# Chapter 2 Marginal Models for Continuous Data

#### 2.1 Simple Methods



- The reason why classical statistical techniques fail in the context of longitudinal data is that observations within subjects are correlated
  - > often the correlation between two repeated measurements decreases as the time span between those measurements increases
- The paired t-test accounts for this by considering subject-specific differences  $\Delta_i = Y_{i1} Y_{i2}$ 
  - by this reduces the number of measurements to just one per subject, which implies that classical techniques can be applied again



- In the case of more than 2 measurements per subject, similar simple techniques are often applied to reduce the number of measurements for the ith subject, from  $n_i$  to 1
  - > Analysis at each time point separately

  - > Analysis of endpoints
  - > Analysis of increments



- Analysis at each time point separately
  - ▶ General idea: The data are analyzed at each occasion separately

#### > Advantages:

- \* simple to interpret
- \* uses all available data

#### **Disadvantages:**

- \* does not consider 'overall' differences
- \* does not allow to study the evolution of differences
- \* problem of multiple testing
- \* possible problems with missing data



- Analysis of area under the curve (AUC)
  - ▶ General idea: For each subject, the area under her curve is calculated

$$AUC_i = (t_{i2} - t_{i1}) \times (y_{i2} + y_{i1})/2 + (t_{i3} - t_{i2}) \times (y_{i3} + y_{i2})/2 + \dots$$

Afterwards, these AUCs are analyzed

#### > Advantages:

- \* no problems of multiple testing
- \* does not explicitly assume balanced data
- \* compares 'overall' differences



- Analysis of area under the curve (AUC)
  - **Disadvantages:** 
    - \* uses only partial information
    - \* possible problems with missing data



#### Analysis of endpoints

▶ General idea: Assess differences only on the last time point

#### > Advantages:

- \* no problems of multiple testing
- \* does not explicitly assume balanced data

#### **Disadvantages:**

- \* applicable only in randomized trials
- \* does not consider 'overall' differences
- \* possible problems with missing data



#### Analysis of increments

 $\triangleright$  General idea: A simple method to compare evolutions between subjects, correcting for differences at baseline, is to analyze the subject-specific changes  $y_{in_i}-y_{i1}$ 

#### > Advantages:

- \* no problems of multiple testing
- \* does not explicitly assume balanced data

#### **Disadvantages:**

- \* uses partial information
- \* possible problems with missing data



- The AUC, endpoints and increments are examples of summary statistics
  - > such summary statistics summarize the vector of repeated measurements for each subject separately
- This leads to the following general procedure:
  - > **Step 1:** Summarize the data of each subject into one statistic
  - ▶ **Step 2:** Analyze the summary statistics, e.g. analysis of covariance to compare groups after correction for important covariates
- This way, the analysis of longitudinal data is reduced to the analysis of independent observations, for which classical statistical procedures are available



• However, all these methods have the disadvantage that (lots of) information is lost

This has led to the development of statistical techniques that overcome these disadvantages

#### 2.2 Review of Linear Regression



- ullet Suppose we have a continuous outcome Y measured cross-sectionally
  - $\triangleright$  Example: The serum bilirubin levels from the PBC dataset at baseline (i.e., time t=0)
- We are interested in making statistical inferences for this outcome, e.g.,
  - ▷ is there any difference between placebo and D-penicil corrected for the age and sex of the patients?
  - ▶ which factors best predict serum bilirubin levels?





• Definition of the linear regression model

$$\begin{cases} y_i = \beta_0 + \beta_1 x_{i1} + \ldots + \beta_p x_{ip} + \varepsilon_i \\ \\ \varepsilon_i \sim \mathcal{N}(0, \sigma^2) \end{cases}$$

#### where

 $\triangleright y_i$  denotes the outcome for subject i

 $\triangleright x_{i1}, \ldots, x_{ip}$  denote the p covariates for subject i

 $\triangleright \beta_0, \beta_1, \dots, \beta_p$  the regression coefficients

 $\triangleright \varepsilon_i$  the error term for subject i



• Example: For the PBC patients we postulate the linear regression model

$$\log(\mathtt{serBilir}_i) = eta_0 + eta_1 \mathtt{Age}_i + eta_2 \mathtt{D-penicil}_i + arepsilon_i, \quad arepsilon_i \sim \mathcal{N}(0, \sigma^2)$$

#### where

- $\triangleright$  serBilir<sub>i</sub> denotes the serum bilirubin of patient i at baseline
- $\triangleright$  Age<sub>i</sub> and D-penicil<sub>i</sub> denote the Age and whether patient i received D-penicil or placebo
- $\triangleright \beta_0$ ,  $\beta_1$ , and  $\beta_2$  are the regression coefficients
- $\triangleright \varepsilon_i$  are the error terms



- Behind this model there are several assumptions, some obvious, some hidden. In particular:
  - > serum bilirubin is assumed to be only related to Age and treatment
  - by the relation between serum bilirubin and Age is linear
  - by the effect of Age is the same whatever the treatment the patient took, and vice versa
  - b the error terms are normally distributed
  - > the variance of the error terms does not depend on neither Age nor D-penicil
  - > measurements are independent with each other



	Estimate	Std. Error	t value	Pr(> t )
(Intercept)	0.5395	0.2824	1.91	0.0570
age	0.0015	0.0056	0.28	0.7817
drugD-penicil	-0.0933	0.1174	-0.79	0.4274

#### Interpretation

- $\triangleright \beta_0 = 0.5$  average log(Ser. Bilir.) for Age = 0 and placebo patients
- $\triangleright \beta_1 = 0.0015$  increase in average log(Ser. Bilir.) for every year increase for patient with the same treatment
- $\triangleright \beta_2 = -0.1$  decrease in average log(Ser. Bilir.) when receiving D-penicil versus placebo for patients of the same age



- Linear regression model with *matrix notation* 
  - $\triangleright$  the linear regression model for the n subjects

$$y_1 = \beta_0 + \beta_1 x_{11} + \ldots + \beta_p x_{1p} + \varepsilon_1$$

$$y_2 = \beta_0 + \beta_1 x_{11} + \ldots + \beta_p x_{1p} + \varepsilon_2$$

:

$$y_n = \beta_0 + \beta_1 x_{n1} + \ldots + \beta_p x_{np} + \varepsilon_n$$



- Linear regression model with *matrix notation* 
  - $\triangleright$  the linear regression model for the n subjects

$$\begin{bmatrix} y_1 \\ y_2 \\ \vdots \\ y_n \end{bmatrix} = \begin{bmatrix} 1 & x_{11} & \dots & x_{1p} \\ 1 & x_{21} & \dots & x_{2p} \\ \vdots & & & \vdots \\ 1 & x_{n1} & \dots & x_{np} \end{bmatrix} \begin{bmatrix} \beta_0 \\ \beta_1 \\ \vdots \\ \beta_p \end{bmatrix} + \begin{bmatrix} \varepsilon_1 \\ \varepsilon_2 \\ \vdots \\ \varepsilon_n \end{bmatrix} \\
\mathbf{y} = \mathbf{X} \qquad \boldsymbol{\beta} + \boldsymbol{\varepsilon}$$



