# Rcourse: Linear model

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Winter semester 2013-14

Background and basics

Analysis of variance

Model checking

#### Contents

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- Analysis of variance
- Model checking

## Intruitive linear regression

What is linear regression?

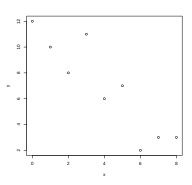
# Intruitive linear regression

What is linear regression?
It is the straight line that best approximates a set of points: y=a+b\*x
a is called the intercept and b the slope.

## Linear regression by eye

I give you the following points:

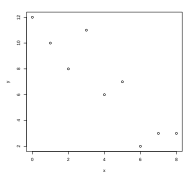
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x \leftarrow 0.8; y \leftarrow c(12,10,8,11,6,7,2,3,3); plot(x,y)
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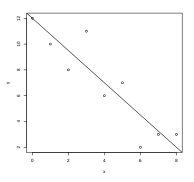


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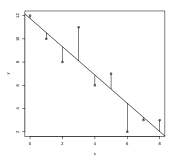
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#### Best fit in R

y is modelled as a function of x. In R this job is done by the function lm(). Lets try on the R console.

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The linear model does not explain all of the variation. The error is called "residual".

The purpose of linear regression is to minimize this error. But do you remember how we do this?

#### **Statistics**

We define the linear regression

$$y = \hat{a} + \hat{b} \cdot x$$

by minimizing the sum of the square of the residuals:

$$(\hat{a},\hat{b}) = \arg\min_{(a,b)} \sum_{i} (y_i - (a+b\cdot x_i))^2$$

This assumes that a, b exist, so that for all  $(x_i, y_i)$ 

$$y_i = a + b \cdot x_i + \varepsilon_i$$

where all  $\varepsilon_i$  are independent and follow the normal distribution with varaince  $\sigma^2$ .

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We can calculate  $\hat{a}$  und  $\hat{b}$  by

$$\hat{b} = \frac{\sum_{i} (y_{i} - \bar{y}) \cdot (x_{i} - \bar{x})}{\sum_{i} (x_{i} - \bar{x})^{2}} = \frac{\sum_{i} y_{i} \cdot (x_{i} - \bar{x})}{\sum_{i} (x_{i} - \bar{x})^{2}}$$

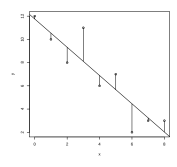
and

$$\hat{a} = \bar{y} - \hat{b} \cdot \bar{x}.$$



## Back to our example

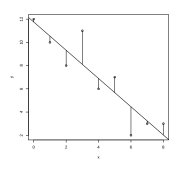
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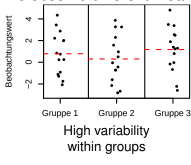


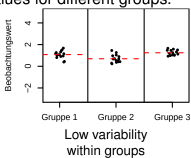
```
regr.obj <- lm(y x)
fitted <- predict(regr.obj)
plot(x,y); abline(regr.obj)
for(i in 1:9)
{
lines(c(x[i],x[i]),c(y[i],fitted[i])
}</pre>
```

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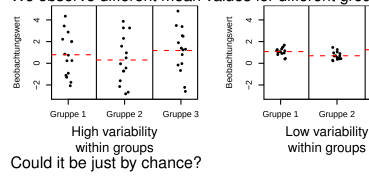
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I am sure you all remember from statistic courses: We observe different mean values for different groups.





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It depends from the variability of the group means and of the values within groups.

Gruppe 3

ANOVA-Table ("ANalysis Of VAriance")

	Degrees of free- dom (DF)	Sum of squares (SS)	Mean sum of squares (SS/DF)	<i>F</i> -Value
Groups	1	88.82	88.82	30.97
Residuals	7	20.07	2.87	

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Under the hypothesis  $H_0$  "the group mean values are equal" (and the values are normally distributed)

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We can reject  $H_0$ .

## **ANOVA** in R

In R ANOVA is performed using summary.aov() and summary().

These functions apply on a regression: result of command lm().

summary.aov() gives you only the ANOVA table whereas summary() outputs other information such as Residuals, R-square etc...

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Lets see a couple of examples with self-generated data in R.

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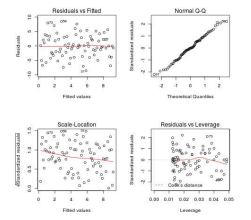
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In R you can do that directly by using the function plot() on your regression object.

Lets try on one example. We will focus on the first two graphs.

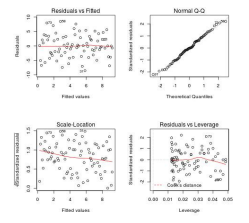
# Model checking: Good example

#### This is how it should look like:



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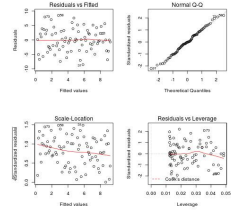
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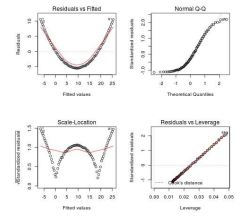
#### This is how it should look like:



- On the first graph, we should see no trend (equal variance).
- On the second graph, points should be close to the line (normality).

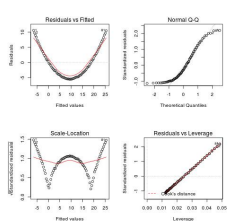
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What do you conclude?