

Lecture 4: Multiple linear regression BIO144 Data Analysis in Biology

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Overview



- ▶ Checking the assumptions of linear regression: the QQ-plot
- ► Multiple linear regression x_1, x_2, \ldots, x_m
- $ightharpoonup R^2$ in multiple linear regression
- t-tests, F-tests and p-values
- Binary and categorical explanatory variables

Course material covered today



The lecture material of today is based on the following literature:

- ► Chapters 3.1, 3.2a-q of *Lineare Regression*
- ► Chapters 4.1 4.2f, 4.3a-e of *Lineare Regression*

Recap of last week I

▶ The linear regression model for the data $y = (y_1, ..., y_n)$ given $x = (x_1, ..., x_n)$ is

$$y_i = \alpha + \beta x_i + \epsilon_i$$
, $\epsilon_i \sim N(0, \sigma^2)$ independent.

- **E**stimate the parameters α , β and σ^2 by least squares.
- ▶ The estimated parameters $\hat{\alpha}$, $\hat{\beta}$ contain uncertainty and are normally distributed around the true values.
- Use the knowledge about the distribution to formulate statistical tests, such as: Is $\beta = 0$?
 - \rightarrow **T-test** with n-2 degrees of freedom.
- ► All this is done automatically by R:

summary(r.bodyfat)\$coef

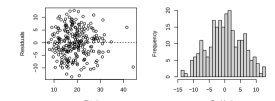
```
## Estimate Std. Error t value Pr(>|t|)
## (Intercept) -26.984368 2.7689004 -9.745518 3.921511e-19
## bmi 1.818778 0.1083411 16.787502 2.063854e-42
```

Recap of last week II

Remember: The assumption in linear regression is that the residuals follow a $N(0, \sigma^2)$ distribution, implying that :

- a) The expected value of ϵ_i is 0: $E(\epsilon_i) = 0$.
- b) All ϵ_i have the same variance: $Var(\epsilon_i) = \sigma^2$.
- c) The ϵ_i are normally distributed.
- d) The ϵ_i are independent of each other.

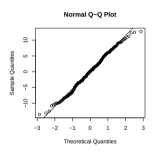
We started to do some residual analysis using the Tukey-Anscombe plot and the histogram of the residuals R_i .





Another useful diagnostic plot: The QQ-plot

Usually, not the histogram of the residuals is plotted, but the so-called quantile-quantile (QQ) plot. The quantiles of the observed distribution are plotted against the quantiles of the respective theoretical (normal) distribution:



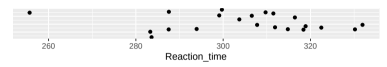
If the points lie approximately on a straight line, the data is fairly normally distributed.

This is often "tested" by eye, and needs some experience.

What on earth is a quantile???



Imagine we make 21 measures of something, say 21 reaction times:



The median of these is 307.8. The median is the 50% or 0.5 quantile, because half the data points are above it, and half below.

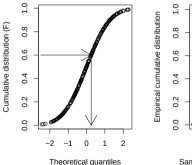
```
quantile(dd$Reaction_time)
```

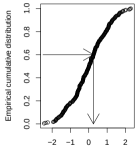
```
## 0% 25% 50% 75% 100%
## 255.7 293.9 307.8 316.4 331.9
```

QQ-plot continued...



The *theoretical quantiles* come from the normal distribution. The *sample quantiles* come from the distribution of our residuals.



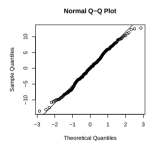


Sample quantiles of the residuals



Another useful diagnostic plot: The QQ-plot

The so-called quantile-quantile (QQ) plot. The quantiles of the observed distribution are plotted against the quantiles of the respective theoretical (normal) distribution:



If the points lie approximately on a straight line, the data is fairly normally distributed.

This is often "tested" by eye, and needs some experience.



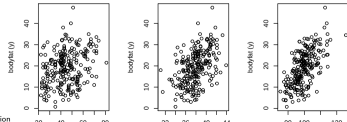
Multiple linear regression

Bodyfat example

We have so far modeled bodyfat in dependence of bmi, that is: $(bodyfat)_i = \alpha + \beta \cdot bmi_i + \epsilon_i$.

However, other explanatory variables might also be relevant for an accurate prediction of bodyfat.

Examples: Age, neck fat (Nackenfalte), hip circumference, abdomen circumference etc.



- 1. Is the **ensemble** of all explanatory variables associated with the response?
- 2. If yes, which explanatory variables are associated with the response?
- 3. What proportion of response variability (σ_y^2) is explained by the model?

Multiple linear regression model



The idea is simple: Just extend the linear model by additional predictors.

▶ Given several influence explanatory variables $x_i^{(1)}, \ldots, x_i^{(m)}$, the straightforward extension of the simple linear model is

$$y_i = \beta_0 + \beta_1 x_i^{(1)} + \beta_2 x_i^{(2)} + \ldots + \beta_m x_i^{(m)} + \epsilon_i$$
 with $\epsilon_i \sim N(0, \sigma^2)$.

▶ The parameters of this model are $\beta = (\beta_0, \beta_1, \dots, \beta_m)$ and σ^2 .

The components of β are again estimated using the **least squares** method. Basically, the idea is (again) to minimize

$$\sum_{i=1}^{n} e_i^2$$

with

$$e_i = y_i - (\beta_0 + \beta_1 x_i^{(1)} + \beta_2 x_i^{(2)} + \ldots + \beta_m x_i^{(m)})$$

It is a bit more complicated than for simple linear regression, see Section 3.4 of the Stahel script.

Some **linear algebra** is needed to understand these sections, but we do not look into this for the moment.





Let us regress the proportion (%) of bodyfat (from last week) on the predictors **bmi** and **age** simultaneously. The model is thus given as

$$(bodyfat)_i = \beta_0 + \beta_1 \cdot bmi_i + \beta_2 \cdot age_i + \epsilon_i$$
,
with $\epsilon_i \sim N(0, \sigma^2)$.



Multiple linear regression with R

Let's now fit the model with R, and quickly glance at the output:

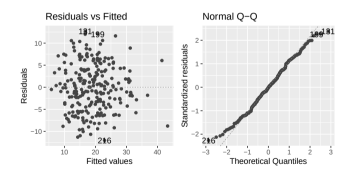
```
r.bodyfatM <- lm(bodyfat ~ bmi + age, d.bodyfat)
summary(r.bodyfatM)</pre>
```

```
##
## Call:
## lm(formula = bodyfat ~ bmi + age, data = d.bodyfat)
## Residuals:
       Min
                 10 Median
                                           Max
## -12.0415 -3.8725 -0.1237 3.9193 12.6599
## Coefficients:
               Estimate Std. Error t value Pr(>|t|)
## (Intercept) -31.25451
                           2.78973 -11.203 < 2e-16 ***
                1.75257
## bmi
                        0.10449 16.773 < 2e-16 ***
## age
                0.13268
                           0.02732
                                    4.857 2.15e-06 ***
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
## Residual standard error: 5.329 on 240 degrees of freedom
## Multiple R-squared: 0.5803, Adjusted R-squared: 0.5768
## F-statistic: 165.9 on 2 and 240 DF, p-value: < 2.2e-16
```

Model checking



Before we look at the results, we have to check if the modelling assumptions are fulfilled:



This seems ok, so continue with answering questions 1-3.



Question 1: Are the explanatory variables associated with the response?

To answer question 1, we need to perform a so-called F-test. The results of the test are displayed in the final line of the regression summary. Here, it says:

So apparently (and we already suspected that) the model has some explanatory power.