- Linear regression model with *matrix notation* 

  - $\triangleright \beta$ : parameter vector
  - ▷ *E*: measurement error vector



Maximum likelihood estimators

$$\begin{cases} \widehat{\beta} &= (X^{\top}X)^{-1}X^{\top}y \\ \\ \widehat{\sigma}^2 &= \frac{1}{n}(y - X\widehat{\beta})^{\top}(y - X\widehat{\beta}) \end{cases}$$

#### where

 $\triangleright X^{\top} \text{ denotes the } \textit{transpose} \text{ of matrix } X$ 

 $\triangleright X^{\top}X$  denotes the *matrix product* of matrices  $X^{\top}$  and X

 $\rhd (X^\top X)^{-1} \text{ denotes the } \textit{matrix inverse} \text{ of matrix } (X^\top X)$ 

#### 2.3 Marginal Models



- Let's go back to the independence assumption
  - by the first five rows of the data are:

id	serBilir	age	drug
1	14.50	58.77	D-penicil
2	1.10	56.45	D-penicil
3	1.40	70.07	D-penicil
4	1.80	54.74	D-penicil
5	3.40	38.11	placebo

Each row represents a different patient, and patients are **independent** of each other



• When we have repeated measurements data, we have the form

id	serBilir	year	age	drug
1	14.50	0.00	58.77	D-penicil
1	21.30	0.53	58.77	D-penicil
2	1.10	0.00	56.45	D-penicil
2	0.80	0.50	56.45	D-penicil
2	1.00	1.00	56.45	D-penicil
2	1.90	2.10	56.45	D-penicil
2	2.60	4.90	56.45	D-penicil



Multiple rows per subject, rows belonging to the same subject are **correlated** 

- Note: Long vs Wide format
  - > wide format can only be used when all subjects are measured at the same time points

  - > (almost) all software accept repeated measurements data in long format



- How correlation affects modeling of the data?
- Say we are interested in the effect of time on serum bilirubin while also correcting for the age of the patients
  - > the corresponding regression equation is

$$\log(\mathtt{serBilir}_{ij}) = \beta_0 + \beta_1 \mathtt{Time}_{ij} + \beta_2 \mathtt{Age}_i + \varepsilon_{ij}$$

where

- \* serBilir $_{ij}$  denotes the level of serum bilirubin of patient i at time point  $Time_{ij}$
- \*  $\varepsilon_{ij}$  is the corresponding error term



- The fact that the responses of each patient are correlated translates to error terms that are correlated
  - ⊳ based on the data of the first two patients (see pp.47) we have

$\boxed{14.5}$		$\lceil 1 \rceil$	0.0	58.8			$oxedsymbol{arepsilon}_{11}$
21.3		1	0.5	58.8			$arepsilon_{12}$
1.1		1	0.0	56.5	$\beta_0$		$arepsilon_{21}$
0.8	=	1	0.5	56.5	$\beta_1$	+	$arepsilon_{22}$
1.0		1	1.0	56.5	$oxed{eta_2}$		$arepsilon_{23}$
1.9		1	1.9	56.5			$arepsilon_{24}$
2.6		$\lfloor 1$	2.6	56.5			$arepsilon_{25}$



ullet The direct approach to account for correlated data  $\Rightarrow$  multivariate regression

$$y_i = X_i \beta + \varepsilon_i, \quad \varepsilon_i \sim \mathcal{N}(0, V_i),$$

#### where

- $\triangleright y_i$  the vector of responses for the *i*-th subject
- $\triangleright X_i$  design matrix describing structural component
- $\triangleright V_i$  covariance matrix describing the correlation structure

The covariance matrix  $V_i$  explicitly accounts for the correlations

#### 2.4 Interpretation



- Interpretation of  $\beta$ 
  - $\triangleright \beta_j$  denotes the change in the average  $y_i$  when  $x_j$  is increased by one unit and all other covariates are fixed
- Example: In the AIDS dataset we are interested in the effect of treatment on the average longitudinal evolutions we fit a marginal model with
  - $\triangleright$  different average longitudinal evolutions per treatment group  $(X\beta)$  part
  - $\triangleright$  compound symmetry covariance matrix  $(V_i \text{ part})$

$$\left\{ \begin{array}{l} \sqrt{\mathtt{CD4}_{ij}} = \beta_0 + \beta_1 \mathtt{Time}_{ij} + \beta_2 \{\mathtt{ddI}_i \times \mathtt{Time}_{ij}\} + \varepsilon_{ij}, \\ \\ \varepsilon_i \sim \mathcal{N}(0, V_i) \end{array} \right.$$



	Value	Std.Err.	t-value	p-value
$\beta_0$	7.189	0.221	32.593	< 0.001
$\beta_1$	-0.156	0.017	-9.247	< 0.001
$\beta_2$	0.016	0.024	0.662	0.508

- $\triangleright$  Coefficient  $\beta_1:$  For patients in the ddC group, every month the average  $\sqrt{\text{CD4}}$  changes by -0.156
- $\triangleright$  Coefficient  $\beta_2$ :
  - \* Is the difference of the time effect between ddl and ddC
  - \* For patients in the ddl group, every month the average  $\sqrt{\text{CD4}}$  changes by  $\left(\text{-0.156} + \text{0.016}\right)$



ullet The estimated covariance matrix  $V_i$  is

$$> \operatorname{corr}(CD4_{t=0}, CD4_{t=2}) = \frac{\operatorname{cov}(CD4_{t=0}, CD4_{t=2})}{\sqrt{\operatorname{var}(CD4_{t=0})}\sqrt{\operatorname{var}(CD4_{t=2})}} = \frac{20.3}{24.15} = 0.84$$



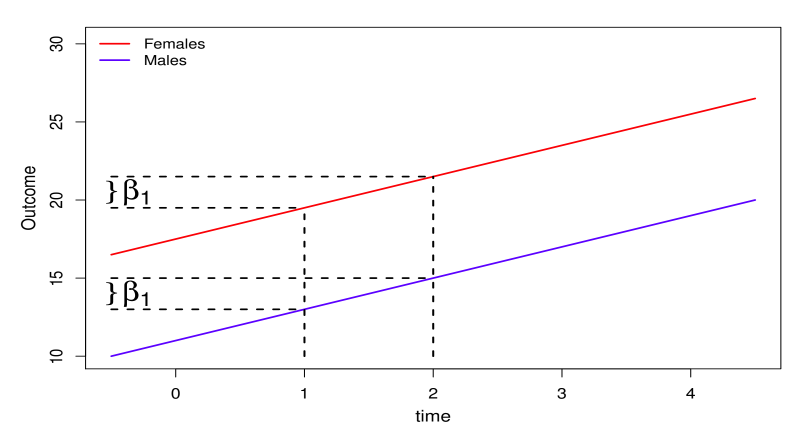
- Note: Interaction terms for longitudinal data
  - Consider the model

$$y_{ij} = \beta_0 + \beta_1 \mathtt{Time}_{ij} + \beta_2 \mathtt{Sex}_i + \varepsilon_{ij}, \quad \varepsilon_i \sim \mathcal{N}(0, V_i)$$

- \* we include the time effect and we also control for sex
- \* the model assumes that the effect of time is the same for the two sexes (parallel lines)



