# Module 3: Linear Regression

TMA4268 Statistical Learning V2021

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January 25 and 26, 2021

Last update: January 19, 2021

# Acknowledgements

- A lot of this material stems from Mette Langaas and her TAs (especiall Julia Debik). I would like to thank Mette for the permission to use her material!
- Some of the figures and slides in this presentation are taken (or are inspired) from James et al. (2013).

#### Introduction

#### Learning material for this module

 James et al. (2013): An Introduction to Statistical Learning, Chapter 3 (skip 3.5).

We need more statistical theory than is presented in the textbook, which you find in this module page.

#### What will you learn?

- Simple linear regression:
  - Model and assumptions
  - Least squares
  - Testing and confidence intervals
- Multiple linear regression:
  - The use of matrix algebra
  - Distribution of estimators
  - Assessing model fit, model selection
  - Confidence and prediction ranges
- Assessing model fit / residual analysis
- Qualitative predictors
- Interactions

# Linear regression

- Very simple approach for *supervised learning*.
- Parametric.
- Quantitative response vs. one or several explanatory variables.
- Aims:
  - Prediction "black box"
  - Explanation understanding the relationship between explanatory variables and the response
- Is linear regression too simple? Maybe, but very useful. Important to *understand* because many learning methods can be seen as generalization of linear regression.

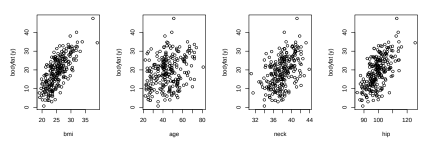
### Motivating example: Prognostic factors for body fat

(From Theo Gasser & Burkhardt Seifert Grundbegriffe der Biostatistik)

Body fat is an important indicator for overweight, but difficult to measure.

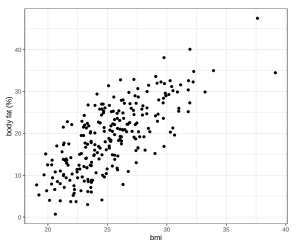
**Question:** Which factors allow for precise estimation (prediction) of body fat?

Study with 243 male participants, where body fat (%) and BMI and other predictors were measured. Some scatterplots<sup>1</sup>:



<sup>&</sup>lt;sup>1</sup>The data to reproduce these plots and analyses can be found here: https://github.com/stefaniemuff/statlearning/tree/master/3LinReg/data

For a good predictive model we need to dive into *multiple linear* regression. However, wer start with the simple case of only one predictor variable:



#### Interesting questions

- 1. How good is BMI as a predictor for body fat?
- 2. How strong is this relationship?
- 3. Is the relationship linear?
- 4. Are also other variables associated with bodyfat?
- 5. How well can we predict the bodyfat of a person?

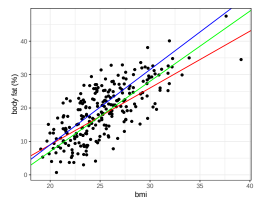
# Simple Linear Regression

- One quantitative response Y is modelled
- from one covariate x (=simple),
- and the relationship between Y and x is assumed to be *linear*.

If the relation between Y and x is perfectly linear, all instances of (x,Y), given by  $(x_i,y_i)$ ,  $i=1,\ldots,n$ , lie on a straight line and fulfill

$$y_i = \beta_0 + \beta_1 x_i .$$

But which is the "true" or "best" line, if the relationship is not exact?



**Task:** Estimate the intercept and slope parameters (by "eye") and write it down (we will look at the "best" answer later).

#### It is obvious that

- the linear relationship does not describe the data perfectly.
- another realization of the data (other 243 males) would lead to a slightly different picture.

⇒ We need a **model** that describes the relationship between BMI and bodyfat.

### The simple linear regression model

In the linear regression model the dependent variable Y is related to the independent variable x as

$$Y = \beta_0 + \beta_1 x + \varepsilon$$
,  $\varepsilon \sim N(0, \sigma^2)$ .

In this formulation Y is a random variable  $Y \sim N(\beta_0 + \beta_1 x, \sigma^2)$  where

$$Y = \underbrace{\text{expected value}}_{\mathsf{E}(Y) = \beta_0 + \beta_1 x} + \underbrace{\text{error}}_{\varepsilon}.$$

#### Note:

- The model for Y given x has three parameters:  $\beta_0$  (intercept),  $\beta_1$  (slope coefficient) and  $\sigma^2$ .
- x is the *independent/ explanatory / regressor* variable.
- ullet Y is the <u>dependent</u> / <u>outcome</u> / <u>response</u> variable.

### Modeling assumptions

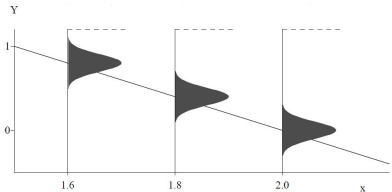
The central assumption in linear regression is that for any pairs  $(x_i, Y_i)$ , the error  $\varepsilon_i \sim N(0, \sigma^2)$ . This implies

- a) The expected value of  $\varepsilon_i$  is 0:  $\mathsf{E}(\varepsilon_i) = 0$ .
- b) All  $\varepsilon_i$  have the same variance:  $\mathsf{Var}(\varepsilon_i) = \sigma^2$ .
- c) All  $\varepsilon_i$  are normally distributed.
- d)  $\varepsilon$  is independent of any variable, observation number etc.
- e)  $\varepsilon_1, \varepsilon_2, \dots, \varepsilon_n$  are independent of each other.

## Visualization of the regression assumptions

The assumptions about the linear regression model lie in the error term

$$\varepsilon \sim N(0,\sigma^2)$$
 .



Note: The true regression line goes through  $\mathsf{E}(Y)$ .

## Parameter estimation ("model fitting")

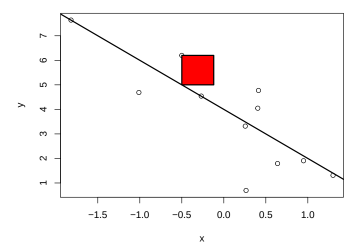
In a regression analysis, the task is to estimate the **regression** coefficients  $\beta_0$ ,  $\beta_1$  and the **residual variance**  $\sigma^2$  for a given set of (x,y) data.

- **Problem:** For more than two points  $(x_i, y_i)$ , i = 1, ..., n, there is generally no perfectly fitting line.
- **Aim**: We want to find the parameters (a, b) of the best fitting line Y = a + bx.
- Idea: Minimize the deviations between the data points  $(x_i,y_i)$  and the regression line.

But what are we actually going to minimize?

#### Least squares

Remember the **Least Squared Method**. Graphically, we are minimizing the sum of the squared distances over all points:



 Mathematically, a and b are estimated such that the sum of squared vertical distances (residual sum of squares)

$$RSS = \sum_{i=1}^{n} e_i^2 , \quad \text{where} \quad e_i = y_i - (a + bx_i)$$

is being minimized.

- The respective "best" estimates are called  $\hat{\beta}_0$  and  $\hat{\beta}_1$ .
- We can predict the value of the response for a (new) observation of the covariate at x.

$$\hat{y} = \hat{\beta}_0 + \hat{\beta_1} x.$$

• The *i*-th *residual* of the model is the difference between the *i*-th *observed* response value and the *i*-th *predicted* value, and is written as:

$$e_i = Y_i - \hat{y}_i.$$

• We may regard the residuals as *predictions* (not estimates) of the error terms  $\varepsilon_i$ .

(The error terms are random variables and can not be estimated - they can be predicted. It is only for parameters that we speak about estimates.)

#### Least squares estimators:

Using n observed independent data points

$$(x_1,y_1),(x_2,y_2),\dots,(x_n,y_n)\ ,$$

the least squares estiamtes for simple linear regression are given as

$$\hat{\beta}_0 = \bar{y} - \hat{\beta}_1 \bar{x} \tag{1}$$

and

$$\hat{\beta}_1 = \frac{\sum_{i=1}^n (x_i - \bar{x})(y_i - \bar{y})}{\sum_{i=1}^n (x_i - \bar{x})^2} = \frac{Cov(x, y)}{Var(x)} , \qquad (2)$$

where  $\bar{y} = \frac{1}{n} \sum_{i=1}^{n} y_i$  and  $\bar{x} = \frac{1}{n} \sum_{i=1}^{n} x_i$  are the sample means.

This is something you should have proven in your previous statistics classes; if you forgot how to get there, please check again, e.g. in chapter 11 of the book by Walepole et al. (2012), see here.

## Do-it-yourself "by hand"

Go to the Shiny gallery and try to "estimate" the correct parameters.

You can do this here:

https://gallery.shinyapps.io/simple\_regression/

## Example continued: Body fat

Assume a linear relationship between the % bodyfat (bodyfat) and the BMI (bmi), we can get the LS estimates using R as follows:

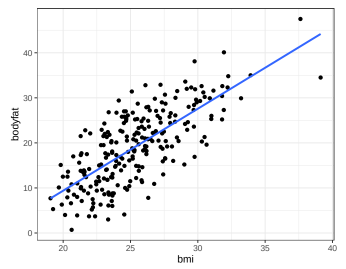
```
r.bodyfat = lm(bodyfat ~ bmi, data = d.bodyfat)
```

The estimates (and more information) can be obtained as follows:

```
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 16.787522 2.063854e-42
```

We see that the model fits the data quite well. It captures the essence. It looks that a linear relationship between bodyfat and bmi is a good approximation.



#### Questions:

- The blue line gives the estimated model. Explain what the line means in practice. Is this result plausible?
- Compare the estimates for  $\beta_0$  and  $\beta_1$  to the estimates you gave at the beginning were you close?
- How does this relate to the *true* (population) model?
- By looking at the spread of the points around the line, can you detect any violations of the modelling assumptions?
- Finally: What could the regression line look like if another set of 243 males were used for estimation?

# Uncertainty in the estimates $\hat{\beta}_0$ and $\hat{\beta}_1$

Note:  $\hat{\beta}_0$  and  $\hat{\beta}_1$  are themselves  $random\ variables$  and as such contain uncertainty!

Let us look again at the regression output, this time only for the coefficients. The second column shows the standard error of the estimate:

```
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 16.787522 2.063854e-42
```

 $\rightarrow$  The logical next question is: what is the distribution of the estimates?

# Distribution of the estimators for $\hat{\beta}_0$ and $\hat{\beta}_1$

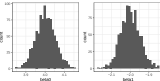
To obtain an intuition, we generate data points according to model

$$y_i = 4 - 2x_i + \varepsilon_i \ , \quad \varepsilon_i \sim N(0, 0.5^2). \label{eq:yi}$$

In each round, we estimate the parameters and store them:

```
set.seed(1)
niter <- 1000
pars <- matrix(NA, nrow = niter, ncol = 2)
for (ii in 1:niter) {
    x <- rnorm(100)
    y <- 4 - 2 * x + rnorm(100, 0, sd = 0.5)
    pars[ii, ] <- lm(y - x)$ccef
}</pre>
```

Doing it 1000 times, we obtain the following distributions for  $\hat{\beta}_0$  and  $\hat{\beta}_1$ :



#### Accuracy of the parameter estimates

 The standard errors of the estimates are given by the following formulas:

$$\mathsf{Var}(\hat{\beta}_0) = \mathrm{SE}(\hat{\beta}_0)^2 = \sigma^2 \Big[ \frac{1}{n} + \frac{\bar{x}^2}{\sum_{i=1}^n (x_i - \bar{x})^2} \Big]$$

and

$$\mathsf{Var}(\hat{\beta}_1) = \mathrm{SE}(\hat{\beta}_1)^2 = \frac{\sigma^2}{\sum_{i=1}^n (x_i - \bar{x})^2}.$$

•  $Cov(\hat{\beta}_0, \hat{\beta}_1)$  is in general different from zero.

**Note**: We will *derive a general version* of these formulas for multiple linear regression, because without matrix notation this is very cumbersome.

Under the assumption that  $\varepsilon \sim N(0, \sigma^2)$ , and assuming  $\hat{\beta}_0$  and  $\hat{\beta}_1$  are estimated as in formulas (1) and (2), we have in addition that

$$\hat{\beta}_0 \sim N(\beta_0, \sigma_{\beta_0}^2) \quad \text{and} \quad \hat{\beta}_1 \sim N(\beta_1, \sigma_{\beta_1}^2) \;.$$

**Again**: We will derive this in the multiple linear regression version in more generality.

## Design issue with data collection

Recall that

$$SE(\hat{\beta}_1)^2 = \frac{\sigma^2}{\sum_{i=1}^n (x_i - \bar{x})^2}$$
,

thus for a given  $\sigma^2$ , the standard error is only dependent on the design of the  $x_i$ 's!

- Would we like the  $SE(\hat{\beta}_1)^2$  large or small? Why?
- If it is possible for us to choose the  $x_i$ 's, which strategy should we use to choose them?
- Assume x can take values from 1 to 10 and we choose n = 10 values. Which is the best design?
  - evenly in a grid: [1, 2, 3, 4, 5, 6, 7, 8, 9, 10].
  - only lower and upper value: [1, 1, 1, 1, 1, 10, 10, 10, 10, 10].
  - randomly drawn from a uniform distribution on [1, 10].

```
x1 = seq(1:10)
x2 = c(rep(1, 5), rep(10, 5))
x3 = runif(10, 1, 10)

ss1 = sum((x1 - mean(x1))^2)
ss2 = sum((x2 - mean(x2))^2)
ss3 = sum((x3 - mean(x3))^2)

print(c(ss1, ss2, ss3))
```

 $\rightarrow$  The second design - all observations at extremes - is best!

## [1] 82.50000 202.50000 49.78026

### Residual standard error (RSE)

- **Problem**:  $\sigma$  is usually not known, but needs to be estimated<sup>2</sup>.
- Remember: The residual sum of squares is  $RSS = \sum_{i=1}^{n} (y_i \hat{\beta}_0 \hat{\beta}_1 x_i)^2.$
- An estimate of  $\sigma$ , the residual standard error, RSE, is given by

$$\hat{\sigma} = \text{RSE} = \sqrt{\frac{1}{n-2} \text{RSS}} = \sqrt{\frac{1}{n-2} \sum_{i=1}^{n} (y_i - \hat{y}_i)^2} \ .$$

• So actually we have

$$\hat{\mathrm{SE}}(\hat{\beta}_1)^2 = \frac{\hat{\sigma}^2}{\sum_{i=1}^n (x_i - \bar{x})^2} \ ,$$

but we usually just write  $SE(\hat{\beta}_1)^2$  (without the extra hat).

