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# Chapter 1

# Using DFBETAs from LME Models to Assess Agreement

## 1.1 Fitting Models with the LME4 R package

Maximum likelihood or restricted maximum likelihood (REML) estimates of the parameters in linear mixed-effects models can be determined using the lmer function in the lme4 package for R. As for most model-fitting functions in R, the model is described in an lmer call by a formula, in this case including both fixed- and random-effects terms. The formula and data together determine a numerical representation of the model from which the profiled deviance or the profiled REML criterion can be evaluated as a function of some of the model parameters. The appropriate criterion is optimized, using one of the constrained optimization functions in R, to provide the parameter estimates. We describe the structure of the model, the steps in evaluating the profiled deviance or REML criterion, and the structure of classes or types that represents such a model. Sufficient detail is included to allow specialization of these structures by users who wish to write functions to fit specialized linear mixed models, such as models incorporating pedigrees or smoothing splines, that are not easily expressible in the formula language used by lmer.

y : Response variable

method: Method of Measurement

subject : Subject

MCSdata

```
library(lme4)

MCS.lme4 <- lmer(y ~ method-1 + (1|subject) , data=MCSdata)</pre>
```

## 1.2 influence.ME

influence.ME allows you to compute measures of influential data for mixed effects models generated by lme4.

influence.ME provides a collection of tools for detecting influential cases in generalized mixed effects models. It analyses models that were estimated using lme4. The basic rationale behind identifying influential data is that when iteratively single units are omitted from the data, models based on these data should not produce substantially different estimates.

To standardize the assessment of how influential a (single group of) observation(s) is, several measures of influence are common practice, such as DFBETAS and Cook's Distance. In addition, we provide a measure of percentage change of the fixed point estimates and a simple procedure to detect changing levels of significance.

#### 1.2.1 Influence

influence() is the workhorse function of the influence.ME package. Based on a priorly estimated mixed effects regression model (estimated using lme4), the influence()

function iteratively modifies the mixed effects model to neutralize the effect a grouped set of data has on the parameters, and which returns returns the fixed parameters of these iteratively modified models. These are used to compute measures of influential data.

## 1.3 Cooks's Distance

Cook's Distance is a measure indicating to what extent model parameters are influenced by (a set of) influential data on which the model is based. This function computes the Cook's distance based on the information returned by the estex() function.

#### 1.4 DFBETAs

DFBETAS (standardized difference of the beta) is a measure that standardizes the absolute difference in parameter estimates between a (mixed effects) regression model based on a full set of data, and a model from which a (potentially influential) subset of data is removed. A value for DFBETAS is calculated for each parameter in the model separately. This function computes the DFBETAS based on the information returned by the estex() function.

## 1.5 Application of DFBETAs in MCS Analysis

When in an MCS study. DFBetas can be used as a proxy measurement, allowing simple techniques to be used for assessing agreement.

# Chapter 2

## Preliminaries: Useful R commands

#### 2.1 coef.lme

The command coef.lme extract lmes coefficients

The estimated coefficients at level i are obtained by adding together the fixed effects estimates and the corresponding random effects estimates at grouping levels less or equal to i. The resulting estimates are returned as a data frame, with rows corresponding to groups and columns to coefficients. Optionally, the returned data frame may be augmented with covariates summarized over groups.

```
> JS.roy1 %>% coef %>% head(10)
     methodJ
               methodS
74
  84.31724 91.08404
   91.54994
            97.05548
36
3
    81.16581
            96.48653
   92.09493 90.89073
62
   88.41411 103.38802
31
   95.56881 104.70922
42
11 103.46092 111.33625
```

```
41 94.97700 108.08384
55 79.44762 110.02055
17 101.53470 112.46964
```

Suppose we refit the model, with a reduced data set ( for example, with subject 68 missing)

```
x=68
datred <- dat[dat$subject !=x,]

# datred is other 84 cases (504 observations)

JS.roy1.red = lme(BP ~ method-1,
    data = datred,
    random = list(subject=pdSymm(~ method-1)),
    weights=varIdent(form=~1|method),
    correlation = corSymm(form=~1 | subject/repl),
    method="ML")</pre>
```

```
> JS.roy1.red %>% coef %>% dim
[1] 84  2
>
> JS.roy1.red %>% coef %>% head(10)
    methodJ methodS
74 84.31425 90.89550
36 91.54526 96.86813
```

- 3 81.17663 96.36665
- 62 92.07933 90.64629
- 31 88.42473 103.27726
- 42 95.57027 104.56000
- 11 103.46071 111.18930
- 41 94.98491 107.96791
- 55 79.48324 110.02963
- 17 101.53939 112.34591

# Chapter 3

## LME commands

vcov returns the variance-covariance matrix of the main parameters of a fitted model object.

```
> vcov(JS.roy1)

methodJ methodS

methodJ 11.01701 9.30105

methodS 9.30105 11.75301
```

getVarCov extract the variance-covariance matrix from a fitted model, such as a mixed-effects model.

