarrow) .

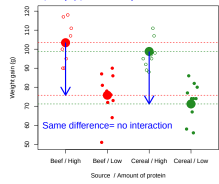
Note: 1-2 and 5 are important, 3-4 not always (as for one-way ANOVA).

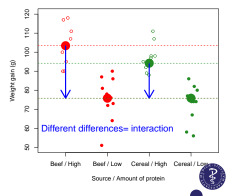
The meaning of "no interaction" (1/2)

No interaction models $\mu_{ij}=\gamma_i+\eta_j$ for the mean for the (i,j)-th combination group. In our example that means that:

The $\left\{\begin{array}{c} \text{source} \\ \text{amount} \end{array}\right\}$ of protein "shifts" the mean of all $\left\{\begin{array}{c} \text{amounts} \\ \text{sources} \end{array}\right\}$ (up or down) in the same way.

Example (hypothetical):



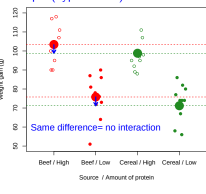


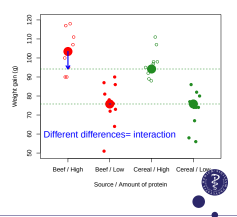
The meaning of "no interaction" (2/2)

No interaction models $\mu_{ij}=\gamma_i+\eta_j$ for the mean for the (i,j)-th combination group. In our example that means that:

The $\left\{\begin{array}{c} \text{source} \\ \text{amount} \end{array}\right\}$ of protein "shifts" the mean of all $\left\{\begin{array}{c} \text{amounts} \\ \text{sources} \end{array}\right\}$ (up or down) in the same way.

Example (hypothetical):





Two-way ANOVA: usual software parametrization (1/4)

Now we come back to the real data and run the analysis!

One-way ANOVA (for comparison)

OneWayRes <- lm(weightgain~source,data=weightgain)
summary(OneWayRes)</pre>

which returns

Coefficients:

```
Estimate Std. Error t value Pr(>|t|)
(Intercept) 89.600 3.669 24.419 <2e-16 ***
sourceCereal -4.700 5.189 -0.906 0.371
---
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Residual standard error: 16.41 on 38 degrees of freedom
Multiple R-squared: 0.02113, Adjusted R-squared: -0.004628
F-statistic: 0.8203 on 1 and 38 DF, p-value: 0.3708
```



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Two-way ANOVA: usual software parametrization (2/4)

Two-way ANOVA

TwoWayRes <- lm(weightgain~type+source,data=weightgain)
summary(TwoWayRes)</pre>

which returns

Coefficients:

	Estimate Std.	Error t value	Pr(> t)	
(Intercept)	95.300	4.255 22.396	3 <2e-16	***
typeLow	-11.400	4.914 -2.320	0.026	*
sourceCereal	-4.700	4.914 -0.957	0.345	
Signif. code	s: 0 '***, 0.	001 '**' 0.01	<pre>'*' 0.05 '</pre>	., 0.1 , 1
Residual star	ndard error: 1	5.54 on 37 des	rees of fr	eedom

Multiple R-squared: 0.1455,Adjusted R-squared: 0.09926 F-statistic: 3.149 on 2 and 37 DF, p-value: 0.05459

Note:

- Variable type in the data weightgain indicates the amount of protein.
- **F-test p-value is not so interesting here** (H_0 : "neither effect of amount of protein nor of source or protein").
- ► Same estimated difference (-4.70) as in the one-way ANOVA because of the balanced factorial design.

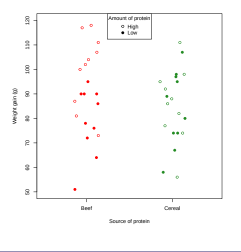
Two-way ANOVA: usual software parametrization (3/4)

- ► (Intercept): est. mean in the reference group (High, Beef).
- ▶ typeLow: est. mean difference between amount (type) of protein, Low vs High, for the same source of protein (any).
- sourceCereal: est. mean difference between sources of protein, Cereal vs Beef, for the same amount (type) of protein (any).
- ► F-statistic: not so interesting (see previous slide).
- ▶ Residual standard error: estimate of σ_{ε} : 15.54



Digression

In this example, the s.e. for the difference of interest is unfortunately only "a little bit" smaller with the two-way ANOVA than with the one-way ANOVA (4.914 vs 5.189). This is because the vertical spread of the observations of the weight gains (i.e. the standard deviation) is not largely different when looking at the entire data or within subgroups.



Two-way ANOVA: usual software parametrization (4/4)

This ANOVA model is also a specific kind of linear model. The mean of each group is given by this regression formula:

$$\mu_{ij} = \alpha + \beta_1 \cdot I(\text{amount}_i = \text{Low}) + \beta_2 \cdot I(\text{source}_i = \text{Cereal})$$

Modeled means and estimates:

Amount	High	Low
Beef	α	$\alpha + \beta_1$
	95.3	95.3 + (-11.4) = 83.9
Cereal	$\alpha + eta_2$	$\alpha + \beta_1 + \beta_2$
	95.3 + (-4.7) = 90.6	95.3 + (-4.7) + (-11.4) = 79.2



(plot of the real data)

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What if we compare more than two groups?

- ▶ In short, everything is very similar to what we have seen before.
- ► For illustration, let's artificially create additional data of 20 more observations from rats fed with Fish (again, 10 receive a Low amount of protein, 10 a High amount).