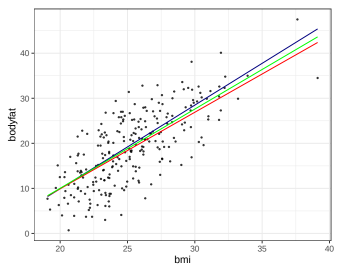
 $<sup>^{2}\</sup>sigma^{2}$  is the *irreducible error* variance.

The estimated standard errors can be seen using the summary() function:

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 16.787522 2.063854e-42
```

To illustrate this point further, fit again the bodyfat example, but each time with only half of the data (randomly selected points each time). See how the model fit varies:



# Testing and Confidence Intervals

After the regression parameters and their uncertainties have been estimated, there are typically two fundamental questions:

- 1. "Are the parameters compatible with some specific value?" Typically, the question is whether the slope  $\beta_1$  might be 0 or not, that is: "Is x an informative predictor or not?"
  - $\rightarrow$  This leads to a **statistical test**.

- 2. "Which values of the parameters are compatible with the data?"
  - $\rightarrow$  This leads us to determine **confidence intervals**.

Let's first go back to the output from the bodyfat example:

#### 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 16.787522 2.063854e-42
```

Besides the estimate and the standard error (which we discussed before), there is a t value and a probability Pr(>|t| that we need to understand.

How do these things help us to answer the two questions above?

#### Testing the effect of a covariate

Remember: in a statistical test you first need to specify the *null hypothesis*. Here, typically, the null hypothesis is

$$H_0: \quad \beta_1 = 0$$
.

In words:  $H_0$  = "There is no relationship between X and Y."

- Note 1: However, you might want to test against another null hypothesis, like  $\beta_1 = c$ .
- Note 2: Included in  $H_0$  is the assumption that the data follow the simple linear regression model!

Here, the alternative hypothesis is given by

$$H_A: \beta_1 \neq 0$$

Remember: To carry out a statistical test, we need a *test statistic*. This is some type of **summary statistic** that follows a known distribution under  $H_0$ . For our purpose, we use the so-called T-statistic

$$T = \frac{\hat{\beta}_1 - 0}{SE(\hat{\beta}_1)} \ .$$

Note: If you want to test against another value than  $\beta_1 = 0$ , the formula is

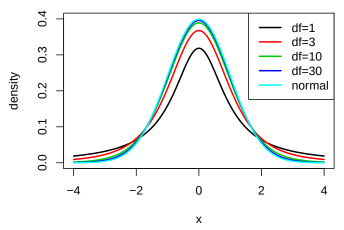
$$T = \frac{\hat{\beta}_1 - c}{SE(\hat{\beta}_1)} \ .$$

## Distribution of parameter estimators

We will *derive a general version* for multiple linear regression! Brief recap:

Under  $H_0$ , T has a t-distribution with n-2 degrees of freedom (n= number of data points; compare to Chapter 8.6 in Walepole et al. (2012)).

### Recap: The t-distribution



- The t-distribution has heavier tails than the normal distribution.
- For df  $\geq$  30 the t and Normal distribution are pretty similar.

### Hypothesis tests for bodyfat example

## bmi

So let's again go back to the bodyfat regression output:

```
summary(r.bodyfat)$coef
## Estimate Std. Error t value Pr(>|t|)
## (Intercept) -26.984368 2.7689004 -9.745518 3.921511e-19
```

1.818778 0.1083411 16.787522 2.063854e-42

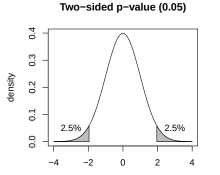
**Task**: Use the above formulas to derive the *T*-statistics.

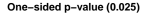
- The last column contains the *p-values* of the tests with  $H_0$ :  $\beta_0 = 0$  and  $\beta_1 = 0$ , respectively.
- The p-value for bmi is very small (p < 0.0001). What does this mean?

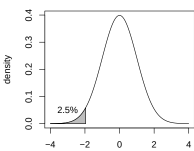
# Recap: Formal definition of the p-value

The *p*-value is the probability to observe a data summary (e.g., an average) that is at least as extreme as the one observed, given that the Null Hypothesis is correct.

**Example** (normal distribution): Assume the observed test-statistic leads to a z-value = -1.96  $\Rightarrow$  P(|z|  $\geq$  1.96) = 0.05 and P(z  $\leq$  -1.96) = 0.025 .







# Recap: Two types of errors

In the testing setup, we typically reject the null hypothesis if the p-value is small enough. Typical cutoffs for the significance level ( $\alpha$ ) are 5% or 1%.

However, this means we can make two types of errors:

- Type I error:
- Type II error:

## Cautionary notes regarding p-values:

- The (mis)use of *p*-values is heavily under critique in the scientific world!
- Simple yes/no decisions do often stand on very wiggly scientific ground.

We will discuss this a bit in the final module 12. The topic is connected to good/bad research practice, problems with "reproducibility" and scientific progress in general. See here:

- The p-value statement by ASA: https://amstat.tandfonline.com/doi/full/10.1080/00031305.2016.1154108#.Xh16iuExnhM
- Ideas to redefine what "statistical significane" means: https://www.nature.com/articles/s41562-017-0189-z
- A blog by the Scientific American: https://blogs.scientificamerican.com/observations/to-fix-the-reproducibility-crisis-rethink-how-we-do-experiments/

### Confidence intervals

- Confidence intervals (CIs) are a much more informative way to report results than *p*-values!
- The t-distribution<sup>3</sup> can be used to create confidence intervals for the regression parameters. The lower and upper limits of a 95% confidence interval for  $\beta_i$  are

$$\hat{\beta}_j \pm t_{(1-\alpha/2),n-2} \cdot \mathrm{SE}(\hat{\beta}_j) \quad j = 0,1.$$

- Interpretation of this confidence interval:
  - There is a 95% probability that the interval will contain the *true* value of  $\beta_i$ .
  - It is the range of parameter estimates that are *compatible* with the data.

 $<sup>^3</sup>$ If n is large, the normal approximation to the t-distribution can be used (and is used in the textbook).

Doing this for the bodfat example "by hand" is not hard. We have 241(=243-2) degrees of freedom:

```
coefs <- summary(r.bodyfat)$coef
beta <- coefs[2, 1]
sdbeta <- coefs[2, 2]
beta + c(-1, 1) * qt(0.975, 241) * sdbeta</pre>
```

## [1] 1.605362 2.032195

Even easier: directly ask R to give you the CIs.

**Interpretation:** for an increase in the bmi by one index point, roughly 1.82 percentage points more bodyfat are expected, and all true values for  $\beta_1$  between 1.61 and 2.03 are compatible with the observed data.

### Confidence and prediction ranges

• Based on the joint distribution of the intercept and slope it is possible to find the distribution for the linear predictor  $\hat{\beta}_0 + \hat{\beta}_1 x$ , and then confidence intervals for  $\beta_0 + \beta_1 x$ .

#### $\rightarrow$ Confidence range

• Accounting for the fact that we also have an error in the equation  $\varepsilon$ , we can also find the distribution of future observations.

### $\rightarrow$ Prediction range

We will discuss confidence and prediction ranges in the (more general) multiple linear regression setup.

# Model accuracy

#### Measured by

- 1. The **residual standard error (RSE)**, which provides an **absolute measure** of *lack of fit* (see above).
- 2. The **coefficient of determination**  $R^2$ , which measures the proportion of y's variance explained by the model (between 0 and 1), is a **relative measure** of *lack of fit*:

$$R^{2} = \frac{\text{TSS} - \text{RSS}}{\text{TSS}} = 1 - \frac{\text{RSS}}{\text{TSS}} = 1 - \frac{\sum_{i=1}^{n} (y_{i} - \hat{y}_{i})^{2}}{\sum_{i=1}^{n} (y_{i} - \bar{y}_{i})^{2}},$$

where

$$TSS = \sum_{i=1}^{n} (y_i - \bar{y})^2$$

is the  $total\ sum\ of\ squares$ , a measure for the total variability in Y.

# $R^2$ in simple linear regression

Note: In simple linear regression,  $\mathbb{R}^2$  is the squared correlation between the independent and the dependent variable.

Verify this by comparing  $\mathbb{R}^2$  from the bodyfat output to the squared correlation between the two variables:

```
summary(r.bodyfat)$r.squared
## [1] 0.5390391
cor(d.bodyfat$bodyfat, d.bodyfat$bmi)^2
## [1] 0.5390391
```

# Multiple Linear Regression

Remember that the bodyfat dataset contained much more information than only bmi and bodyfat:

- bodyfat: % of body fat.
- age: age of the person.
- weight: body weighth.
- height: body height.
- bmi: bmi.
- abdomen: circumference of abdomen.
- hip: circumference of hip.

#### Model

We assume

$$Y = \beta_0 + \beta_1 X_1 + \beta_2 X_1 + \dots + \beta_p X_p + \varepsilon , \qquad (3)$$

where  $X_j$  is the jth predictor and  $\beta_j$  the respective regression coefficient.

Assume we have n sampling units  $(x_{1i}, \ldots, x_{pi}, y_i)$ ,  $1 \le i \le n$ , such that each represent an instance of equation (3), we can use the data matrix

$$X = \begin{bmatrix} 1 & x_{11} & \dots & x_{1p} \\ 1 & x_{21} & \dots & x_{2p} \\ \vdots & \dots & \dots & \vdots \\ 1 & x_{n1} & \dots & x_{np} \end{bmatrix}$$

to write the model in matrix form:

$$Y = X\beta + \varepsilon$$

### Notation

- $Y:(n \times 1)$  vector of responses [e.g. one of the following: rent, weight of baby, pH of a lake, volume of a tree]
- $X : (n \times (p+1))$  design matrix, and  $x_i^T$  is a (p+1)-dimensional row row vector for observation i.
- $\beta: ((p+1)\times 1)$  vector of regression parameters  $(\beta_0,\beta_1,\dots,\beta_p)^{\top}$ .
- $\varepsilon : (n \times 1)$  vector of random errors.
- We assume that pairs  $(x_i^T, y_i)$  (i=1,...,n) are measured from independent sampling units.

Remark: other books, including the book in TMA4267 and TMA4315 define p to include the intercept. This may lead to some confusion about p or p+1 in formulas...

#### Classical linear model

$$Y = X\beta + \varepsilon$$

Assumptions:

- 1.  $\mathsf{E}(\varepsilon) = 0$ .
- 2.  $Cov(\varepsilon) = E(\varepsilon \varepsilon^T) = \sigma^2 I$ .
- 3. The design matrix has full rank, rank(X) = p + 1. (We assume n >> (p + 1).)

The classical normal linear regression model is obtained if additionally

4.  $\varepsilon \sim N_n(0, \sigma^2 I)$  holds. Here  $N_n$  denotes the *n*-dimensional multivarate normal distribution.

# Design matrix: Getting it in R

r.bodyfat = lm(bodyfat ~ bmi + age, data = d.bodyfat)

```
head(model.matrix(r.bodyfat))
##
     (Intercept) bmi age
## 1
              1 23.65 23
             1 23.36 22
## 2
## 3
             1 24.69 22
             1 24.91 26
## 4
## 5
             1 25 54 24
## 6
             1 26.48 24
head(d.bodyfat$bmi)
## [1] 23.65 23.36 24.69 24.91 25.54 26.48
head(d.bodyfat$age)
## [1] 23 22 22 26 24 24
```

## Distribution of the response vector

Assume that

$$Y = X\beta + \varepsilon \ , \quad \varepsilon \sim N_n(0,\sigma^2 I) \ .$$

 $\mathbf{Q}$ :

- What is the mean  $\mathsf{E}(Y)$ ?
- The covariance matrix Cov(Y) given X?
- Thus what is the distribution of Y?

 $\mathbf{A}$ :

$$Y \sim N_n(X\beta, \sigma^2 I)$$

### Parameter estimation for $\beta$

In multiple linear regression, the parameter vector  $\beta$  is estimated with *maximum likelihood* and *least squares*. These two methods give the same estimator when we assume the normal linear regression model.

With least sugres, we minimize the RSS for a multiple linear regression model:

$$\begin{aligned} \text{RSS} &= \sum_{i=1}^{n} (y_i - \hat{y}_i)^2 = \sum_{i=1}^{n} (y_i - \hat{\beta}_0 - \hat{\beta}_1 x_{i1} - \hat{\beta}_2 x_{i2} - \dots - \hat{\beta}_p x_{ip})^2 \\ &= \sum_{i=1}^{n} (y_i - x_i^T \hat{\beta})^2 = (Y - X \hat{\beta})^T (Y - X \hat{\beta}) \end{aligned}$$

The estimator is found by solving the system of (p+1) equations

$$\frac{\partial \mathrm{RSS}}{\partial \beta} = 0 \ .$$

 $\rightarrow$  Derivation on the board. See also here (from Mette in TMA4267).

Summing up:

The least squares and maximum likelihood estimator for  $\beta$ : is given like

$$\hat{\beta} = (X^T X)^{-1} X^T Y \ .$$

## Example continued

```
r.bodyfat3 <- lm(bodyfat ~ bmi + age + neck + hip + abdomen, data = d.bodyfat)
summary(r.bodyfat3)
##
## Call ·
## lm(formula = bodyfat ~ bmi + age + neck + hip + abdomen, data = d.bodyfat)
##
## Residuals:
      Min
               10 Median
## -9.3727 -3.1884 -0.1559 3.1003 12.7613
##
## Coefficients:
##
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) -7.74965
                       7.29830 -1.062 0.28939
## bmi
             0.42647
                        0.23133 1.844 0.06649 .
             0.01457 0.02783 0.524 0.60100
## age
## neck -0.80206 0.19097 -4.200 3.78e-05 ***
            -0.31764
                       0.10751 -2.954 0.00345 **
## hip
## abdomen
             0.83909
                       0.08418 9.968 < 2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 4.392 on 237 degrees of freedom
## Multiple R-squared: 0.7185, Adjusted R-squared: 0.7126
## F-statistic: 121 on 5 and 237 DF, p-value: < 2.2e-16
```

Reproduce the values under Estimate by calculating without the use of lm.

```
X = model.matrix(r.bodyfat3)
Y = d.bodyfat$bodyfat
betahat = solve(t(X) %*% X) %*% t(X) %*% Y
print(betahat)
```

```
## [,1]
## (Intercept) -7.74964673
## bmi 0.42647368
## age 0.01457356
## neck -0.80206081
## hip -0.31764315
## abdomen 0.83909391
```

# Distribution of the regression parameter estimator

Given

$$\hat{\boldsymbol{\beta}} = (X^T X)^{-1} X^T Y \ ,$$

#### what are

- The mean  $\mathsf{E}(\hat{\beta})$ ?
- The covariance matrix  $Cov(\hat{\beta})$ ?
- The distribution of  $\hat{\beta}$ ?

#### Hint: Use that

- $\hat{\beta} = CY$  with  $C = (X^T X)^{-1} X^T$ .
- $Y \sim N_n(X\beta, \sigma^2 I)$ .

#### Answers:

See Problem 3 of recommended exercise 3 (and the respective solutions) to derive that

$$\hat{\boldsymbol{\beta}} \sim N_{p+1}(\boldsymbol{\beta}, \underbrace{\sigma^2(X^TX)^{-1}}_{\text{covariance matrix}}) \ .$$

# The covariance matrix of $\hat{\beta}$ in R

The covariance matrix for the  $\hat{\beta}$  can be obtained as follows: vcov(r.bodyfat3)

```
##
              (Intercept)
                                   bmi
                                                age
                                                             neck
## (Intercept) 53.26521684 0.6774596810 -0.0780438125 -0.7219656479
## bmi
               0.67745968 0.0535131152 0.0005729015 -0.0120408637
## age
             -0.07804381 0.0005729015 0.0007745054 -0.0003432518
## neck
            -0.72196565 -0.0120408637 -0.0003432518 0.0364680351
## hip
            -0.54820573 -0.0058040729 0.0015239515 -0.0027159299
## abdomen
          0.16457979 -0.0110809165 -0.0011917596 -0.0007706161
##
                                abdomen
                      hip
## (Intercept) -0.548205733 0.1645797895
## bmi
            -0.005804073 -0.0110809165
## age
             0.001523951 -0.0011917596
## neck
             -0.002715930 -0.0007706161
## hip
            0.011558850 -0.0045707222
## abdomen
           -0.004570722 0.0070861066
```

How does this compare to simple linear regression? Not so easy to see a connection!

$$\begin{split} \hat{\beta}_0 &= \bar{Y} - \hat{\beta}_1 \bar{x} \text{ and } \hat{\beta}_1 = \frac{\sum_{i=1}^n (x_i - \bar{x})(Y_i - \bar{Y})}{\sum_{i=1}^n (x_i - \bar{x})^2}, \\ \hat{\beta} &= (X^T X)^{-1} X^T Y \ . \end{split}$$

**Exercise:** Verify the connection using  $\beta = (\beta_0, \beta_1)^{\top}$  and

$$X = \begin{bmatrix} 1 & x_{11} \\ 1 & x_{21} \\ 1 & \vdots \\ 1 & x_{n1} \end{bmatrix}.$$

# Four important questions

- 1. Is at least one of the predictors  $X_1,\dots,X_p$  useful in predicting the response?
- 2. Do all the predictors help to explain Y, or is only a subset of predictors useful?
- 3. How well does the model fit the data?
- 4. Given a set of predictor variables, what response value should we predict, and how accurate is our prediction?

### 1. Relationship between predictors and response?

Question is whether we could as well omit all predictor variables at the same time, that is

$$H_0: \beta_1 = \beta_2 = \ldots = \beta_p = 0$$
 vs.

 $H_1$ : at least one  $\beta_j$  is non-zero.

To answer this, we need the F-statistic

$$F = \frac{(\mathrm{TSS} - \mathrm{RSS})/p}{\mathrm{RSS}/(n-p-1)} \sim F_{p,(n-p-1)} \ ,$$

where total sum of squares TSS =  $\sum_i (y_i - \bar{y})^2$ , and residual sum of squares RSS =  $\sum_i (y_i - \hat{y}_i)^2$ . Under the Normal regression assumptions, F follows an  $F_{p,(n-p-1)}$  distribution (see Walepole et al. (2012), Chapter 8.7).

- If  $H_0$  is true, F is expected to be 1.
- Otherwise, we expect that the numerator is larger than the denominator (because the regression then explains a lot of variation) and thus F is greater than 1. For an observed value  $f_0$ , the p-value is given as

$$p = P(F_{p,n-p-1} > f_0)$$
.

#### Checking the F-value in the R output:

summary(r.bodyfat)

```
##
## Call:
## lm(formula = bodyfat ~ bmi + age, data = d.bodyfat)
##
## Residuals:
##
       Min
               1Q Median
                                3Q
                                       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 ***
## bmi
              1.75257 0.10449 16.773 < 2e-16 ***
## age
             ## ---
## Signif. codes: 0 '***' 0.001 '**' 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
```

#### Conclusion?

## More complex hypotheses

Sometimes we don't want to test if all  $\beta$ 's are zero at the same time, but only a subset  $1, \dots, q$ :

$$H_0:\beta_1=\beta_2=\dots=\beta_q=0$$

VS.

 $H_1$ : at least one different from zero.

Again, the F-test can be used, but now F is calculated like

$$F = \frac{(\mathrm{RSS}_0\text{-RSS})/(q)}{\mathrm{RSS}/(n-p-1)} \sim F_{q,n-p-1} \ ,$$

where

- Large model: RSS with p+1 regression parameters
- Small model:  $RSS_0$  with q+1 regression parameters

### Example in R

- Question: Do weight and height explain something of bodyfat, on top of the variables bmi and age?
- Fit both models and use the anova() function to carry out the F-test:

```
r.bodyfat.small = lm(bodyfat ~ bmi + age, data = d.bodyfat)
r.bodyfat.large = lm(bodyfat ~ bmi + age + weight + height, data = d.bodyfat)
anova(r.bodyfat.small, r.bodyfat.large)

## Analysis of Variance Table
##
## Model 1: bodyfat ~ bmi + age
## Model 2: bodyfat ~ bmi + age + weight + height
## Res.Df RSS Df Sum of Sq F Pr(>F)
## 1 240 6816.2
## 2 238 6702.9 2 113.28 2.0112 0.1361
```

# Inference about a single predictor $\beta_j$

A special case is

$$H_0: \beta_j = 0$$
 vs.  $H_1: \beta_j \neq 0$ 

- Nothing new: We did it for simple linear regression!
- This makes sense: it is known that (or you can try to show it yourself)

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

thus we can use a T-statistics with (n-p-1) degrees of freedom to get the p-value.

#### Going back again:

```
summary(r.bodyfat)$coef
```

```
## Estimate Std. Error t value Pr(>|t|)
## (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
```

#### However:

- Only checking the individual *p*-values is dangerous. **Why?**
- Not possible if  $n > p \to \text{need other approaches}$  (see e.g., Module 6).

# Inference about $\beta_i$ : confidence interval

• Using that

$$T_j = \frac{\hat{\beta}_j}{\mathrm{SE}(\hat{\beta}_j)} \sim t_{n-p-1} \ ,$$

we can create confidence intervals for  $\beta_j$  in the same manner as we did for simple linear regression (see slide 43). For example, when using the typical confidence level  $\alpha=0.05$  we have

$$\hat{\beta}_j \pm t_{0.975,n-p-2} \cdot \text{SE}(\hat{\beta}_j) \ . \label{eq:beta_j}$$

• Using R, this is very easy:

```
confint(r.bodyfat)
```

```
## 2.5 % 97.5 %
## (Intercept) -36.7499929 -25.7590185
## bmi 1.5467413 1.9583996
## age 0.0788673 0.1864861
```

## 2. Deciding on important variables

#### Overarching question:

#### Which model is the best?

#### But:

- Not clear what best means  $\rightarrow$  we need an objective criterion, like AIC, BIC, Mallows  $C_p$ , adjusted  $R^2$ .
- There are usually **many** possible models. For p predictors, we can build  $2^p$  different models.
- Cautionary note: Model selection can also lead to biased parameters estimates.
- $\rightarrow$  This topic is the focus of Module 6.

#### 3. Model Fit

We can again look at the two measures from simple linear regression:

• An absolute measure of lack of fit is again given by the estimate of  $\sigma$ , the residual standard error (RSE)

$$\hat{\sigma} = \text{RSE} = \sqrt{\frac{\text{RSS}}{n - p - 1}} \ .$$

•  $R^2$  is again the fraction of variance explained (no change from simple linear regression)

$$R^2 = \frac{\text{TSS} - \text{RSS}}{\text{TSS}} = 1 - \frac{\text{RSS}}{\text{TSS}} = 1 - \frac{\sum_{i=1}^n (y_i - \hat{y}_i)^2}{\sum_{i=1}^n (y_i - \bar{y}_i)^2} \ .$$

Simply speaking: "The higher  $R^2$ , the better."

#### However: Caveat with $R^2$

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

model 1:  $y \sim bmi$ 

model 2:  $y \sim bmi + age$ 

model 3:  $y \sim bmi + age + neck + hip + abdomen$ 

```
summary(r.bodyfatM1)$r.squared
## [1] 0.5390391
summary(r.bodyfatM2)$r.squared
## [1] 0.5802956
summary(r.bodyfatM3)$r.squared
## [1] 0.718497
```

The models explain 54%, 58% and 72% of the total variability of y. 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.

# 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 covariates:

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

 $R_a^2$  penalizes for adding more variables if they do not really improve the model!

 $\rightarrow R_a$  may decrease when a new variable is added.

## Model fit – in a broader sense

We will look at model validation / model checking later.

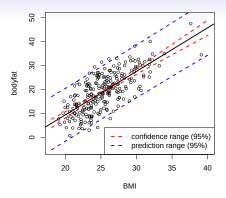
### 4. Predictions: Two questions

- 1. Which other regression lines are compatible with the observed data?
- $\rightarrow$  We can use  $\hat{\beta}_0, \dots, \hat{\beta}_p$  to estimate the  $\mathit{least\ squares\ plane}$

$$\hat{Y} = \hat{\beta}_0 + \hat{\beta}_1 X_1 + \ldots + \hat{\beta}_p X_p$$

as an approximation of  $f(X) = \beta_0 + \beta_1 X_1 + ... + \beta_p X_p$ . This leads to the *confidence interval*.

- 2. Where do future observations with a given x coordinate lie?
- $\rightarrow$  Even if we could predict  $\hat{Y} = f(X)$ , the *true* value Y varies around  $\hat{Y}$ . We can compute a *prediction interval* for new observations Y.



 Plotting the confidence and prediction intervals around all predicted values \(\hat{Y}\_0\) one obtains the confidence range or confidence band for the expected values of Y.

Note: The prediction range is much broader than the confidence range. Why?

# Calculation of the confidence intervals/range

Given a realization of  $X_1,\dots,X_p,$  say  $x_1^{(0)},\dots x_p^{(0)}.$  The question is:

Where does  $\hat{y}_0 = \hat{\beta}_0 + \hat{\beta}_1 x_1^{(0)} + ... \hat{\beta}_p x_p^{(0)}$  lie with a certain confidence (i.e., 95%)?

This question is not trivial, because  $\hat{\beta}_0, \dots \hat{\beta}_p$  are estimates from the data and contain uncertainty.

 $\to$  For the confidence range, only the uncertainty in the estimates  $\hat\beta_0,\dots\hat\beta_p$  matters.

# Calculation of the prediction intervals/range

Given a new value of  $X_1, \dots, X_p$ , say  $x_1^{(0)}, \dots x_p^{(0)}$ . The question is:

Where does a future observation lie with a certain confidence (i.e., 95%)?

To answer this question, we have to sum uncertainty over two components:

- 1. the uncertainty in the predicted value  $\hat{y}_0 = \hat{\beta}_0 + \hat{\beta}_1 x_1^{(0)} + \dots \hat{\beta}_p x_p^{(0)}$  (due to uncertainty in  $\hat{\beta}$ ).
- 2. the *irreducible error*  $\varepsilon_i \sim N(0, \sigma^2)$ .
- $\rightarrow$  The prediction intervals and range are always wider than the confidence intervals and range.

• Confidence and prediction intervals for given data points can be found in R using predict on an lm object (make sure that newdata is a data.frame with the same names as the original data).

```
fit = lm(bodyfat ~ bmi + age + abdomen, data = d.bodyfat)
newobs = d.bodyfat[1, ]
predict(fit, newdata = newobs, interval = "confidence", type = "response")

## fit lwr upr
## 1 13.17595 11.99122 14.36069
predict(fit, newdata = newobs, interval = "prediction", type = "response")

## fit lwr upr
## fit lwr upr
## 1 13.17595 3.951613 22.4003
```

Finally, we need to keep in mind that the model we work with is only an approximation of the reality. In fact,

#### In 2014, David Hand wrote:

In general, when building statistical models, we must not forget that the aim is to understand something about the real world. Or predict, choose an action, make a decision, summarize evidence, and so on, but always about the real world, not an abstract mathematical world: our models are not the reality – a point well made by George Box in his often-cited remark that "all models are wrong, but some are useful".

(Box 1979)

### Challenges - for model fit

- 1. Non-linearity of data
- 2. Correlation of error terms
- 3. Non-constant variance of error terms
- 4. Non-Normality of error terms
- 5. Outliers
- 6. High leverage points
- 7. Collinearity

## Recap of modelling assumptions in linear regression

To make valid inference from our model, we must check if our model assumptions are fulfilled! $^4$ 

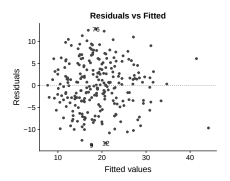
The assumption in linear regression is that the residuals follow a  $N(0,\sigma^2)$  distribution, implying that :

- 1. The expected value of  $\varepsilon_i$  is 0:  $\mathsf{E}(\varepsilon_i) = 0$ .
- 2. All  $\varepsilon_i$  have the same variance:  $Var(\varepsilon_i) = \sigma^2$ .
- 3. The  $\varepsilon_i$  are normally distributed.
- 4. The  $\varepsilon_i$  are independent of each other.

<sup>&</sup>lt;sup>4</sup>What is the problem if the assumptions are violated?

## Model checking tool I: Tukey-Anscombe diagram

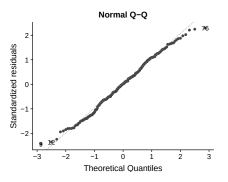
The *Tukey-Anscombe* diagram plots the residuals against the fitted values. For the bodyfat data it looks like this:



This plot is ideal to check if assumptions 1. and 2. (and partially 4.) are met. Here, this seems fine.

# Model checking tool II: The QQ-diagram

To check assumption 3., the quantiles of the observed distribution are plotted against the quantiles of the respective theoretical (normal) distribution:

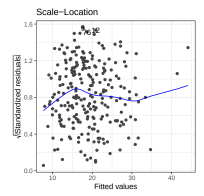


If the points lie approximately on a straight line, the data is fairly normally distributed. This is often "tested" by eye, and needs some experience.

### Model checking tool III: The scale-location plot

The scale-location plot is particularly suited to check the assumption of equal variances (homoscedasticity; assumption 2.).

The idea is to plot the square root of the (standardized) residuals  $\sqrt{|\tilde{r}_i|}$  against the fitted values  $\hat{y}_i$ . There should be *no trend*:



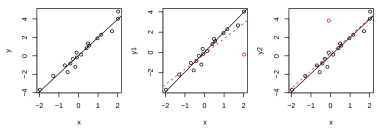
### Model checking tool IV: The leverage plot

- Mainly useful to determine outliers.
- To understand the leverage plot, we need to introduce the idea of the leverage.
- In simple regression, the leverage of individual i is defined as

$$H_{ii} = \frac{1}{n} + \frac{(x_i - \overline{x})^2}{\sum_{i'} (x_{i'} - \overline{x})^2} \ . \tag{4}$$

**Q:** When are leverages expected to be large/small?

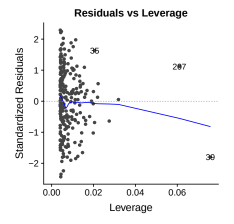
**Illustration**: Data points with  $x_i$  values far from the mean have a stronger leverage effect than when  $x_i \approx \overline{x}$ :



The outlier in the middle plot "pulls" the regression line in its direction and biases the slope.

Click here to do it manually!

In the leverage plot, (standardized) residuals  $\tilde{r_i}$  are plotted against the leverage  $H_{ii}$  (still for the bodyfat):



Critical ranges are the top and bottom right corners! Why?

## Leverages in multiple regression

- Leverage is defined as the diagonal elements of the so-called *hat*  $matrix \mathbf{H}^5$ , i.e., the leverage of the *i*-th data point is  $H_{ii}$  on the diagonal of  $\mathbf{H} = \mathbf{X}(\mathbf{X}^T\mathbf{X})^{-1}\mathbf{X}^T$ .
- Exercise: Verify that formula (4) comes out in the special case of simple linear regression.
- A large leverage indicates that the observation (i) has a large influence on the estimation results, and that the covariate values  $(x_i)$  are unusual.

<sup>&</sup>lt;sup>5</sup>Do you remember why **H** is called *hat matrix*?

### Different types of residuals?

It can be shown that the vector of residuals,  $e=(e_1,e_2,\ldots,e_n)$  have a normal (singular) distribution with

- E(e) = 0
- $Cov(e) = \sigma^2(I H),$

where  $H = X(X^TX)^{-1}X^T$ .

This means that the residuals (possibly) have different variance, and may also be correlated.

**Q:** Why is that a problem?

#### $\mathbf{A}$ :

- To check the model assumptions we want to look at the distribution of the error terms  $\varepsilon_i$ , to check that our errors are independent, homoscedastic (same variance for each observation), and not dependent on our covariates.
- However, we only have the residuals  $e_i$ , the "predictions" for  $\epsilon_i$ .
- It would have been great if the  $e_i$  have the same properties as  $\epsilon_i$ .
- $\rightarrow$  To make the  $e_i$  more "like  $\epsilon_i$ ", we use standardized or studentized residuals.

#### Standardized residuals:

$$r_i = \frac{e_i}{\hat{\sigma}\sqrt{1 - H_{ii}}}$$

where  $H_{ii}$  is the *i*th diagonal element of the hat matrix H.

In R you can get the standardized residuals from an lm-object (named fit) by rstandard(fit).

#### Studentized residuals:

$$r_i^* = \frac{e_i}{\hat{\sigma}_{(i)} \sqrt{1 - H_{ii}}}$$

where  $\hat{\sigma}_{(i)}$  is the estimated error variance in a model with observation number i omitted. It can be shown that it is possible to calculated the studentized residuals directly from the standardized residuals.

In R you can get the studentized residuals from an lm-object (named fit) by rstudent(fit).

### Diagnostic plots in R

See exercises: We use autoplot() from the ggfortify package in R to plot the diagnostic plots.

### Collinearity

In brief, collinearity refers to the situation when two or more predictors are correlated, thus encode (partially) for the same information.

#### Problems:

- Reduces the accuracy of the estimated coefficients  $\hat{\beta}_j$  (large SE!).
- Consequently, reduces power in finding effects (*p*-values become larger).

#### Solutions:

- Detect it by calculating the variance inflation factor (VIF).
- Remove the problematic variable.
- Or combine the collinear variables into a single new one.

**Todo:** Read in the course book p.99-102 (self-study).

# Other considerations in the regression model

- 1. Qualitative predictors  $(X_i)$ :
  - Binary covariate (e.g., male/female, smoker/non-smoker)
  - Categorical covariate (e.g., black/white/green)?
- 2. Extensions of the linear model
  - Interactions
  - Non-linear terms

#### Binary predictors

So far, the covariates X were always continuous.

In reality, there are no restrictions assumed with respect to the X variables.

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

If the binary variable x is the only variable in the model  $Y_i=\beta_0+\beta_1x_i+\varepsilon_i$ , the model has only two predicted outcomes (plus error):

$$Y_i = \left\{ \begin{array}{ll} \beta_0 + \varepsilon_i & \text{if } x_i = 0 \ , \\ \beta_0 + \beta_1 + \varepsilon_i & \text{if } x_i = 1 \ . \end{array} \right.$$

**Example**: Credit card data analysis in Section 3.3.1 in the ISLR book.

# Qualitative predictors with more than 2 levels

More generally, a covariate may indicate a **category**, for instance the species of an animal or a plant. This type of covariate is called a **factor**. The trick: convert a factor variable X with k levels (for instance 3 species) into k dummy variables  $X_j$  with

$$x_{ij} = \left\{ \begin{array}{ll} 1, & \text{if the $i$th observation belongs to group $j$.} \\ 0, & \text{otherwise.} \end{array} \right.$$

Each of the covariates  $x_1,\dots,x_k$  can then be included as a binary variable in the model

$$y_i = \beta_0 + \beta_1 x_1 + \ldots + \beta_k x_k + \varepsilon_i \ .$$

However: this model is *not identifiable*.<sup>6</sup>

<sup>&</sup>lt;sup>6</sup>What does that mean? I could add a constant to  $\beta_1, \beta_2, ... \beta_k$  and subtract it from  $\beta_0$ , and the model would fit equally well to the data, so it cannot be decided which set of the parameters is best.

**Solution:** 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 factor levels, such that (assuming  $\beta_1 = 0$ )

$$y_i = \left\{ \begin{array}{ll} \beta_0 + \varepsilon, & \text{if } x_{i1} = 1 \\ \beta_0 + \beta_2 + \varepsilon, & \text{if } x_{i2} = 1 \\ \dots & \\ \beta_0 + \beta_k + \varepsilon, & \text{if } x_{ik} = 1 \end{array} \right..$$

#### !Important to remember!

(Common aspect that leads to confusion!)

A factor covariate with k factor levels requires k-1 parameters!

 $\rightarrow$  The degrees of freedom of the fitted model are therefore reduced by k-1.

#### Example

library(ISLR)

We are now using the Credit dataset from the ISLR library.

```
data(Credit)
head(Credit)
##
         Income Limit Rating Cards Age Education Gender Student Married
## 1
     1
         14.891
                 3606
                         283
                                    34
                                              11
                                                    Male
                                                              No
                                                                     Yes
## 2
     2 106.025
                 6645
                         483
                                    82
                                              15 Female
                                                             Yes
                                                                     Yes
## 3
     3 104.593 7075
                        514
                                 4 71
                                              11
                                                   Male
                                                              Nο
                                                                      Nο
## 4
     4 148.924 9504
                       681
                                 3 36
                                              11 Female
                                                                      Nο
                                                              Nο
                                 2 68
## 5
    5 55.882 4897
                         357
                                              16
                                                   Male
                                                              No
                                                                     Yes
## 6
     6 80.180
                 8047
                         569
                                 4 77
                                              10
                                                   Male
                                                              Nο
                                                                      Nο
##
     Ethnicity Balance
## 1 Caucasian
                   333
## 2
         Asian
                   903
## 3
         Asian
                   580
## 4
         Asian
                   964
## 5 Caucasian
                   331
## 6 Caucasian
                  1151
```

Question: Do the Balances differ for different Ethnicities?

In R, a factor covariate can be used in the same way as a continuous predictor:

r.lm <- lm(Balance ~ Ethnicity, data = Credit)

```
summary(r.lm)
##
## Call:
## lm(formula = Balance ~ Ethnicity, data = Credit)
##
## Residuals:
##
      Min
              10 Median
                                    Max
## -531.00 -457.08 -63.25 339.25 1480.50
##
## Coefficients:
##
                    Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                     531.00 46.32 11.464 <2e-16 ***
## EthnicitvAsian
                  -18.69 65.02 -0.287 0.774
## EthnicityCaucasian -12.50 56.68 -0.221 0.826
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 460.9 on 397 degrees of freedom
## Multiple R-squared: 0.0002188, Adjusted R-squared: -0.004818
## F-statistic: 0.04344 on 2 and 397 DF, p-value: 0.9575
```

Interpretation? Do the ethnicities really differ? Check also the F-test in the last line of the summary output.

#### The "reference category"

In the above example we do not see a result for the EthnicityAfrican American. Why?

- African American is chosen to be the reference category.
- The results for EthnicityAsian and EthnicityCaucasian are differences with respect to the reference cateogry.
- R chooses the reference category in alphabetic order! This is sometimes not a relevant category.
- You can change the reference category:

```
library(dplyr)
Credit <- mutate(Credit, Ethnicity = relevel(Ethnicity, ref = "Caucasian"))
r.lm <- lm(Balance ~ Ethnicity, data = Credit)
summary(r.lm)$coef</pre>
```

Note: The differences are now with respect to the Caucasian category – the model is however exactly the same!

# Testing for a categorical predictor

**Question**: Is a qualitative predictor needed in the model?

For a predictor with more than two levels (like Ethnicity above), the Null Hypothesis is whether

$$\beta_1 = \dots = \beta_{k-1} = 0$$

at the same time.

 $\rightarrow$  We again need the F-test<sup>7</sup>, as always when we test for more than one  $\beta_j = 0$  simultaneously!

In R, this is done by the anova() function:

```
anova(r.lm)
## Analysis of Variance Table
##
## Response: Balance
## Df Sum Sq Mean Sq F value Pr(>F)
## Ethnicity 2 18454 9227 0.0434 0.9575
## Residuals 397 84321458 212397
```

<sup>&</sup>lt;sup>7</sup>remember that the F-test is a generalization of the t-test!

### Interactions: Removing the additivity assumption

We again look at the Credit dataset. We want to model the Balance as a function of Income and wheter the person is a student or not.

The model is given as

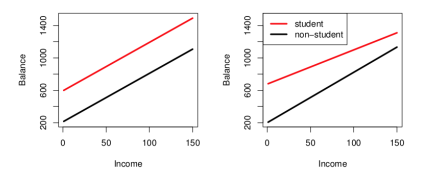
$$\text{Balance}_i = \beta_0 + \beta_1 \cdot \text{Income}_i + \beta_2 \cdot \text{Student}_i + \varepsilon_i \ ,$$

where Student is a binary variable. Thus we have a model that looks like

$$\text{Balance}_i = \left\{ \begin{array}{l} \beta_0 + \beta_2 + \beta_1 \cdot \text{Income}_i + \varepsilon_i \ , & \text{if $i$ is a student,} \\ \beta_0 + \beta_1 \cdot \text{Income}_i + \varepsilon_i & \text{otherwise.} \end{array} \right.$$

In R, we simply add Student to the model:

Caveat: This model assumes that students and non-students have the same slope for Income. Realistic? Let's look at the graphs:



 $\rightarrow$  We want a model that allows for different slopes!

#### Interaction terms

We formulate a new model that includes the interaction term (Income  $\cdot$  Student):

$$\mathrm{Balance}_i = \beta_0 + \beta_1 \cdot \mathrm{Income}_i + \beta_2 \cdot \mathrm{Student}_i + \beta_3 \cdot \mathrm{Income}_i \cdot \mathrm{Student}_i + \varepsilon_i \ ,$$

Thus we have a model that allows for different intercept and slope for the two groups:

$$\text{Balance}_i = \left\{ \begin{array}{l} \beta_0 + \beta_2 + (\beta_1 + \beta_3) \cdot \text{Income}_i + \varepsilon_i \ , & \text{if $i$ is a student,} \\ \beta_0 + \beta_1 \cdot \text{Income}_i + \varepsilon_i & \text{otherwise.} \end{array} \right.$$

In R, this is again quite simple:

#### Interpretation:

We allow the model to depend on the binary variable Student, such that

For a student:  $\hat{y} = 200.6 + 476.7 + (6.2 + -2.0) \cdot \text{Income}$ 

For a non-Student:  $\hat{y} = 200.6 + (6.2) \cdot \text{Income}$ 

**Question:** Is the interaction relevant here?

## The hierarchical principle

If we include an interaction in a model, we should also include the main effects, even if the p-values associated with the coefficients of the main effects are large (see p.89 in ISLR book).

#### More interactions

We can include interactions also between

- two continuous variables.
- a categorical variable with more than 2 levels and a continuous variable.
- $\rightarrow$  See exercises!

#### Non-linear terms

#### Linear regression is even more powerful!

- We have seen that it is possible to include continuous, binary or factorial covariates in a regression model.
- Even transformations of covariates can be included in (almost) any form. For instance the square of a variable  $X^2$

$$y_i = \beta_0 + \beta_1 x_i + \beta_2 x_i^2 + \varepsilon_i ,$$

which leads to a **quadratic** or **polynomial** regression (if higher order terms are used).

- Other common transformations are:
  - log
  - $\sqrt{..}$
  - sin, cos,

How can a *quadratic* regression be a *linear regression*??

#### Note:

The word *linear* refers to the *linearity in the coefficients*, and not on a linear relationship between Y and  $X_1, \ldots, X_n!$ 

**Question**: When would we need such a regression? Well, sometimes the world is not linear. In particular, if

- there is a theoretical/biological/medical reason to believe in a non-linear relationship, or
- the residual analysis indicates that there are non-linear associations in the data,

it can sometimes help to use transformations of a variable X.

 $\rightarrow$  In the later modules, we will discuss other more advanced non-linear approaches for addressing this issue.

# Further reading

• Videoes on YouTube by the authors of ISL, Chapter 3

#### References

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