# Chapter 4

 ${\bf Model\ Diagnostics\ for\ MCS}$ 

#### Abstract

Model diagnostic techniques, well established for classical models, have since been adapted for use with linear mixed effects models. However, diagnostic techniques for LME models are inevitably more difficult to implement, due to the increased complexity.

- ? describes the examination of model-data agreement as comprising several elements;
  - residual analysis,
  - goodness of fit,
  - collinearity diagnostics
  - influence analysis.

This chapter is comprised of two sections:

- 1. Residual Diagnostics
- 2. Influence Diagnostics

## 4.1 Introduction

In statistical modelling, the process of model validation is a critical step, but also a step that is too often overlooked. A very simple procedure is to examine well known metrics, such as the AIC and  $R^2$  measures. However, using a small handful of simple measures and methods is insufficient to properly assess the quality of a fitted model. To do so properly, a full and comprehensive analysis that tests of all of the assumptions, as far as possible, must be carried out.

A statistical model, whether of the fixed-effects or mixed-effects variety, represents how you think your data were generated. Following model specification and estimation, it is of interest to explore the model-data agreement by raising pertinent questions. Further to the analysis of residuals, ? recommends the examination of the following questions.

- Does the model-data agreement support the model assumptions?
- Should model components be refined, and if so, which components? For example, should certain explanatory variables be added or removed, and is the covariance of the observations properly specified?
- Are the results sensitive to model and/or data? Are individual data points or groups of cases particularly influential on the analysis?

## 4.2 Residual diagnostics

#### 4.2.1 Introduction to Residual Analysis

Residual analysis is a widely used model validation technique. A residual is simply the difference between an observed value and the corresponding fitted value, as predicted by the model. The rationale is that, if the model is properly fitted to the model, then the residuals would approximate the random errors that one should expect; if the residuals behave randomly, with no discernible trend. If some sort of non-random trend is evident in the model, then the model can be considered to be poorly fitted.

For classical linear models, residual diagnostics are typically implemented as a plot of the observed residuals and the predicted values. A visual inspection for the presence of trends inform the analyst on the validity of distributional assumptions, and to detect outliers and influential observations. Statistical software environments, such as the R programming language, provides a suite of tests and graphical procedures for appraising a fitted linear model, with several of these procedures analysing the model residuals.

However, for LME models the matter of residual is more complex, both from a theoretical point of view and from the practical matter of implementing a comprehensive analysis using statistical software. As the LME model can be tailored to the needs of the particular research question, the rationale behind the model appraisal must follow accordingly.

## 4.2.2 Residuals in the Blood Data Example

The fitted model used in the Blood data example, JS.roy1, was fitted using the lme() function from the nlme package, and as such, is stored as an lme object. The residual functions extracts residuals of a fitted LME model, depending on the type of residual required.

For an lime object, the residuals at level i are obtained by subtracting the fitted levels at that level from the response vector (and dividing by the estimated within-

group standard error, if type="pearson"). The Pearson residual is the raw residual divided by the square root of the variance function (here, the Within-group standard error for both methods, 6.11 and 9.11 respectively). The fitted values at level i are obtained by adding together the population fitted values (based only on the fixed effects estimates) and the estimated contributions of the random effects to the fitted values at grouping levels less or equal to i.

"response": the raw residuals (observed - fitted) are used. This is the default option.