Two-way ANOVA (results with additional, artificial, data)

Two-way ANOVA

TwoWayRes <- lm(weightgain~type+source,data=weightgain)
summary(TwoWayRes)</pre>

which returns

Coefficients:

```
Estimate Std. Error t value Pr(>|t|)
(Intercept)
              95.250
typeLow
             -11.300
                                 -2.984
                                          0.0042 **
sourceCereal
             -4.700
                          4.637
                                -1.014
                                          0.3152
sourceFish
             -10.050
                                         0.0345 *
                          4.637 -2.167
Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
Residual standard error: 14.66 on 56 degrees of freedom
Multiple R-squared: 0.1955, Adjusted R-squared: 0.1524
```

F-statistic: 4.537 on 3 and 56 DF, p-value: 0.006454





F-test with two-way ANOVA

An F-test can be performed for the null hypothesis $H_0: \beta_2 = \beta_3 = 0$, with model formula

$$\mu_{ij} = \alpha + \beta_1 \cdot I(\mathsf{amount}_j = \mathsf{Low}) + \beta_2 \cdot I(\mathsf{source}_i = \mathsf{Cereal}) + \beta_3 \cdot I(\mathsf{source}_i = \mathsf{Fish})$$

that is

 H_0 : "All sources of proteins give the same average weight gain, when comparing rats fed with the same amount of protein".

It compares the mean weight gain from all sources of proteins "adjusted" on the amount of protein received (i.e. within groups of rats receiving the same amount of proteins).

- ▶ This is a very commonly used, well-known and "old" method.
- Pros and cons: similar to that of F-test for one-way ANOVA.



R code and results

R code for the F-test (two-way ANOVA)

```
Full.lm <- lm(weightgain~type+source, data=weightgain) # "full" model (same as TwoWayRes)
Cons.lm <- lm(weightgain~type, data=weightgain) # "constrained" model
anova(Cons.lm,Full.lm) # F-test (compares the 2 models)</pre>
```

which returns

```
Analysis of Variance Table

Model 1: weightgain ~ type

Model 2: weightgain ~ type + source

Res.Df RSS Df Sum of Sq F Pr(>F)

1 58 13054

2 56 12042 2 1011.4 2.3517 0.1045
```



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R code and results

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```
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Cons.lm <- lm(weightgain~type, data=weightgain)  # "constrained" model
anova(Cons.lm,Full.lm)  # F-test (compares the 2 models)
```

which returns

```
Analysis of Variance Table

Model 1: weightgain ~ type

Model 2: weightgain ~ type + source

Res.Df RSS Df Sum of Sq F Pr(>F)

1 58 13054

2 56 12042 2 1011.4 2.3517 0.1045
```

Comments:

- ► F-test p-value=0.1045 is not significant.
- ► To avoid coding mistakes and misunderstandings of the R output, compare the two models and do not instead use "anova(Full.lm)", since the order of the variables in the formula would generally matter in that case.

Recommended analysis (see R-demo for code)

Statistical methods:

Comparisons between sources of proteins were made using a two-way ANOVA model (without interaction), to adjust for the amount of proteins received. P-values and 95% confidence intervals were adjusted for multiple testing using the max-t test method (aka min-p method) as implemented in the multcomp-package [ref. 17] of R [ref. 18] and described in [ref. 19].

Results (adjusted for multiple testing):

Comparison	Est. Diff	95% CI	p-value
Cereal - Beef	-4.7	[-15.9; 6.5]	0.572
Fish - Beef	-10.1	[-21.2; 1.1]	0.086
Fish - Cereal	-5.4	[-16.5; 5.8]	0.486

Notes:

- ▶ p-values ≤ 3 times the non-adjusted ones obtained from the default summary (Bonferroni).
- results for comparisons, including "Fish Cereal" (unlike in default summary)

Bretz, Hothorn, & Westfall (2016). Multiple comparisons using R. CRC Pres



¹⁷ Hothorn, Bretz & Westfall (2008). Simultaneous Inference in General Parametric Models. Biometrical Journal 50(3), 346–363.

¹⁸ R Core Team (2022). R: A language and environment for statistical computing. R Foundation for Statistical Computing, Via Austria. URL https://www.R-project.org/.

Assuming no interaction

- ► This assumption can be important.
- ▶ It simplifies the interpretation of the results.
- ▶ It should be supported by subject-matter knowledge.
- ▶ This assumption can (most often) be checked with the data.
- ▶ Usually, the smaller the sample size the more assumptions we need to compensate. This applies to the assumption of no interaction.
- ▶ More on interactions in Lecture 7.



..., ...

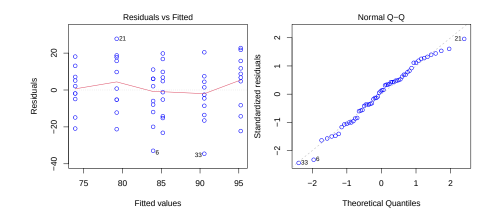
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(A few) Take home messages

- ▶ ANOVA is useful to compare the mean outcome in several groups.
- ▶ One-way: one categorical variable used in the analysis, Two-way: two categorical variables.
- A two-way ANOVA can be better than a one-way ANOVA (gain in power and precision), but not necessarily (more assumptions are required).
- ▶ Pairwise comparisons are often relevant, but adjusting for multiple testing is necessary to control the risk of false discoveries.
- F-tests are often used, sometimes useful, but not always.
- ► All model assumptions are not equally important.
- ► Think about your research question first, then choose your statistical method (one- or two-way ANOVA, F-test, pairwise comparisons, which multiple testing adjustment: none, all-pairwise, many-to-one?)
- ► Check the main modeling assumptions, tune your conclusions appropriately.



Model checking (default) plots



Note: these are similar plots to those of the linear models.



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