#### **Interaction Terms**



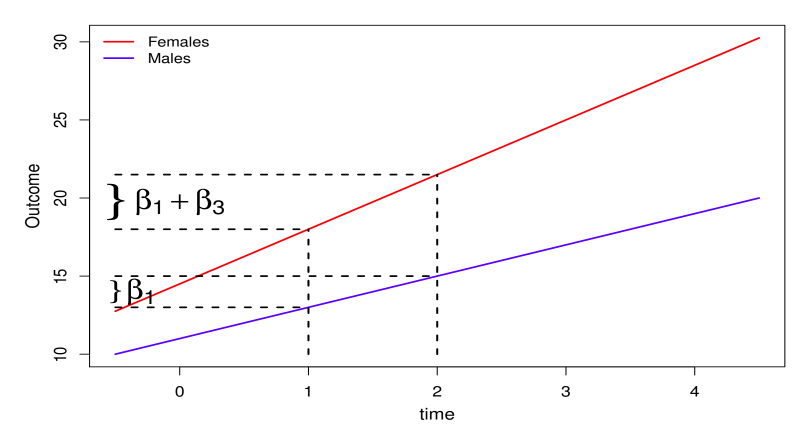


- Note: Interaction terms for longitudinal data
  - if we would like different longitudinal evolutions for the two sexes we need to include the interaction term

$$y_{ij} = \beta_0 + \beta_1 \mathtt{Time}_{ij} + \beta_2 \mathtt{Sex}_i + \beta_3 \{\mathtt{Sex}_i \times \mathtt{Time}_{ij}\} + \varepsilon_{ij}, \quad \varepsilon_i \sim \mathcal{N}(0, V_i)$$



#### **Interaction Terms**





- Communicating a model with complex terms: Due to the elaborate structure of repeated measurements data it is often required to include complex terms in a model

  - ▷ nonlinear terms (e.g., nonlinear evolutions in times modeled with polynomials or splines)
- ullet In such cases the regression coefficients eta we obtain in the output do not often have a straightforward interpretation



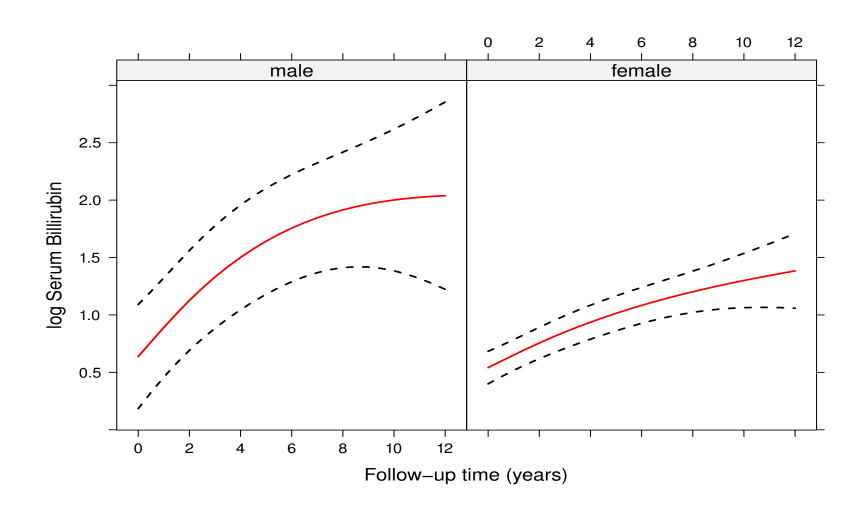
- To overcome this issue we can use **effect plots** 
  - ▶ this is a figure that depicts the average outcome along with 95% confidence intervals for specific combinations of the predictors levels
- Example: We have fitted the following model to the PBC dataset:

$$\begin{cases} \log(\text{serBilir}_{ij}) = \beta_0 + \beta_1 N(\text{Time}_{ij})_1 + \beta_2 N(\text{Time}_{ij})_2 + \beta_3 \text{Female}_i + \beta_4 \text{Age}_i + \\ \beta_5 \{\text{Female}_i \times N(\text{Time}_{ij})_1\} + \beta_6 \{\text{Female}_i \times N(\text{Time}_{ij})_2\} + \\ \beta_7 \{\text{Female}_i \times \text{Age}_i\} + \varepsilon_{ij} \end{cases}$$
 
$$\varepsilon_i \sim \mathcal{N}(0, V_i) \qquad V_i \text{ has a continuous AR1 structure}$$



- The terms  $N(\text{Time}_{ij})_1$  and  $N(\text{Time}_{ij})_2$  denote the basis for a natural spline with two degrees of freedom to model possible nonlinearities in the time effect
- In this model not all coefficients have a direct interpretation in isolation
- Hence to understand the model we depict
  - > how the average longitudinal profiles evolve over time time,
  - > separately for males and females, and
  - be for the average age of 49 years old (in the app different ages can be selected).
  - ▷ including also the corresponding 95% pointwise confidence intervals





#### 2.5 Estimation



- Estimation of model parameters
  - $\triangleright$  For known covariance matrix  $V_i$ , the regression coefficients are estimated using generalized least squares

$$\hat{\beta} = \left(\sum_{i=1}^{n} X_i^{\top} V_i^{-1} X_i\right)^{-1} \sum_{i=1}^{n} X_i^{\top} V_i^{-1} y_i$$

- $\triangleright$  Variance Components matrix  $V_i$ :
  - \* Maximum Likelihood (ML)
  - \* restricted maximum likelihood (REML)



- What's the difference between ML and REML?
  - > ML estimates of variances are known to be biased in small samples
  - b the simplest case: Sample variance

$$var(x) = \frac{1}{n-1} \sum_{i=1}^{n} (x_i - \bar{x})^2$$

\* to obtain an unbiased estimate we need to divide by n-1



#### The REML estimation is a generalization of this idea

- ullet It provides unbiased estimates of the parameters in the covariance matrix  $V_i$  in small samples
- Example: To illustrate the difference between REML and ML we consider fitting the same model for the AIDS dataset we have seen before but using only the first 50 rows



#### **▶ REML Estimation**

	t = 0	t=2	t = 6	t = 12	t = 18
t = 0	16.03	13.48	13.48	13.48	13.48
t = 2	13.48	16.03	13.48	13.48	13.48
t = 6	13.48	13.48	16.03	13.48	13.48
t = 12	13.48	13.48	13.48	16.03	13.48
t = 18	13.48	13.48	13.48	13.48	16.03



#### **▶ ML Estimation**

	t = 0	t=2	t = 6	t = 12	t = 18
t = 0	14.97	12.56	12.56	12.56	12.56
t = 2	12.56	14.97	12.56	12.56	12.56
t = 6	12.56	12.56	14.97	12.56	12.56
t = 12	12.56	12.56	12.56	14.97	12.56
t = 18	12.56	12.56	12.56	12.56	14.97

<sup>\*</sup> We observe some visible differences because of small n

<sup>\*</sup> In the full dataset the differences are negligible



- Features of REML estimation:
  - > Available in all software that fit marginal and mixed effects models
  - $\triangleright$  The way it works is by applying a transformation in the longitudinal outcome y based on the chosen structure of the design matrix X (i.e., which predictors you have included in the model)
  - ▶ Hence, we <u>cannot</u> compare the likelihoods of models fitted with REML and have different  $X\beta$  part

#### 2.6 Fitting Marginal Models in R



- R> Marginal models can be fitted using function gls() from the **nlme** package
- R> It has four basic arguments
  - ▷ model: a formula specifying the response vector and the covariates to include in the model

  - > correlation: an object describing the assumed correlation structure
  - ▷ weights: an object describing the assumed describing the within-group heteroscedasticity structure

# 2.6 Fitting Marginal Models in R (cont'd)



R> The data frame that contains all variables should be in the *long format* 

Subject	У	time	gender	age
1	5.1	0.0	male	45
1	6.3	1.1	male	45
2	5.9	0.1	female	38
2	6.9	0.9	female	38
2	7.1	1.2	female	38
2	7.3	1.5	female	38
ŧ	i	i	:	:

### 2.6 Fitting Marginal Models in R (cont'd)



R> Using formulas in R

R> Note: the intercept term is included by default

### 2.6 Fitting Marginal Models in R (cont'd)



R> The following code fits a marginal model for CD4 cell count with an AR1 correlation structure

#### 2.7 Covariance Matrix



• Reminder: What is a variance-covariance matrix?

b we have the dataset:

Subject	$Y_1$	$Y_2$	$Y_3$	$Y_4$
1	2.1	3.2	2.9	3.3
2	1.8	3.1	4.2	5.1
3	3.1	3.2	3.5	3.3
÷	÷	÷	÷	÷



• The variance-covariance matrix is the matrix whose element in the i, j-th position is the covariance between  $Y_i$  and  $Y_j$ , e.g.,

$$\begin{bmatrix} \mathsf{var}(Y_1) & \mathsf{cov}(Y_1,Y_2) & \mathsf{cov}(Y_1,Y_3) & \mathsf{cov}(Y_1,Y_4) \\ \mathsf{cov}(Y_2,Y_1) & \mathsf{var}(Y_2) & \mathsf{cov}(Y_2,Y_3) & \mathsf{cov}(Y_2,Y_4) \\ \mathsf{cov}(Y_3,Y_1) & \mathsf{cov}(Y_3,Y_2) & \mathsf{var}(Y_3) & \mathsf{cov}(Y_3,Y_4) \\ \mathsf{cov}(Y_4,Y_1) & \mathsf{cov}(Y_4,Y_2) & \mathsf{cov}(Y_4,Y_3) & \mathsf{var}(Y_4) \end{bmatrix}$$

#### Properties

> on the diagonal the variances, of diagonal covariances

$$\triangleright$$
 symmetric  $\Rightarrow$  cov $(Y_1, Y_2) = \text{cov}(Y_2, Y_1)$ 



- Variances, covariances and correlations
  - > variance measures how far a set of numbers is spread out (always positive)
  - covariance is a measure of how much two random variables change together (positive or negative)
  - $\triangleright$  correlation a measure of the linear correlation (dependence) between two variables (between -1 and 1; 0 no correlation)

$$\operatorname{corr}(Y_1,Y_2) = \frac{\operatorname{cov}(Y_1,Y_2)}{\sqrt{\operatorname{var}(Y_1)}\sqrt{\operatorname{var}(Y_2)}}$$



ullet Due to the fact that the magnitude of the covariance between  $Y_1$  and  $Y_2$  depends on their variability, we translate the covariance matrix to a correlation matrix



Coming back to our model

$$y_i = X_i \beta + \varepsilon_i, \quad \varepsilon_i \sim \mathcal{N}(0, V_i)$$

- ullet We need an appropriate choice for  $V_i$  in order to appropriately describe the correlations between the repeated measurements

> autoregressive process

- ▷ exponential spatial correlation
- ▷ ...



- Let's see some of those
  - □ General/Unstructured

$$\begin{bmatrix} \sigma_1^2 & \sigma_{12} & \sigma_{13} \\ \sigma_{12} & \sigma_2^2 & \sigma_{23} \\ \sigma_{13} & \sigma_{23} & \sigma_3^2 \end{bmatrix}$$

Diagonal

$$\begin{bmatrix} \sigma_1^2 & 0 & 0 \\ 0 & \sigma_2^2 & 0 \\ 0 & 0 & \sigma_3^2 \end{bmatrix} \qquad \text{or} \qquad \begin{bmatrix} \sigma^2 & 0 & 0 \\ 0 & \sigma^2 & 0 \\ 0 & 0 & \sigma^2 \end{bmatrix}$$



$$egin{bmatrix} \sigma^2 & 
ho\sigma^2 & 
ho^2\sigma^2 \ 
ho\sigma^2 & \sigma^2 & 
ho\sigma^2 \ 
ho^2\sigma^2 & 
ho\sigma^2 & \sigma^2 \end{bmatrix}$$

$$\begin{bmatrix} \sigma_1^2 & \rho_1 \sigma_1 \sigma_2 & \rho_2 \sigma_1 \sigma_3 \\ \rho_1 \sigma_1 \sigma_2 & \sigma_2^2 & \rho_1 \sigma_2 \sigma_3 \\ \rho_2 \sigma_1 \sigma_3 & \rho_1 \sigma_2 \sigma_3 & \sigma_3^2 \end{bmatrix} \quad \text{or} \quad \begin{bmatrix} \sigma^2 & \sigma_{12} & \sigma_{13} \\ \sigma_{12} & \sigma^2 & \sigma_{12} \\ \sigma_{13} & \sigma_{12} & \sigma^2 \end{bmatrix}$$



- The aforementioned structure for the covariance matrix are applicable in cases we have discrete and equally spaced time points
- For continuous time and unbalanced data, alternative options are:

  - ▷ exponential serial correlation



- These serial correlation structures are defined using the semi-variogram
  - by which we are not going to cover here because it is a bit technical (more info in any standard text for mixed models / longitudinal data analysis)
- the basic assumption is that correlations decay with the time lag  $|t_i t_j| \Rightarrow$  measurements at closer time points are more strongly correlated than measurements at more distant time points
  - > each of these structure how one parameter that controls how correlation decay in time



- Notes: on building covariance matrices
  - ► variance function: in some cases, and especially for longitudinal data quite often, it may not be reasonable to assume that the variance of the outcome remains constant in time
    - \* we have seen versions of heteroscedastic covariance matrices, but these are only applicable when we have balanced data and few time points
    - \* for unbalanced designs we can specify other variance functions, e.g., that variances increase linearly or exponentially with time
  - *correlation at the same point:* is it **always** reasonable that the correlation of the outcome at the same point is set to 1?



• Let's try the app...

#### 2.8 Model Building



- We have seen that marginal models consist of two parts:
  - $\triangleright$  Mean part  $X\beta$ : that describes how covariates we have put in the model explain the average of the repeated measurements
  - $\triangleright$  Covariance part  $V_i$ : assumed covariance structure between the repeated measurements

• In the majority of the cases scientific interest focuses on the mean part

However, to obtain valid and efficient inferences for this part, the covariance part need to be adequately specified

### 2.8 Model Building (cont'd)



- Hence, the general strategy for building models for repeated measurements data proceeds as follows:
  - 1. Put all the covariates of interest in the mean part, considering possible interactions between them do NOT remove the ones that are not significant
  - 2. Then select an appropriate covariance matrix  $V_i$  that adequately describes the correlations in the repeated measurements
    - \* in this step you should be a bit conservative, i.e., do not favor a simpler covariance matrix if the p-value is just non-significant
  - 3. Finally, return to the mean part and exclude non significant covariates
    - \* first start by testing the interaction terms

### 2.8 Model Building (cont'd)



- How many covariates we can put in the mean part?
- It depends on how strong are the correlation between the repeated measurements
  - $\triangleright$  weak correlations  $\Rightarrow N/10$  (N total number of measurements)
  - $\triangleright$  strong correlations  $\Rightarrow n/10$  (n number of subjects)

#### 2.9 Hypothesis Testing



- Having fitted a marginal model using maximum likelihood we can use standard inferential tools for performing hypothesis testing

  - ▷ Likelihood ratio tests

- Following the model building strategy described above, we will

  - > and following focus on hypothesis testing for the mean part of the model



- Hypothesis testing for  $V_i$ : Assuming the same mean structure we can fit a series of model and choose the that best describes the covariances
- In general, we distinguish between two cases
  - > comparing two models with *nested* covariance matrices
  - > comparing two models with *non-nested* covariance matrices
- Note: Model A is nested in Model B, when Model A is a special case of Model B, i.e.,
  - by setting some of the parameters of Model B at some specific value we then obtain Model A



• For **nested** models the preferable test for selecting  $V_i$  is the likelihood ratio test (LRT):

$$LRT = -2 \times \{\ell(\hat{\theta}_0) - \ell(\hat{\theta}_a)\} \sim \chi_p^2$$

#### where

- $\triangleright \ell(\hat{\theta}_0)$  the value of the log-likelihood function under the null hypothesis, i.e., the special case model
- $\triangleright \ell(\hat{\theta}_1)$  the value of the log-likelihood function under the alternative hypothesis, i.e., the general model
- $\triangleright p$  denotes the number of parameters being tested
- Note: Provided that the mean structure in the two models is the same, we can either compare the REML of ML likelihoods of the models (preferable is REML)



- Example: In the model we fitted for the AIDS dataset (see pp.52) we had assumed a compound symmetry covariance matrix we would like to see if this option was sufficient
  - ▷ we will compare the compound symmetry model:

$$H_0: V_i = \begin{bmatrix} t = 0 & t = 2 & t = 6 & t = 12 & t = 18 \\ \sigma^2 & \tilde{\sigma} & \tilde{\sigma} & \tilde{\sigma} & \tilde{\sigma} \\ & \sigma^2 & \tilde{\sigma} & \tilde{\sigma} & \tilde{\sigma} \\ & & \sigma^2 & \tilde{\sigma} & \tilde{\sigma} \\ & & & \sigma^2 & \tilde{\sigma} & \tilde{\sigma} \\ & & & & \sigma^2 & \tilde{\sigma} \\ & & & & & \sigma^2 \end{bmatrix}$$



> versus the unstructured model

$$H_a: V_i = \begin{bmatrix} t = 0 & t = 2 & t = 6 & t = 12 & t = 18 \\ \sigma_1^2 & \sigma_{12} & \sigma_{13} & \sigma_{14} & \sigma_{15} \\ & \sigma_2^2 & \sigma_{23} & \sigma_{24} & \sigma_{25} \\ & & \sigma_3^2 & \sigma_{34} & \sigma_{35} \\ & & & \sigma_4^2 & \sigma_{45} \\ & & & & \sigma_5^2 \end{bmatrix}$$



We can rewrite the two hypothesis as

$$H_0: \begin{cases} \sigma_1^2 = \sigma_2^2 = \dots = \sigma_5^2 = \sigma^2 \\ \sigma_{12} = \sigma_{13} = \dots = \sigma_{45} = \tilde{\sigma} \end{cases}$$

 $H_a$ : at least one variance of covariance is not equal to the others

• The likelihood ratio test gives:

	df	logLik	LRT	p-value
Comp Symm	5	-3586.91		NA
General	18	-3547.72	78.39	< 0.0001



- When we have **non-nested** models we **cannot** use standard tests anymore
- As an alternative for this case we use information criteria the two standard ones are:

$$\begin{aligned} \mathsf{AIC} &= -2\ell(\hat{\theta}) + 2n_{par} \\ \mathsf{BIC} &= -2\ell(\hat{\theta}) + n_{par}\log(n) \end{aligned}$$

#### where

- $riangleright \ell(\hat{ heta})$  is the value of the log-likelihood function
- $\triangleright n_{par}$  the number of parameters in the model
- $\triangleright n$  the number of subjects (independent units)



When we compare two **non-nested** models we choose the model that has the **lowest** AIC/BIC value

• Example: For the Prothrobin data we compare the exponential and Gaussian serial correlation structures – the model are:

$$\begin{cases} \textit{$M_1$: $\operatorname{pro}_{ij} = \beta_0 + \beta_1 \operatorname{Time}_{ij} + \beta_2 \{\operatorname{predn}_i \times \operatorname{Time}_{ij}\} + \varepsilon_{ij}, & \varepsilon_i \sim \mathcal{N}(0, V_i^{Exp}) \\ \\ \textit{$M_2$: $\operatorname{pro}_{ij} = \beta_0 + \beta_1 \operatorname{Time}_{ij} + \beta_2 \{\operatorname{predn}_i \times \operatorname{Time}_{ij}\} + \varepsilon_{ij}, & \varepsilon_i \sim \mathcal{N}(0, V_i^{Gauss}) \end{cases}$$



• The AIC and BIC values for the two models are:

	df	logLik	AIC	BIC
Exp	5	-13468.84	26947.67	26977.65
Gauss	5	-13750.88	27511.76	27541.73

▶ Both AIC and BIC suggest that the model with the exponential correlation structure is better



- The models we have assumed for the Prothrobin data assumed constant variance in time as we have mentioned (see pp. 82), this assumption is not often justified for longitudinal data
- We extend models  $M_1$  and  $M_2$  by assuming that the variances are an exponential function of time, i.e.,

$$\mathsf{var}(arepsilon_{ij}) = \sigma^2 \exp ig( \delta \mathtt{Time}_{ij} ig)$$

where

 $hd \delta$  is a parameter that controls how fast the variance changes with time

- \* if  $\delta < 0$ , the variance decreases with time
- \* if  $\delta = 0$ , the variance remains constant
- \* if  $\delta > 0$ , the variance increases with time



ullet This means that models  $M_1$  and  $M_2$  are nested within their heteroscedastic cousins, i.e.,

 $H_0: \delta = 0$  homoscedastic model

 $H_a: \delta \neq 0$  heteroscedastic model

• This implies that we can perform a likelihood ratio test

	df	logLik	AIC	BIC	LRT	p-value
Exp - homoscedastic	5	-13468.84	26947.67	26977.65		NA
Exp - heteroscedastic	6	-13459.99	26931.97	26967.94	17.70	< 0.0001
Gauss - homoscedastic	5	-13750.88	27511.76	27541.73		NA
Gauss - heteroscedastic	6	-13748.10	27508.21	27544.18	17.70	0.0185



- ullet Notes: Hypothesis testing for the covariance matrix  $V_i$ 
  - > The unstructured covariance matrix is the most general matrix we can assume:
    - \* all other covariance matrices are a special case of the unstructured matrix
    - \* **but** realistically it can only be fitted when we have balanced data and relatively few time points
  - ▷ The AIC and BIC do not always select the same model when they disagree
    - \* AIC typically selects the more elaborate model, whereas
    - \* BIC the more parsimonious model



- Hypothesis testing for the regression coefficients  $\beta$ : We assume that first a suitable choice for the covariance matrix has been made
- In the majority of the cases we compare nested models, and hence the standard test can be used
- We distinguish between two cases
  - > tests for individual coefficients
  - bets for groups of coefficients



- Tests for individual coefficients are based on the Wald-type statistic but assume the t distribution for calculating p-values
  - betto the betto t

$$H_0: \beta = 0$$

$$H_a:\beta\neq 0$$

 $\triangleright$  and we use the t test statistic

$$\frac{\hat{eta}}{s.e.(\hat{eta})} \sim t_{df}$$

where  $\hat{\beta}$  is the MLE,  $s.e.(\hat{\beta})$  is the standard error of the MLE, and df are specified according to the number of subjects and number of repeated measurements per subject



- Tests for groups of coefficients are based on the F-test
  - b the set of hypotheses is:

$$H_0: L\beta = 0$$

$$H_a: L\beta \neq 0$$

where L is the contrasts matrix

 $\triangleright$  the F test statistic is

$$\frac{\hat{\beta}^{\top}L^{\top}\bigg\{L\bigg(\sum\limits_{i=1}^{n}X_{i}^{\top}V_{i}^{-1}X_{i}\bigg)^{-1}L^{\top}\bigg\}^{-1}L\hat{\beta}}{\operatorname{rank}(L)}\sim F_{d\!f_{1},d\!f_{2}}$$



- Tests for groups of coefficients are based on the F-test
  - $\triangleright$  The numerator degrees of freedom are always equal to the rank of the contrast matrix L
  - Denominator degrees of freedom need to be estimated from the data:
    - \* Containment method
    - \* Satterthwaite approximation
    - \* Kenward and Roger approximation



• Example: We have fitted the following model to the PBC dataset:

$$\begin{cases} \log(\texttt{serBilir}_{ij}) = \beta_0 + \beta_1 \texttt{Time}_{ij} + \beta_2 \texttt{Female}_i + \beta_3 \texttt{Age}_i + \\ \beta_4 \{\texttt{D-penicil}_i \times \texttt{Time}_{ij}\} + \beta_5 \{\texttt{Female}_i \times \texttt{Time}_{ij}\} + \varepsilon_{ij} \end{cases}$$
 
$$\varepsilon_i \sim \mathcal{N}(0, V_i)$$

where  $V_i$  has a continuous AR1 structure

- We are interested in
  - b the effect of Age, and
  - b the overall effect of Sex



• For the effect of Age we set the hypotheses:

$$H_0: \beta_3 = 0$$

$$H_a: \beta_3 \neq 0$$

• The output of the model gives: ...



	Value	Std.Err.	t-value	p-value
$\beta_0$	0.940	0.395	2.382	0.017
$\beta_1$	0.154	0.034	4.546	< 0.001
$\beta_2$	-0.281	0.218	-1.291	0.197
$\beta_3$	-0.002	0.006	-0.361	0.718
$\beta_4$	-0.014	0.020	-0.670	0.503
$\beta_5$	-0.064	0.034	-1.862	0.063

• Hence, a non-significant Age effect

 $\triangleright$  the t-value in the output is the estimated coefficient divided by its standard error



• For the overall effect of Sex we set the hypotheses:

$$H_0: \beta_2 = \beta_5 = 0$$

 $H_a$  : either  $eta_2$  or  $eta_5$  are not equal to 0

- We cannot obtain the p-value for this test directly from the output
- ullet We have six parameters, the contrast matrix L is

$$L = \begin{bmatrix} \beta_0 & \beta_1 & \beta_2 & \beta_3 & \beta_4 & \beta_5 \\ 0 & 0 & 1 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 & 1 \end{bmatrix}$$



• We obtain

$$\frac{F\text{-value}}{4.458} \ \frac{df_1}{2} \ \frac{df_2}{1939} \ \frac{p\text{-value}}{0.0117}$$

- Hence, a significant overall sex effect
- We could also test the same hypotheses using a likelihood ratio test
  - ▷ in this case we compare the models under the null and alternative hypothesis



• The two models are:

$$H_0: \log(\mathtt{serBilir}_{ij}) = \beta_0 + \beta_1 \mathtt{Time}_{ij} + \beta_3 \mathtt{Age}_i + \beta_4 \{\mathtt{D-penicil}_i \times \mathtt{Time}_{ij}\} + \varepsilon_{ij}$$

$$\begin{split} H_a: \log(\texttt{serBilir}_{ij}) &= \beta_0 + \beta_1 \texttt{Time}_{ij} + \beta_2 \texttt{Female}_i + \beta_3 \texttt{Age}_i + \\ & \beta_4 \{\texttt{D-penicil}_i \times \texttt{Time}_{ij}\} + \beta_5 \{\texttt{Female}_i \times \texttt{Time}_{ij}\} + \varepsilon_{ij} \end{split}$$

 $\triangleright$  for both models  $V_i$  has a continuous AR1 structure

• If we compare the two models we again end up in the same hypotheses:

$$H_0: \beta_2 = \beta_5 = 0$$

 $H_a$ : either  $eta_2$  or  $eta_5$  are not equal to 0



• The likelihood ratio test gives

	df	logLik	AIC	BIC	LRT	p-value
without Sex	6	-1618.23	3248.46	3281.90		NA
with Sex	8	-1613.76	3243.52	3288.10	8.94	0.0114

• Hence, again the same conclusion, i.e., a significant overall sex effect



- Notes: Hypothesis testing for the regression coefficients  $\beta$ 
  - $\triangleright$  The likelihood ratio test, and the classical univariate and multivariate Wald tests (i.e., using the  $\chi^2$  distribution instead of the t or F distributions) are 'liberal' \* they give smaller p-values than the ones they should give, especially in small
    - $^{f k}$  they give smaller p-values than the ones they should give, especially in small samples
  - ▶ Important: The likelihood ratio test for comparing models with different  $X\beta$  parts is only valid when the models have been fitted using maximum likelihood and <u>not</u> REML (see also pp. 64–68)

#### 2.10 Confidence Intervals



• Confidence intervals for model parameters are obtained from the approximate distribution of the maximum likelihood estimates (MLEs)

$$\hat{\beta} \sim \mathcal{N}(\beta^*, \mathsf{var}(\hat{\beta}))$$

where

 $\triangleright \hat{\beta}$  are the MLEs

 $\triangleright \beta^*$  the true parameter values

$$> \mathrm{var}(\hat{\beta}) = \left(\sum_{i=1}^n X_i^\top V_i^{-1} X_i\right)^{-1} \text{ is the covariance matrix of the MLEs}$$

## 2.10 Confidence Intervals (cont'd)



• For example, for the k-th regression coefficient  $\beta_k$ , the 95% Cl is

$$\hat{\beta} \pm 1.96 \times \text{s.e.}(\hat{\beta})$$

ullet To obtain confidence intervals for the whole mean evolution we need to multiply with a corresponding design matrix X (see pp. 43–44), i.e.,

$$X \hat{\beta} \pm 1.96 \times \sqrt{\mathsf{diag} \big\{ X \mathsf{var}(\hat{\beta}) X^\top \big\}}$$

b this type of confidence intervals have been used in the effects plots we have seen earlier (see pp. 59–62)

#### 2.11 Residuals



All statistical models are based on assumptions

• Hence, to extract meaningful conclusions we need to check whether these assumptions are (crudely) violated



• The marginal model for continuous data makes analogous assumptions as the linear regression model

$$y_i = X_i \beta + \varepsilon_i, \quad \varepsilon_i \sim \mathcal{N}(0, V_i)$$

#### namely

- $\triangleright$  the error terms  $\varepsilon_i$  follow the normal distribution  $\mathcal{N}(0,V_i)$
- $\triangleright$  the error terms are independent from the covariates X
- $\triangleright$  the covariates act linearly on the average outcome (here 'linearly' means with respect to the parameters  $\beta$ )



- ullet To validate these assumptions we need an estimate of the error terms  $arepsilon_{ij}$
- Based on the fitted model we obtain the estimate

$$r_{ij} = y_{ij} - x_{ij}^{\mathsf{T}} \hat{\beta}$$

- $\triangleright \hat{\beta}$  are the (restricted) maximum likelihood estimates
- $\triangleright$  the  $r_{ij}$  are called *residuals*

When the model is correctly specified, we expect these residuals to have a  $\mathcal{N}(0, V_i)$  distribution



- Hence, we expect these residuals to be correlated and possibly also heteroscedastic
  - ▷ 'heteroscedastic' means that they exhibit non-constant variance
- This feature complicates matters because it is not easy to assess if the residuals exhibit the assumed properties
- ullet To overcome this problem we need to transform  $r_{ij}$  to a scale that has easier to check properties



• To achieve this we multiply the residual with the inverse Choleski factor

$$r_i^{norm} = \widehat{H}_i^{-1} r_i = \widehat{H}_i^{-1} (y_i - X_i \hat{\beta})$$

#### where

- $\triangleright \widehat{H}_i$  is an upper-triangular matrix with the property  $\widehat{H}_i^{\top}\widehat{H}_i = \widehat{V}_i$ , with  $\widehat{V}_i$  denoting the estimated covariance matrix
- $ightharpoonup rate} rate rate rate rate and rate rate rate and when the covariance matrix is correctly specified, they should be approximately distributed as <math>\mathcal{N}(0,1)$  random variables



• When we have assumed a homoscedastic covariance matrix (i.e., variance remains constant), another transformation that it is often used is

$$r_i^{Pears} = \hat{\sigma}^{-1} r_i = \sigma^{-1} (y_i - X_i \hat{\beta})$$

#### where

- $\triangleright \hat{\sigma}$  denotes the estimated standard deviation of the error term, i.e.,  $V_i$  has the structure  $\sigma^2 R_i$ , with  $R_i$  denoting a correlation matrix
- ightharpoonup residuals and when the covariance matrix is correctly specified, they should be approximately distributed as  $\mathcal{N}(0,R_i)$  random variables



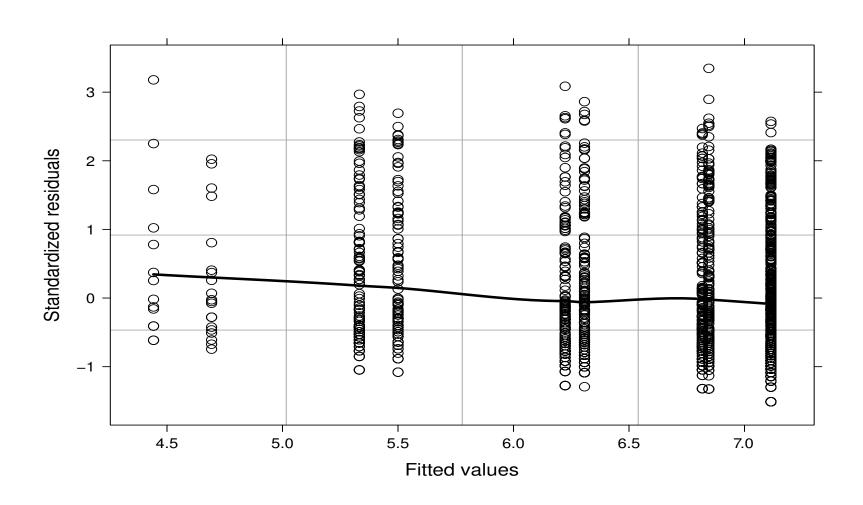
• Example: We evaluate the assumptions behind the following model fitted to the AIDS dataset:

$$\begin{cases} \sqrt{\text{CD4}_{ij}} = \beta_0 + \beta_1 \text{Time}_{ij} + \beta_2 \{\text{ddI}_i \times \text{Time}_{ij}\} + \varepsilon_{ij}, \\ \\ \varepsilon_i \sim \mathcal{N}(0, V_i), \quad V_i \text{ is unstructured} \end{cases}$$

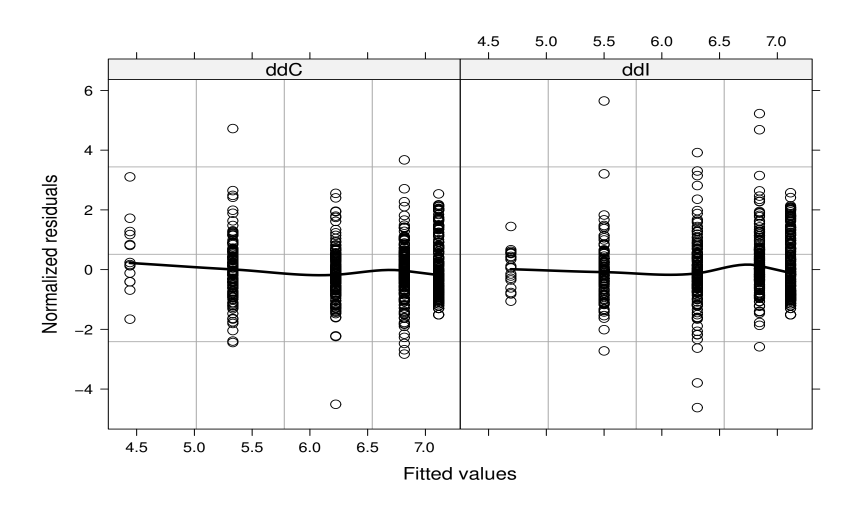
#### by plotting

- b the standardized residuals versus fitted values
- > the normalized residuals versus fitted values per treatment group

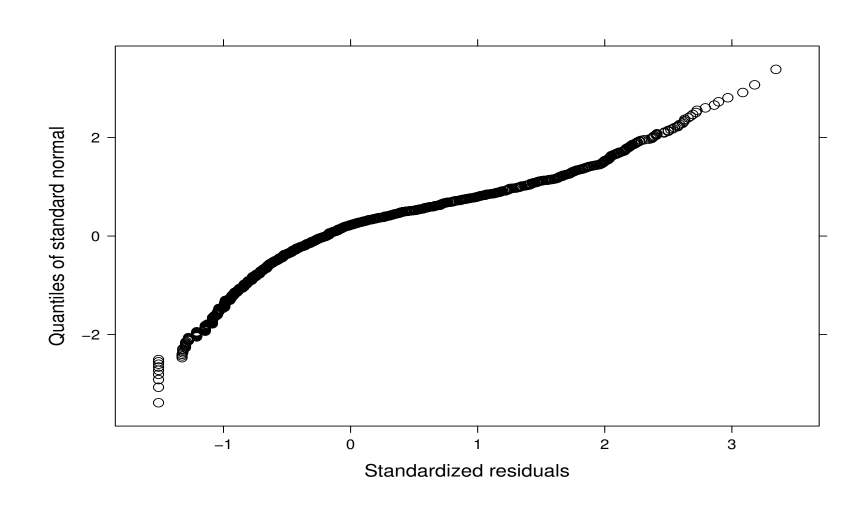














#### Observations

- behavior with more positive residuals in the range of low fitted values
- b the QQ-plot is not perfect, but does not show a big discrepancy from normality



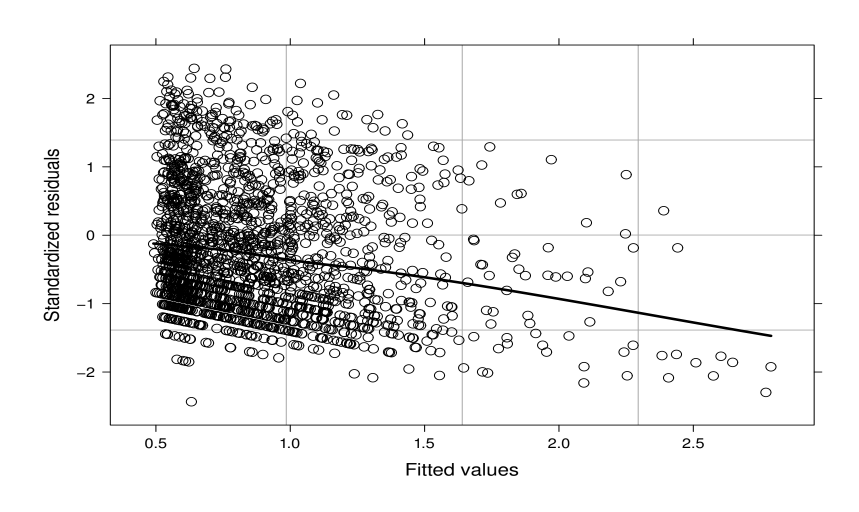
• Example: We continue by evaluating the assumptions of the model we have fitted to the PBC dataset:

$$\begin{cases} \log(\text{serBilir}_{ij}) = \beta_0 + \beta_1 \text{Time}_{ij} + \beta_2 \text{Female}_i + \beta_3 \text{Age}_i + \\ \beta_4 \{\text{D-penicil}_i \times \text{Time}_{ij}\} + \beta_5 \{\text{Female}_i \times \text{Time}_{ij}\} + \varepsilon_{ij} \end{cases}$$
 
$$\varepsilon_i \sim \mathcal{N}(0, V_i) \qquad V_i \text{ has a continuous AR1 structure}$$

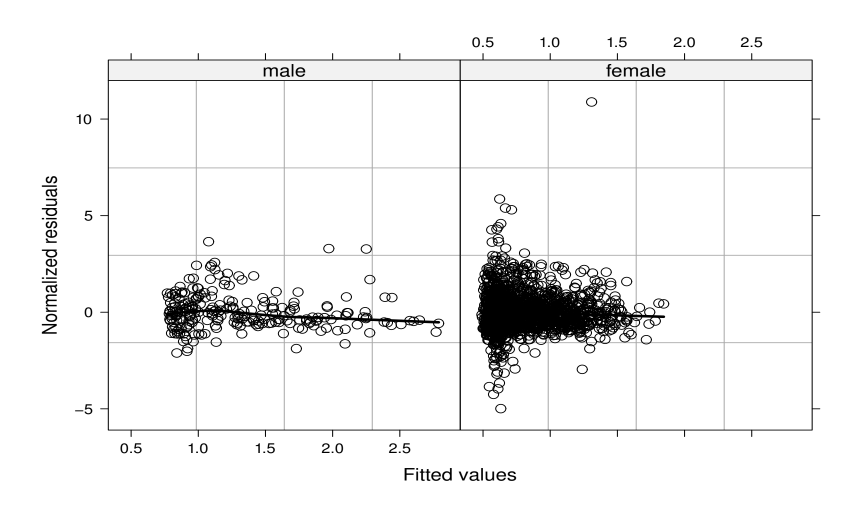
by plotting again

- > the standardized residuals versus fitted values
- > the normalized residuals versus fitted values per gender
- ▷ QQ-plot of the standardized residuals

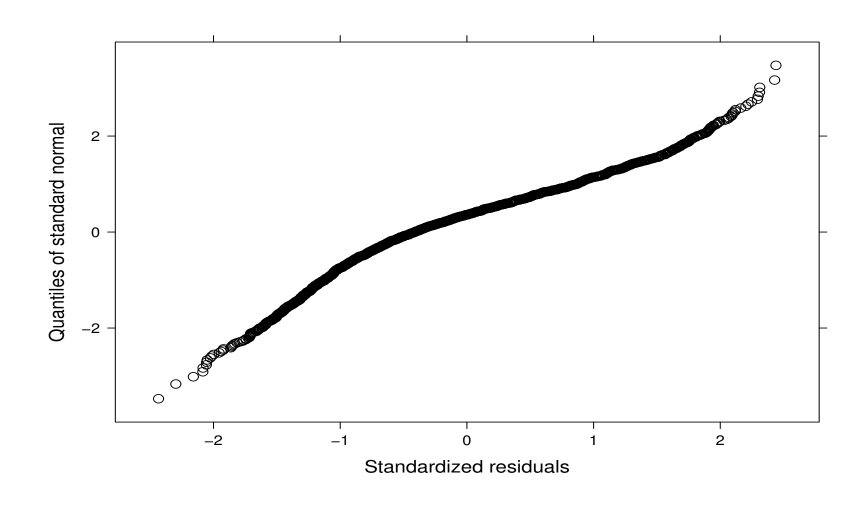














#### Observations

- by the plot of the standardized residuals versus fitted values shows a clear systematic trend with more negative residuals in the range of high fitted values
- b the plot of normalized residuals versus fitted values shows an outlying observation
   for female and some slight heteroscedasticity (higher spread of residuals for low
   fitted values than for high)
- by the QQ-plot suggests a good fit of the normal distribution

#### 2.12 Review of Key Points



- Methods for analyzing grouped/correlated data
  - $\triangleright$  naive approached working on parts or summaries of the data  $\Rightarrow$  loss of information
  - ▷ marginal models ⇒ extension of simple linear regression to the context of correlated data
- Marginal models: Features
  - $\triangleright$  error terms are assumed correlated  $\Rightarrow$  we need to make an appropriate assumption
  - ▶ mean structure is build as in standard regression models however, need to account for potential nonlinear effects of time and/or interaction terms
  - ▶ model building: we start from a 'fully' specified mean structure, we select an
     appropriate covariance structure, and then the return to make inference for the
     mean

## 2.12 Review of Key Points (cont'd)



#### Hypothesis testing

- be for the covariance structure and for nested models likelihood ratio tests are most often used, for non-nested models AIC/BIC
- $\triangleright$  for the mean structure t and F tests with appropriate degrees of freedom

#### Residuals

- > standard residuals plots are used to check the model assumptions
- > standardized and normalized residuals