"pearson": the standardized residuals (raw residuals divided by the corresponding standard errors) are used;

"normalized": the normalized residuals (standardized residuals pre-multiplied by the inverse square-root factor of the estimated error correlation matrix) are used.

```
response pearson normalized

1 -4.65805902 -0.761587227 -0.7615872269

2 -0.88701342 -0.145025661 0.0776238081

3 -5.16580898 -0.844603753 -0.8446037530

4 2.29041830 0.374480726 0.6450898404

5 7.87508366 1.287567009 1.2875670086

6 -6.57048659 -1.074266908 -1.5090772378
```

For the J observations, the variance is 6.116252 whereas for the S observations, the denominator is 9.118144. (with the expected ratio of 1.490806)

```
> pearson %>%
   as.numeric %>%
   matrix(nrow=85) %>%
   round(4)
               [,1]
                       [,2]
                              [,3]
                                     [,4]
                                             [,5]
                                                    [,6]
        [1,] -0.7616 0.2194 0.3829 -0.2983 0.3597 -0.0790
        [2,] -0.1450  0.1820 -0.1450 -0.5014  0.1567
                                                 0.2663
        [3,] -0.8446  0.4634  0.1364  -0.1630  -0.2727
                                                  0.1660
        [4,] 0.3745 -0.2795 -0.2795 -0.2658 -0.2658
        [5,] 1.2876 -0.6744 -0.6744 0.8935 -0.0935 -0.8612
        [6,] -1.0743 1.8687 -0.7473 -0.0383 0.2908 -0.3673
```

We can plot the residuals against the fitted values, to assess the assumption of constant variance.

```
# standardized residuals versus fitted values
plot(JS.roy1, resid(., type = "pearson") ~ fitted(.) ,
    abline = 0, id = 0.05)
```

## 4.2.3 Normality of Residuals in the Blood Data Example

LME models assume that the residuals of the model are normally distributed. The residuals can be divided according to groups according to the method of measurement. In the following examples, we separately assess normality the J method residuals (the

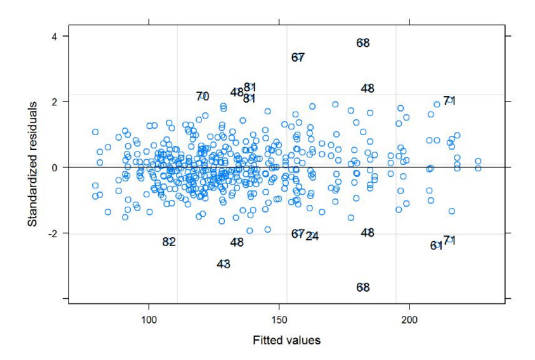


Figure 4.2.1:

first 255 residuals) and S method residuals (the remaining 255). Importantly the residuals from the J method are normally distributed, but there is non-normality of the residuals according to the S method.

```
> shapiro.test(resid(JS.roy1)[1:255])
Shapiro-Wilk normality test

data: resid(JS.roy1)[1:255]
W = 0.9931, p-value = 0.2852
```

> shapiro.test(resid(JS.roy1)[256:510])

Shapiro-Wilk normality test

data: resid(JS.roy1)[256:510]

W = 0.9395, p-value = 9.503e-09

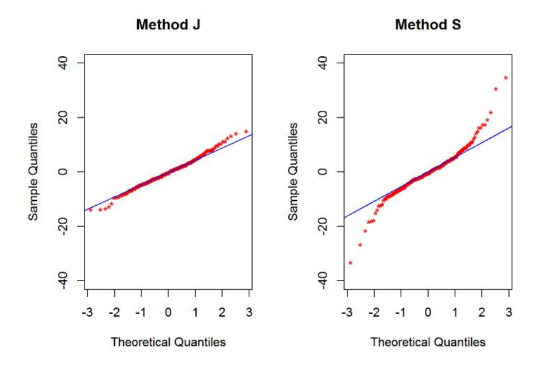


Figure 4.2.2:

#### 4.2.4 Conditional and Marginal Residuals

A residual is the difference between an observed quantity and its estimated or predicted value. For LME models, ? describes two types of residuals, marginal residuals and conditional residuals.

- A marginal residual is the difference between the observed data and the estimated (marginal) mean,  $r_{mi} = y_i x_0' \hat{b}$
- A conditional residual is the difference between the observed data and the predicted value of the observation,  $r_{ci} = y_i - x_i' \hat{b} - z_i' \hat{\gamma}$

In a model without random effects, both sets of residuals coincide (?) . We shall revert to this matter in due course.

In linear mixed effects models, diagnostic techniques may consider 'conditional' residuals. A conditional residual is the difference between an observed value  $y_i$  and the conditional predicted value  $\hat{y}_i$ .

$$\hat{\epsilon}_i = y_i - \hat{y}_i = y_i - (X_i \hat{\beta} + Z_i \hat{b}_i)$$

However, using conditional residuals for diagnostics presents difficulties, as they tend to be correlated and their variances may be different for different subgroups, which can lead to erroneous conclusions.

## 4.2.5 Residual Analysis for MCS

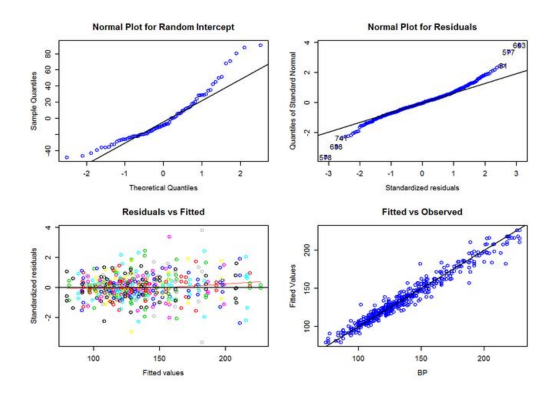


Figure 4.2.3:

## 4.2.6 PRESS Residuals and PRESS Statistic

An (unconditional) predicted value is  $\hat{y}_i = x_i' \hat{\boldsymbol{\beta}}$ , where the vector  $x_i$  is the *i*th row of  $\boldsymbol{X}$ . For an lme object, such as our fitted model JS.roy1, the predicted values for each subject can be determined using the coef.lme function.

```
> JS.roy1 %>% coef %>% head(5)

methodJ methodS

74 84.31724 91.08404

36 91.54994 97.05548
```

```
3 81.16581 96.48653
62 92.09493 90.89073
31 88.41411 103.38802
```

The (raw) residual is given as  $\varepsilon_i = y_i - \hat{y}_i$ . The PRESS residual is similarly constructed, using the predicted value for observation i with a model fitted from reduced data.

$$\varepsilon_{i(U)} = y_i - x_i' \hat{\boldsymbol{\beta}}_{(U)}$$

The PRESS statistic is the sum of the squared PRESS residuals:

$$PRESS = \sum \varepsilon_{i(U)}^2$$

where the sum is over the observations in U.

## 4.3 Introduction to Influence analysis

Model diagnostic techniques determine whether or not the distributional assumptions are satisfied, and to assess the influence of unusual observations. In classical linear models model diagnostics have been become a required part of any statistical analysis, and the methods are commonly available in statistical packages and standard text-books on applied regression. However it has been noted by several papers that model diagnostics do not often accompany LME model analyses. For linear models for uncorrelated data, it is not necessary to refit the model after removing a data point in order to measure the impact of an observation on the model. The change in fixed effect estimates, residuals, residual sums of squares, and the variance-covariance matrix of the fixed effects can be computed based on the fit to the full data alone. By contrast, in mixed models several important complications arise. Data points can affect not only the fixed effects but also the covariance parameter estimates on which the fixed-effects estimates depend.

#### 4.3.1 What is Influence

Broadly defined, influence is understood as the ability of a single or multiple data points, through their presence or absence in the data, to alter important aspects of the analysis, yield qualitatively different inferences, or violate assumptions of the statistical model. The goal of influence analysis is not primarily to mark data points for deletion so that a better model fit can be achieved for the reduced data, although this might be a result of influence analysis (?).

## 4.3.2 Importance of Influence

The influence of an observation can be thought of in terms of how much the predicted values for other observations would differ if the observation in question were not included in the model fit. Likelihood based estimation methods, such as ML and REML, are sensitive to unusual observations. Influence diagnostics are formal techniques that

assess the influence of observations on parameter estimates for  $\beta$  and  $\theta$ . A common technique is to refit the model with an observation or group of observations omitted. The basic procedure for quantifying influence is simple as follows:

- 1. Fit the model to the data and obtain estimates of all parameters.
- 2. Remove one or more data points from the analysis and compute updated estimates of model parameters.
- 3. Based on full- and reduced-data estimates, contrast quantities of interest to determine how the absence of the observations changes the analysis.

#### Measures of Influence 4.3.3

The impact of an observation on a regression fitting can be determined by the difference between the estimated regression coefficient of a model with all observations and the estimated coefficient when the particular observation is deleted. DFBETA and DFFITS are well known measures of influenc. The measure DFBETA is the studentized value of this difference. DFFITS is a statistical measured designed to a show how influential an observation is in a statistical model. DFFITS is closely related to the studentized residual.

$$DFBETA_a = \hat{\beta} - \hat{\beta}_{(a)} \tag{4.1}$$

$$= B(Y - Y_{\bar{a}} \tag{4.2})$$

$$DFBETA_{a} = \beta - \beta_{(a)}$$

$$= B(Y - Y_{\bar{a}}$$

$$DFFITS = \frac{\widehat{y}_{i} - \widehat{y}_{i(k)}}{s_{(k)}\sqrt{h_{ii}}}$$

$$(4.1)$$

The prediction residual sum of squares (PRESS) is an value associated with this calculation. When fitting linear models, PRESS can be used as a criterion for model selection, with smaller values indicating better model fits.

$$PRESS = \sum (y - y^{(k)})^2$$

#### **4.3.4 DFBETAs**

The measure that measures how much impact each observation has on a particular predictor is DFBETAs The DFBETA for a predictor and for a particular observation is the difference between the regression coefficient calculated for all of the data and the regression coefficient calculated with the observation deleted, scaled by the standard error calculated with the observation deleted.

DFBETA is a measure found for each observation in a dataset. The DFBETA for a particular observation is the difference between the regression coefficient for an included variable calculated for all of the data and the regression coefficient calculated with the observation deleted, scaled by the standard error calculated with the observation deleted.

The cut-off value for DFBETAs is  $\frac{2}{\sqrt{n}}$ , where n is the number of observations. However, another cut-off is to look for observations with a value greater than 1.00. Here cutoff means, this observation could be overly influential on the estimated coefficient.

#### **DFFITS**

DFFITS is a diagnostic meant to show how influential a point is in a statistical regression. It was proposed in 1980. It is defined as the change ("DFFIT"), in the predicted value for a point, obtained when that point is left out of the regression, "Studentized" by dividing by the estimated standard deviation of the fit at that point:

DFFITS = 
$$\frac{\widehat{y_i} - \widehat{y_{i(i)}}}{s_{(i)}\sqrt{h_{ii}}}$$

#### DFbetas for Blood Data

```
plot(JS.roy1.dfbeta$all.res1[1:255],JS.roy1.dfbeta$all.res2[256:510],
pch=16,col="blue")
abline(v=JS.roy1.dfbeta$all.res1[256],col="red")
abline(h=JS.roy1.dfbeta$all.res2[1],col="red")
```

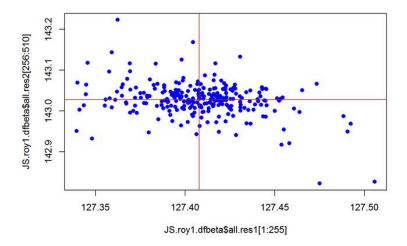


Figure 4.3.4:

## 4.4 Case Deletion Diagnostics

Case-deletion diagnostics provide a useful tool for identifying influential observations and outliers.

The computation of case deletion diagnostics in the classical model is made simple by the fact that estimates of  $\beta$  and  $\sigma^2$ , which exclude the ith observation, can be computed without re-fitting the model. Such update formulas are available in the mixed model only if you assume that the covariance parameters are not affected by the removal of the observation in question. This is rarely a reasonable assumption.

Linear models for uncorrelated data have well established measures to gauge the influence of one or more observations on the analysis. For such models, closed-form update expressions allow efficient computations without refitting the model.

Since the pioneering work of Cook in 1977, deletion measures have been applied to many statistical models for identifying influential observations. Case-deletion diagnostics provide a useful tool for identifying influential observations and outliers.

The key to making deletion diagnostics useable is the development of efficient com-

putational formulas, allowing one to obtain the case deletion diagnostics by making use of basic building blocks, computed only once for the full model.

The computation of case deletion diagnostics in the classical model is made simple by the fact that estimates of  $\beta$  and  $\sigma^2$ , which exclude the *i*-th observation, can be computed without re-fitting the model.

? describes two type of diagnostics. When the set consists of only one observation, the type is called 'observation-diagnostics'. For multiple observations, Preisser describes the diagnostics as 'cluster-deletion' diagnostics. When applied to LME models, such update formulas are available only if one assumes that the covariance parameters are not affected by the removal of the observation in question. However, this is rarely a reasonable assumption.

#### 4.4.1 Local Influence Analysis

Beckman, Nachtsheim and Cook (1987) applied the local influence method of Cook (1986) to the analysis of the linear mixed model.

While the concept of influence analysis is straightforward, implementation in mixed models is more complex. Update formulae for fixed effects models are available only when the covariance parameters are assumed to be known.

If the global measure suggests that the points in U are influential, the nature of that influence should be determined. In particular, the points in U can affect the following

- the estimates of fixed effects,
- the estimates of the precision of the fixed effects,
- the estimates of the covariance parameters,
- the estimates of the precision of the covariance parameters,
- fitted and predicted values.

#### 4.4.2 Likelihood Distances

The likelihood distance is a global summary measure that expresses the joint influence of the subsets of observations, U, on all parameters in  $\phi$  that were subject to updating. For classical linear models, the implementation of influence analysis is straightforward. ? points out the likelihood distance gives the amount by which the log-likelihood of the model fitted from the full data changes if one were to estimate the model from a reduced-data estimates. Importantly  $LD(\psi_{(U)})$  is not the log-likelihood obtained by fitting the model to the reduced data set. It is obtained by evaluating the likelihood function based on the full data set (containing all n observations) at the reduced-data estimates.

$$LD((\boldsymbol{U})) = 2[l(\hat{\boldsymbol{\phi}}) - l\hat{\boldsymbol{\phi}}_{\boldsymbol{\omega}}]$$

$$RLD((\boldsymbol{U})) = 2[l_R(\hat{\boldsymbol{\phi}}) - l_R(\hat{\boldsymbol{\phi}})_{\boldsymbol{\omega}}]$$

An overall influence statistic measures the change in the objective function being minimized. For example, in OLS regression, the residual sums of squares serves that purpose. In linear mixed models fit by maximum likelihood (ML) or restricted maximum likelihood (REML), an overall influence measure is the likelihood distance (?). In LME models, fitted by either ML or REML, an important overall influence measure is the likelihood distance (?). The procedure requires the calculation of the full data estimates  $\hat{\psi}$  and estimates based on the reduced data set  $\hat{\psi}_{(U)}$ . The likelihood distance is given by determining

$$LD_{(U)} = 2\{l(\hat{\psi}) - l(\hat{\psi}_{(U)})\} \tag{4.4}$$

$$RLD_{(U)} = 2\{l_R(\hat{\psi}) - l_R(\hat{\psi}_{(U)})\}$$
 (4.5)

For classical linear models, the implementation of influence analysis is straightforward. However, for LME models, the problem is more complex. Update formulas for the fixed effects are available only when the covariance parameters are assumed to be known. A measure of total influence requires updates of all model parameters. This can only be achieved in general is by omitting observations or cases, then refitting the model. This is a very simplistic approach, and computationally expensive.

? examines a group of methods that examine various aspects of influence diagnostics for LME models. For overall influence, the most common approaches are the *likelihood distance* and the *restricted likelihood distance*.

#### The logLik Function

logLik.lme returns the log-likelihood value of the linear mixed-effects model represented by object evaluated at the estimated coefficients. It is also possible to determine the restricted log-likelihood, if relevant, using this function. For the Blood Data Example, the loglikelihood of the JS.roy1 model can be computed as follows.

```
> logLik(JS.roy1)
'log Lik.' -2030.736 (df=8)
```

#### 4.4.3 Cook's Distance

? develops case deletion diagnostics, in particular the equivalent of Cook's distance, a well-known metric, for diagnosing influential observations when estimating the fixed effect parameters and variance components. Deletion diagnostics provide a means of assessing the influence of an observation (or groups of observations) on inference on the estimated parameters of LME models. Cook's Distance is a good measure of the influence of an observation that is a measure of aggregate impact of each observation on the group of regression coefficients, as well as the group of fitted values. If the predictions are the same with or without the observation in question, then the observation has no influence on the regression model. If the predictions differ greatly when the observation is not included in the analysis, then the observation is influential.

## Cook's Distance for Blood Data

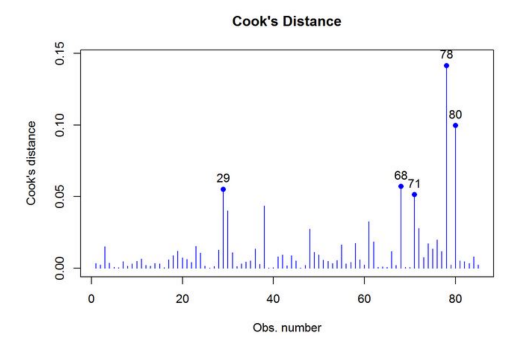


Figure 4.4.5: