Experimental Design and Data Analysis:

Calcium, inorganic phosphorus and alkaline phosphatase levels in elderly patients

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1 Introduction

In this paper the process of analysing a certain dataset is laid out. The dataset used is calcium.dat, which can be found at http://www.amstat.org/publications/jse/jse_data_archive.htm. This dataset is used for this project, because it has a good number of entries and enough features to be able to do data analysis on. In the rest of this paper the experiment is explained, the data is analyzed, the results are shown and the process is discussed. In the appendix the used R code is shown.

2 The Experiment

The experiment was set up with the goal to see if age and sex has an influence on certain concentrations in the body. The concentrations that were measured are:

- Alkaline Phosphatase International Units/Liter (Alksphos)
- Calcium mmol/L (Cammol)
- Inorganic Phosphorus mmol/L (Phosmmol)

There are 6 different labs from which the data is extracted. Next to these features, the sex, age, agegroup and patient observation number are recorded. In the calcium.dat the original data is stored with errors, in calciumgood.dat the data is already cleaned up. In this project only the calcium.dat data is used to explain how the cleaning up of the data is done. The research question we want to answer is: What influence does age have on the given concentrations in the body?

3 Data Analysis

3.1 Preparation

The data needed to be prepared to be able to read it in R. This preparation exists of replacing the empty fields by an underscore, with this the data can be read in R. First off a pairs plot was made of the data to be able to see how the data relates to each other. In Fig1(a) the foremost problem is clear, the outliers in the data are big and not logical. Furthermore has Lab more than 6 categories, these problems can be explained by human error. And lastly the lab 3 has strange measurements with the cammol, this might be because of confusion of the measurement unit. The following values were changed:

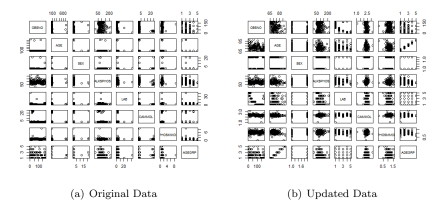


Figure 1: Pairsplot of the data

- Removed Ages over 110
- Removed Sex which is not in the category 1 or 2
- Removed Lab categories which are over 6
- Removed Phosmmol over 2
- Divided Cammol of Lab 3 by 10 (this is visually tested by using a pairs plot)

After removal the same plot is made, see: Fig1(b). Here you can see the correlations between the features better than in Fig1(a).

3.2 Analysis

The feature that needs to be analyzed is age, this can be done by either using the feature age or the feature age group. The age group exists of 5 levels and is thus categorical. The different age groups are: 1=65-69, 2=70-74, 3=75-79, 4=80-84, 5=85-89 Years. In Fig2 the relation between the different concentrations and the age groups are visually represented.

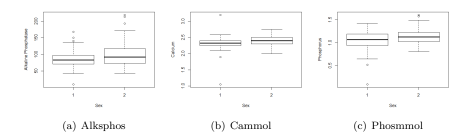


Figure 2: Boxplots of the concentrations between sexes

Analysing whether the data in CAMMOL, ALKSPHOS and PHOSMMOL is normal:

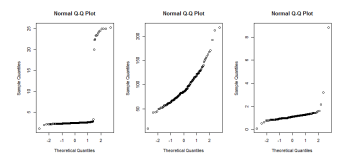


Figure 3: QQ plot of raw data

Obviously these are not normal, most likely due to outliers. After removing outliers we obtain:

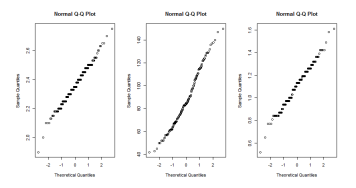


Figure 4: QQ plot after removing outliers

Clearly these are more likely normal, even though we do have a step type pattern, likely due to rounding.

There is some difference between the different age groups with all of the concentrations. But with Alksphos the difference only seems to be in the variety of the values, the same holds for Cammol. For Phosmmol there does seems to be a difference in the age groups, the median drops with the older ages. The difference in the first two concentrations is because of the number of people in the different age groups, see Table 1. This shows that there are only three patients in the last age group and not too much in the group before that. This explains that the variety seems to lower with the higher age groups.

Table 1: The number of patients per age group

Groups	1	2	3	4	5
Patients	56	70	38	10	3

To see if sex, lab and age have an influence on the concentrations, the interaction plots in Fig5 and Fig6 are made. This shows that all three features have an influence on the measured concentrations. The age has an influence over the concentrations, which is good for the research question. But the lab and sex also influence the outcome, which means that these need to be taken into account with modeling the data.

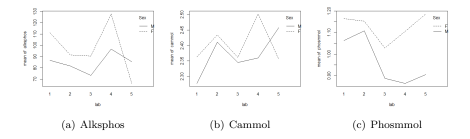


Figure 5: Interaction plots of the sex and labs of the different concentrations

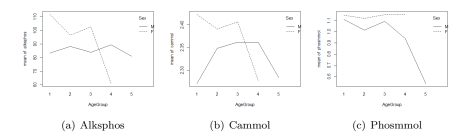


Figure 6: Interaction plots of the sex and age groups of the different concentrations

Table 2: Kruskal-Wallis rank sum test

Variable	p
CAMMOL~SEX	0.001539
CAMMOL~LAB	1.433e-05
CAMMOL~AGEGRP	0.5451
ALKSPHOS~SEX	0.007631
ALKSPHOS~LAB	0.001325
ALKSPHOS~AGEGRP	0.6927
PHOSMMOL~SEX	0.007711
PHOSMMOL~LAB	0.159
PHOSMMOL~AGEGRP	0.006144

Analysis of Variance Table

Response: CAMMOL

CAMMOL~SEX*LAB

Residuals 153 2.32036 0.015166

CAMMOL~SEX*AGEGRP

CAMMOL~LAB*AGEGRP

Signif. codes:

 $0 \quad *** \quad 0.001 \quad ** \quad 0.01 \quad * \quad 0.05 \quad . \quad 0.1 \qquad \qquad 1$

Figure 7: Summary of computed linear model

Analysis of Variance Table

Signif. **codes**: 0 *** 0.0

0.001

Response: ALKSPHOS ALKSPHOS~SEX*LAB Df Sum Sq Mean Sq F value Pr(>F) SEX1939 1938.76 1 $3.6568 \ 0.05781$. LAB 6806 1701.42 $3.2092 \ 0.01471 *$ 4 SEX:LAB 4 1686 421.61 $0.7952 \ 0.53010$ Residuals 145 76875530.18ALKSPHOS~SEX*AGEGRP Df Sum Sq Mean Sq F value Pr(>F) SEX1 1702 1701.75 $3.1397 \ 0.07838$. **AGEGRP** 1375 343.84 $0.6344 \ 0.63871$ 4 SEX:AGEGRP 3 3591 1197.11 $2.2087 \ 0.08933$. Residuals 154 83468 542.00ALKSPHOS~LAB*AGEGRP Df Sum Sq Mean Sq F value Pr(>F) LAB 4 6057 1514.19 $2.7606 \ 0.03025 *$ **AGEGRP** 4 1123 280.79 $0.5119\ 0.72707$ 475.42 $0.8668 \ 0.58935$ LAB:AGEGRP 6180 1374048 548.50Residuals 135

Figure 8: alk 2 way anova

0.05

0.1

1

0.01

Analysis of Variance Table

 $Response: \ P\!H\!O\!S\!M\!M\!O\!L$

PHOSMMOL~SEX*LAB

	Df	Sum Sq	Mean Sq	F value	$\Pr(>F)$	
SEX	1	0.2500	0.250022	9.1038	0.002991	**
LAB	4	0.4122	0.103060	3.7526	0.006101	**
SEX:LAB	4	0.1639	0.040972	1.4919	0.207392	

Residuals 152 4.1744 0.027463

PHOSMMOL~SEX*AGEGRP

	Df	Sum Sq	Mean Sq	F value	$\Pr(>F)$	
SEX	1	0.2392	0.239170	8.9478	0.003218	**
AGEGRP	4	0.3847	0.096177	3.5981	0.007732	**
${\bf SEX:} {\bf AGEGRP}$	3	0.0788	0.026266	0.9827	0.402506	
Residuals	160	4.2767	0.026730			

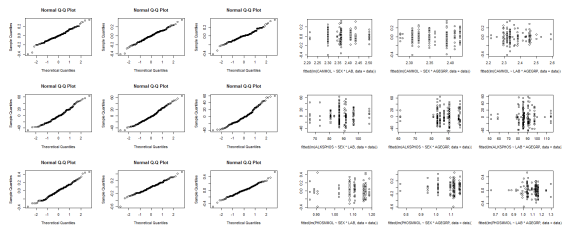
PHOSMMOL~LAB*AGEGRP

	Df	Sum Sq	Mean Sq	F value	Pr(>F)	
LAB	4	0.1703	0.042569	1.5556	0.18946	
AGEGRP	4	0.3204	0.080096	2.9270	0.02311	*
LAB:AGEGRP	13	0.5003	0.038486	1.4064	0.16321	
Residuals	142	3.8858	0.027365			

Signif. codes:

0 *** 0.001 0.01* 0.05 . 0.1 1

Figure 9: phos 2 way anova



- (a) QQ plot of residuals for all 9 linear models
- (b) Plots of fitted residuals for all 9 linear models

Figure 10: Graphical tests for normality of residuals

3.3 Modeling

Modeling the data can be done in several ways. The first one tried is a step down linear regression approach for each of the concentrations. For Alksphos it follows the following steps:

- $ALKSPHOS \sim AGE + SEX + LAB + AGEGRP$
- $ALKSPHOS \sim AGE + SEX + LAB$
- $ALKSPHOS \sim SEX + LAB$
- $\bullet \ \ ALKSPHOS = 88.96 + 17.91 * SEX2 11.69 * LAB2 16.25 * LAB3 + 18.03 * LAB4 30.20 * LAB5 + 18.03 * LAB4 10.20 * LAB5 + 10.20 * L$

For Cammol it follows the following steps:

- $CAMMOL \sim AGE + SEX + LAB + AGEGRP$
- $CAMMOL \sim AGE + SEX + LAB$
- $CAMMOL \sim SEX + LAB$
- CAMMOL = 2.29 + 0.06 * SEX2 + 0.11 * LAB2 + 0.03 * LAB3 + 0.14 * LAB4 + 0.07 * LAB5

For Phosmmol it follows the following steps:

- $PHOSMMOL \sim AGE + SEX + LAB + AGEGRP$
- $PHOSMMOL \sim AGE + SEX + LAB$
- $PHOSMMOL \sim AGE + SEX$
- PHOSMMOL = 2.02 0.01 * AGE + 0.17 * SEX2

As our research question is about what influence age has on the concentration and only Phosmmol has a model with age in it, this is not a good approach for this data.

4 Discussion

The data was realistic to work with, it contained some errors. At the start of the project, this was a challenge. To get all the data to correspond to realistic data. The good data was also supplied, but we decided against using it for the experience of working with error prone data. After the preprocessing the real data analysis could begin. The research question that was decided on was: What influence does age have on the given concentrations in the body? The reason to not only look at age and the three concentrations was to be able to see if more features had an influence on the concentrations. If this was not done the results would not be correct, because the features lab and sex also had an effect on the concentrations. In the models created it was clear that age did not have as big an influence as first thought. In two of the three models, age was not included. This means that age has a small influence on the measured concentrations, which is the answer to our research question.

5 R-Code

```
data = read.table('calcium.dat.txt', na.strings="_", header=TRUE)
# exploration
pairs (data)
# data preparation
data[!is.na(data$AGE) & !(data$AGE <= 110),]$AGE<-NA
data[!is.na(data\$SEX) \& !(data\$SEX == 1|data\$SEX == 2),]\$SEX < -NA
data[!is.na(data$LAB) \& !(data$LAB < 6),]$LAB < -NA
data[!is.na(data$PHOSMMOL) & !(data$PHOSMMOL < 2),]$PHOSMMOL<-NA
\mathbf{data\$CAMMOL[!is.na(data\$CAMMOL) \& (data\$CAMMOL>10)]} < - (\mathbf{data\$CAMMOL[!is.na(data\$CAMMOL)}) = - (\mathbf{data\$CAMMOL[!is.na(data§CAMMOL)}) = - (\mathbf{data\$CAMMOL[!is.na(data§CAMMOL)}) = - (\mathbf{data\$CAMMOL[!is.na(data§CAMMOL)}) = - (\mathbf{data§CAMMOL[!is.na(data§CAMMOL)}) = - (\mathbf{data§CAMMOL[!is.na(data§CAMMOL]}) = - (\mathbf{data§CAMMOL[!i
data$SEX <- as.factor(data$SEX)
data$LAB <- as.factor(data$LAB)
data$AGEGRP <- as.factor(data$AGEGRP)
# exploration
pairs (data)
summary(data$AGEGRP)
# data preparation, replace NA with mean
sex = data$SEX
sex[is.na(sex)]<-mean(na.omit(data$SEX))
lab = data$LAB
lab[is.na(lab)] < -mean(na.omit(data$LAB))
cammol = data$CAMMOL
cammol [is.na(cammol)]<-mean(na.omit(data$CAMMOL))
alksphos = data$ALKSPHOS
alksphos [is.na(alksphos)] <-mean(na.omit(data$ALKSPHOS))
phosmmol = data PHOSMMOL
[is.na(phosmmol)] \le mean(na.omit(data$PHOSMMOL))
\# data preperation, make sex M and F instead of 1 and 2
Sex = as.character(sex)
Sex[sex == 1] = "M"
Sex[sex == 2] = "F"
Sex = as.factor(Sex)
AgeGroup = AGEGRP
# interaction plots
interaction.plot(lab, Sex, cammol)
```

```
interaction.plot(lab,Sex,alksphos)
interaction.plot(lab, Sex, phosmmol)
interaction.plot(AgeGroup, Sex, cammol)
interaction.plot(AgeGroup, Sex, alksphos)
interaction.plot(AgeGroup, Sex, phosmmol)
# step down linear regression
#ALKSPHOS
summary(lm(ALKSPHOS~AGE+SEX+LAB+AGEGRP, data=data))
summary(lm(ALKSPHOS~AGE+SEX+LAB, data=data))
summary(lm(ALKSPHOS~SEX+LAB, data=data))
#CAMMOL
summary(lm(CAMMOL~AGE+SEX+LAB+AGEGRP, data=data))
\mathbf{summary}(\mathbf{lm}(CAMMOL^*AGE+SEX+LAB}, \mathbf{data} = \mathbf{data}))
summary(lm(CAMMOL~SEX+LAB, data=data))
#PHOSMMOL
summary(lm(PHOSMMOL~AGE+SEX+LAB+AGEGRP, data=data))
\mathbf{summary}(\mathbf{lm}(\mathtt{PHOSMMOL}^{\mathtt{A}}\mathtt{A}\mathtt{G}\mathtt{E}\!+\!\mathtt{SEX}\!+\!\mathtt{L}\mathtt{A}\mathtt{B},\mathbf{data}\!=\!\mathbf{data}))
summary(lm(PHOSMMOL~AGE+SEX, data=data))
```