*The F-statistic and -test is briefly recaptured in 3.1.f) of the Stahel script, but see also Mat183 chapter 6.2.5. It uses the fact that

$$\frac{SSQ^{(R)}/m}{SSQ^{(E)}/(n-p)} \sim F_{m,n-p}$$

follows an F-distribution with m and (n-p) degrees of freedom, where m are the number of variables, n the number of data points, p the number of β -parameters (typically m+1). $SSQ^{(E)} = \sum_{i=1}^n R_i^2$ is the squared sum of the residuals, and $SSQ^{(R)} = SSQ^{(Y)} - SSQ^{(E)}$ with $SSQ^{(y)} = \sum_{i=1}^n (y_i - \overline{y})^2$.



n is the number of data points

m is the number of explanatory variables in the regression model

p is the number of beta parameters estimated (e.g. intercept, plus a slope for each explanatory variable, hence p=m+1)

And the degrees of freedom for error are n-p



Question 2: Which variables are associated with the response?

summary(r.bodyfatM)\$coef

```
## (Intercept) -31.2545057 2.78973238 -11.203406 1.039096e-23
## bmi 1.7525705 0.10448723 16.773060 2.600646e-42
## age 0.1326767 0.02731582 4.857137 2.149482e-06
```

To answer this question, again look at the *t*-tests, for which the *p*-values are given in the final column. Each *p*-value refers to the test for the null hypothesis $\beta_0^{(j)}=0$ for explanatory variable $x^{(j)}$.

As in simple linear regression, the T-statistic for the j-th explanatory variable is calculated as

$$T_j = \frac{\hat{\beta}_j}{s_{\mathbf{P}}(\beta_j)} , \qquad (1)$$

with $se^{(\beta_j)}$ given in the second column of the regression output.

The distribution of this statistic is $T_j \sim t_{n-p}$.

Therefore: A "small" p-value indicates that the variable is relevant in the model Zurich Zurich

Here, we have

- \triangleright p < 0.001 for bmi
- ightharpoonup p < 0.001 for age

Thus both, bmi and age seem to be associated with bodyfat.

Again, a 95% CI for $\hat{\beta}_i$ can be calculated with R:

confint(r.bodyfatM)

```
## (Intercept) -36.7499929 -25.7590185
## bmi 1.5467413 1.9583996
## age 0.0788673 0.1864861
```

(The CI is again $[\hat{\beta} - c \cdot \sigma^{(\beta)}; \hat{\beta} + c \cdot \sigma^{(\beta)}]$, where c is the 97.5% quantile of the t-distribution with n - p degrees of freedom; compare to slides 38-40 of last week).



!However!:

The p-value and T-statistics should only be used as a **rough guide** for the "significance" of the coefficients.

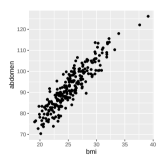
For illustration, let us extend the model a bit more, including also neck, hip and abdomen:

	Coefficient	95%-confidence interval	<i>p</i> -value
Intercept	-7.75	from -22.13 to 6.63	0.29
bmi	0.43	from -0.03 to 0.88	0.066
age	0.015	from -0.04 to 0.07	0.60
neck	-0.80	from -1.18 to -0.43	< 0.0001
hip	-0.32	from -0.53 to -0.11	0.003
abdomen	0.84	from 0.67 to 1.00	< 0.0001

It is now much less clear how strongly age (p=0.60) and bmi (p=0.07) are associated with bodyfat.

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Basically, the problem is that the variables in the model are correlated and therefore explain similar aspects of bodyfat. **Example:** Abdomen (Bauchumfang) seems to be a relevant predictor and it is obvious that abdomen and BMI are correlated:



This problem of collinearity is at the heart of many confusions of regression analysis, and we will talk about such issues later in the course (lectures 8 and 9).

Please see also IC: practical 4 (milk example) for an analysis and more thoughts.

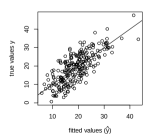


Question 3: Which proportion of variability is explained?

To answer this question, we can look at the multiple R^2 (see Stahel 3.1.h). It is a generalized version of R^2 for simple linear regression:

 R^2 for multiple linear regression is defined as the squared correlation between (y_1, \ldots, y_n) and $(\hat{y}_1, \ldots, \hat{y}_n)$, where the \hat{y} are the fitted values

$$\hat{y}_i = \hat{\beta}_0 + \hat{\beta}_1 x^{(1)} + \ldots + \hat{\beta}_m x^{(m)}$$





 R^2 is also called the *coefficient of determination* or "Bestimmtheitsmass", because it measures the proportion of the reponse's variability that is explained by the ensemble of all explanatory variables:

$$R^2 = SSQ^{(R)}/SSQ^{(Y)} = 1 - SSQ^{(E)}/SSQ^{(Y)}$$

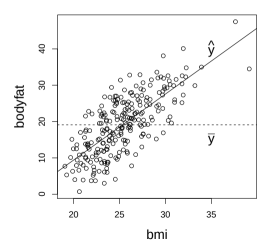
With

total variability = explained variability + residual variability

$$\sum_{i=1}^{n} (y_i - \overline{y})^2 = \sum_{i=1}^{n} (\hat{y}_i - \overline{y})^2 + \sum_{i=1}^{n} (y_i - \hat{y}_i)^2$$

$$SSQ^{(Y)} = SSQ^{(R)} + SSQ^{(E)}$$

This can be visualized for a model with only one predictor:



Let us look at the R^2 s from the three bodyfat models



(model r.bodyfat: $y \sim bmi$

model r.bodyfatM: $y \sim bmi + age$

model r.bodyfatM2: $v \sim bmi + age + neck + hip + abdomen$):

summary(r.bodyfat)\$r.squared

[1] 0.5390391

summary(r.bodyfatM)\$r.squared

summary(r.bodyfatM2)\$r.squared

[1] 0.718497

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[1] 0.5802956

The models explain 54%, 58% and 72% of the total variability of ν . It thus seems that larger models are "better". However, R^2 does always increase when new variables are included, but this does not mean that the model is more reasonable.

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Adjusted R^2



When the sample size n is small with respect to the number of variables m included in the model, an adjusted R^2 gives a better ("fairer") estimation of the actual variability that is explained by the explanatory variables:

$$R_a^2 = 1 - (1 - R^2) \frac{n-1}{n-m-1}$$

Why R_a^2 ?

It penalizes for adding more variables if they do not really improve the model!

Note: R_a may decrease when a new variable is added.

Interpretation of the coefficients



Apart from model checking and thinking about questions 1-3, it is probably even **more important to understand what you** *see.* Look at the output and ask yourself:

What does the regression output actually mean?

	Coefficient	95%-confidence interval	<i>p</i> -value
Intercept	-31.25	from -36.75 to -25.76	< 0.0001
bmi	1.75	from 1.55 to 1.96	< 0.0001
age	0.13	from 0.08 to 0.19	< 0.0001

Table 1: Parameter estimates of model 2.

Task in teams: Interpret the coefficients, 95% CIs and p-values.

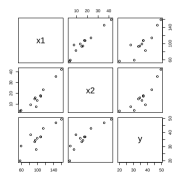


Example: Catheter Data

Catheter length (y) for heart surgeries depending on two characteristic variables $x^{(1)}$ and $x^{(2)}$ of the patients.

Aim: estimate y from $x^{(1)}$ and $x^{(2)}$ (n = 12).

Again look at the data first $(x^{(1)})$ and $x^{(2)}$ are highly correlated!):



Regression results with both variables: $R^2=0.81$, $R_a^2=0.76$, F-test p=0.0006. Zurich Turiersity of Lucient Turier Turi

	Coefficient	95%-confidence interval	<i>p</i> -value
Intercept	21.09	from 1.25 to 40.93	0.04
×1	0.077	from -0.25 to 0.40	0.61
×2	0.43	from -0.41 to 1.26	0.28

With
$$x_1$$
 only: $R^2 = 0.78$, $R_a^2 = 0.75$, F -test $p = 0.0002$

	Coefficient	95%-confidence interval	<i>p</i> -value
Intercept	12.13	from 2.66 to 21.59	0.017
×1	0.24	from 0.15 to 0.33	0.0002
	0.24	110111 0.13 to 0.33	0.000

With
$$x_2$$
 only: $R^2 = 0.80$, $R_a^2 = 0.78$, F-test $p = 0.0001$

	Coefficient	95%-confidence interval	<i>p</i> -value
Intercept	25.63	from 21.16 to 30.09	< 0.0001
x2	0.62	from 0.40 to 0.83	< 0.0001

- 1. Is x_1 an important explanatory variable?
- 2. Is x_2 an important explanatory variable?
- 3. Are both explanatory variables needed in the model?
- 4. Interpretation of the results?

Binary explanatory variables



So far, the explanatory variables x were always continuous.

In reality, there are no restrictions assumed with respect to the \boldsymbol{x} variables.

One very frequent data type are **binary** variables, that is, variables that can only attain values 0 or 1.

See section 3.2c of the Stahel script:

If the binary variable x is the only variable in the model $y_i = \beta_0 + \beta_1 x_i + \epsilon_i$, the model has only two predicted outcomes (plus error):

$$y_i = \begin{cases} \beta_0 + \epsilon_i & \text{if } x_i = 0\\ \beta_0 + \beta_1 + \epsilon_i & \text{if } x_i = 1 \end{cases}$$

Example: Smoking variable in Hg Study



For the 59 mothers in the Hg study, check if their smoking status (0=no,1=yes) influences the Hg-concentration in their urine.

We fit the following linear regression model:

$$\log(Hg_{urin})_{i} = \beta_{0} + \beta_{1} \cdot x_{i}^{(1)} + \beta_{2} \cdot x_{i}^{(2)} + \beta_{3} \cdot x_{i}^{(3)} + \epsilon_{i} ,$$

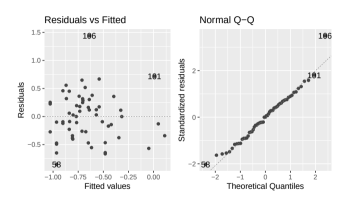
Where

- $ightharpoonup \log(Hg_{urin})$ is the urine mercury concentration.
- \triangleright $x^{(1)}$ is the binary smoking indicator (0/1), denoted as **dummy variable**.
- \triangleright $x^{(2)}$ the number of amalgam fillings.
- \triangleright $x^{(3)}$ the monthly number of marine fish meals.

(Remember from week 1 that the log of Hg concentrations is needed to obtain "useful" distributions.)



First check the modelling assumptions:



It seems ok, apart from one point (106) that could be categorized as an outlier. We ignore it for the moment.



The results table is given as follows:

	Coefficient	95%-confidence interval	<i>p</i> -value
Intercept	-1.03	from -1.24 to -0.81	< 0.0001
smoking	0.26	from -0.03 to 0.55	0.073
amalgam	0.098	from 0.05 to 0.14	< 0.0001
fish	0.032	from 0.01 to 0.06	0.019

There is some weak (p = 0.12) indication that smokers have an increased Hg concentration in their body. Their $log(Hg_{urin})$ is in average by 0.22 higher than for nonsmokers.

In principle, we have now – at the same time – fitted **two models:** one for smokers and one for non-smokers, assuming that the slopes of the remaining explanatory variables are the same for both groups.

Smokers:
$$y_i = -1.03 + 0.26 + 0.098 \cdot amalgam_i + 0.032 \cdot fish_i + \epsilon_i$$

Non-smokers: $y_i = -1.03 + 0.098 \cdot amalgam_i + 0.032 \cdot fish_i + \epsilon_i$

Categorical explanatory variables



Some explanatory variables indicate a **category**, for instance the species of an animal or a plant. This type of explanatory variable is termed **categorical**. For this there is trick: we can convert a categorical variable with k levels (for instance 3 species) into k dummy variables $x_i^{(j)}$ with

$$x_i^{(j)} = \begin{cases} 1, & \text{if the } i \text{th observation belongs to group } j. \\ 0, & \text{otherwise.} \end{cases}$$

Each of the explanatory variables $x^{(1)}, \ldots, x^{(k)}$ can then be included as a binary variable in the model

$$y_i = \beta_0 + \beta_1 x^{(1)} + \ldots + \beta_k x^{(k)} + \epsilon_i$$
.

However: this model is not identifiable. We could add a constant to $\beta_1, \beta_2, ... \beta_k$ and subtract it from β_0 , and the model would fit equally well to the data, so it cannot be decided which set of the parameters is best.



One of the k categories must be selected as a reference category and is not included in the model. Typically: the first category is the reference, thus $\beta_1 = 0$.

The model thus discriminates between the categories, such that (assuming $\beta_1 = 0$)

$$\hat{y}_i = \left\{ egin{array}{ll} eta_0, & ext{if } x_i^{(1)} = 1 \\ eta_0 + eta_2, & ext{if } x_i^{(2)} = 1 \\ ... \\ eta_0 + eta_k, & ext{if } x_i^{(k)} = 1 \end{array} \right..$$

!!Important to remember!!



(Common aspect that leads to confusion!)

Please note that a categorical variable with k categories requires k-1 parameters!

 \rightarrow The degrees of freedom are also reduced by k-1.

Example: Earthworm study

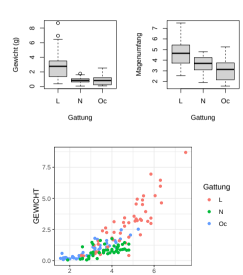


(Angelika von Förster und Burgi Liebst)

Die Dachse im Sihlwald ernähren sich zu einem grossen Prozentsatz von Regenwürmern. Ein Teil des Muskelmagens der Regenwürmer wird während der Passage durch den Dachsdarm nicht verdaut und mit dem Kot ausgeschieden. Wenn man aus der Grösse des Muskelmagenteilchens auf das Gewicht des Regenwurms schliessen kann, ist die Energiemenge berechnenbar, die der Dachs aufgenommen hat.

Frage: Besteht eine Beziehung zwischen dem Umfang des Muskelmagenteilchens und dem Gewicht des Regenwurms?

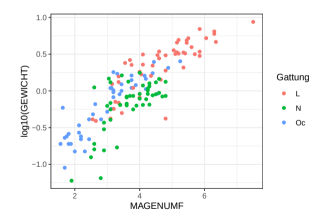
Data set of n = 143 worms with three species (Lumbricus, Nicodrilus, Octolasion), weight, stomach circumference (Magenumfang).





However, data inspection also suggests that there is not really a linear relationship between weight and stomach size – rather it looks exponential!

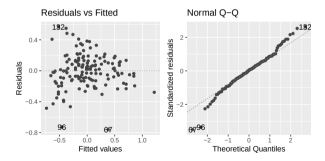
Therefore, log-transform the response (weight), and it looks much better:



Formulate a model with $\log_{10}(\text{Gewicht})$ as response and Magenumfang and Gabrurgh as explanatory variables. This is simple in R:

```
r.lm <- lm(log10(GEWICHT) ~ MAGENUMF + Gattung,d.wurm)
```

Before doing anything else, check the modeling assumptions:



 \rightarrow This seems ok (although the TA plot is a bit funnel-like).



Results:

	Coefficient	95%-confidence interval	<i>p</i> -value
Intercept	-1.10	from -1.29 to -0.91	< 0.0001
MAGENUMF	0.31	from 0.27 to 0.35	< 0.0001
GattungN	-0.22	from -0.32 to -0.13	< 0.0001
GattungOc	-0.039	from -0.15 to 0.07	0.48

$$R^2 = 0.76$$
, $R_a^2 = 0.75$.

- ▶ Question: Why is Gattung Lumbricus (L) not in the results table?
- ▶ Answer: L was chosen as the "reference category", thus $\beta_L = 0$.

Degrees of freedom: We had 143 data points. How many degrees of freedom are left for the residual error?

Answer: We estimated 4 parameters, thus 143 - 4 = 139.

Interpreting the results I



- $\beta_0 = -1.10$ is the intercept.
- $ightharpoonup eta_1 = 0.31$ is the slope for MAGENUMF.
- ho $\beta_2 = -0.22$ is the coefficient for Gattung=Nicodrilus.
- $ightharpoonup eta_3 = -0.039$ is the coefficient for Gattung =Octolasion.
- ▶ No coefficient is needed for Gattung Lumbricus, because $\beta_L = 0$.

We have now actually fitted three models, one model for each species:

Lumbricus: $\hat{y}_i = -1.10 + 0.31 \cdot MAGENUMF$

Nicodrilus: $\hat{y}_i = -1.10 + (-0.22) + 0.31 \cdot MAGENUMF$

Octolasion: $\hat{y}_i = -1.10 + (-0.039) + 0.31 \cdot MAGENUMF$

Interpreting the results II



Main question: Is there a relation between stomach size and body mass?

Results: MAGENUMF has a positive slope estimate with p < 0.0001, thus very strong evidence that the relation exists. Increasing MAGENUMF by 1 unit increases $\log_{10}(\text{GEWICHT})$ by +0.31.

Moreover, the $R^2 = 0.76$ is relatively high and almost identical to R_a^2 .

Interpreting the results III



Question: Is the "Gattung" explanatory variable relevant in the model, that is, do the model intercepts differ for the three species?

Problem: The *p*-values of the worm species are not very meaningful. They belong to tests that compare the intercept of a category with the intercept of the reference level (i.e., the *difference* in intercepts!). However, the question is whether the variable Gattung has an effect in total.

Solution: When a categorical explanatory variable with k levels is in the model, it occupies k-1 parameters. Therefore, the t-test needs to be replaced by the F-test.

F-test to compare models



F-Test zum Vergleich von Modellen. Die Frage sei, ob die q Koeffzienten $\beta_{j_1}, \beta_{j_2}, ..., \beta_{j_n}$ in einem linearen Regressionsmodell gleich null sein könnten.

- Null hypothese: $\beta_{j_1}=0\, \text{und}\, \beta_{j_2}=0\, \text{und} \dots \text{und}\, \beta_{j_e}=0$
- Teststatistik:

$$T = \frac{(SSQ^{(E)*} - SSQ^{(E)})/q}{SSQ^{(E)}/(n-p)};$$

 $\mathrm{SSQ}^{(E)*}$ ist die Quadratsumme des Fehlers im "kleinen" Modell, die man aus einer Regression mit den verbleibenden m-q X-Variablen erhält, und p die Anzahl Koeffizienten im "grossen" Modell (= m+1, falls das Modell einen Achsenabschnitt enthält, = m sonst).

• Verteilung von T unter der Nullhypolthese: $T \sim \mathcal{F}_{q,n-p}$, F-Verteilung mit q und n-p Freiheitsgraden.

Der Test heisst F-Test zum Vergleich von Modellen. Allerdings kann nur ein kleineres Modell mit einem grösseren verglichen werden, in dem alle X-Variablen des kleinen wieder vorkommen, also mit einem "umfassenderen" Modell. Der früher besprochene F-Test für das gesamte Modell (3.1.e) ist ein Spezialfall: das "kleine" Modell besteht dort nur aus dem Achsenabschnitt β_0 .

Remember: $F_{1,n-p} = t_{n-p}^2$

F-test for the earthworms



The function anova() in R does the F-test for categorical variables.

```
## Analysis of Variance Table

##
## Response: log10(GEWICHT)

## Df Sum Sq Mean Sq F value Pr(>F)

## MAGENUMF 1 19.7790 19.7790 409.69 < 2.2e-16 ***

## Gattung 2 1.3537 0.6768 14.02 2.842e-06 ***

## Residuals 139 6.7106 0.0483

## ---

## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

Note: Here, the F-value for Gattung is distributed as $F_{2,139}$ under the Null-Hypothesis.

This gives p = 2.842e - 06, thus a clear difference in the regression models for the three species ("Gattung is relevant").

anova(r.lm)



Plotting the earthworms results

All species have the same slope (this is a modeling assumption), but different intercepts:

