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STAT6180-Applied Statistic

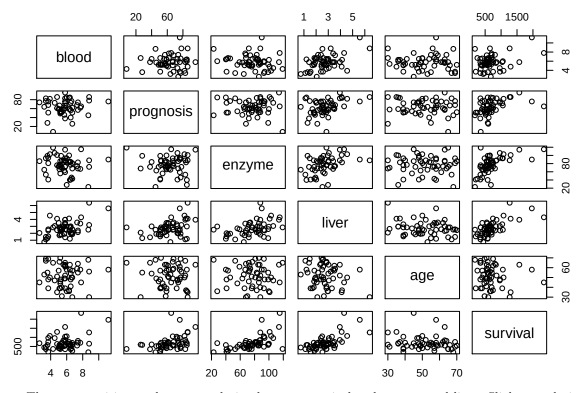
Assignment Semester 1

Question 1

A medical research team wants to investigate the survival time of patients that have a particular type of liver operation as part of their treatment. For each patient in the study, the following variables were recorded:

blood	Blood clotting Index
prognosis	Prognosis Index
enzyme	Enzyme function Index
liver	Liver function Index
age	Age of the patient, in years
gender	Gender of the patient, (Male of Female)
survival	Survival time of the patient after surgery (in days)

a. Produce a scatterplot of the data and comment on the features of the data and possible relationships between the response and predictors and relationships between the predictors themselves.



There are positive moderate correlation between survival and enzyme and liver. Slight correlation between

survival and prognosis.

Why it is necessary to remove the gender variable to compute the correlation matrix?

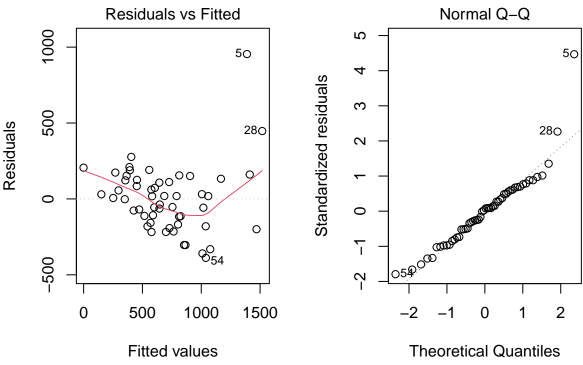
Because the gender variable is a categorical variables. To compute the correlation matrix, every variables must be numeric.

b. Compute the correlation matrix of the dataset and comment.

##		blood	prognosis	enzyme	liver	age	survival
##	blood	1.00	0.09	-0.15	0.50	-0.02	0.35
##	prognosis	0.09	1.00	-0.02	0.37	-0.05	0.42
##	enzyme	-0.15	-0.02	1.00	0.42	-0.01	0.58
##	liver	0.50	0.37	0.42	1.00	-0.21	0.67
##	age	-0.02	-0.05	-0.01	-0.21	1.00	-0.12
##	survival	0.35	0.42	0.58	0.67	-0.12	1.00

The correlation matrix shows that there are moderate correlation between survival and liver(0.67) and enzyme(0.58). Low correlation between survival and blood(0.35) and prognosis(0.42). The correlation between survival and age is -0.12, which is close to 0, indicates that no linear relationship between these variables.

c. Fit a model using all the predictors to explain the survival response.Conduct an F-test for the overall regression i.e. is there any relationship between the response and the predictors. In your answer:



There is a significant pattern in the residuals vs fitted plot and the normal Q-Q plot of residuals close to linear.

• Write down the mathematical multiple regression model for this situation, defining all appropriate parameters.

##	(Intercept)	blood	prognosis	enzyme	liver
##	-1179.1888797	86.6437068	8.5012606	11.1245627	38.5068155
##	age	gender			

-2.3408883 -0.2201138

$$sur\hat{v}ival = -1179.1888797 + 86.6437068blood + 8.5012606prognosis + 11.1245627enzyme \\ +38.5068155liver - 0.2201138age - 0.2201138gender$$

• Write down the Hypotheses for the Overall ANOVA test of multiple regression.

$$H_0: \beta_1=\beta_2=\beta_3=\beta_4=\beta_5=0;$$
 $H_1: \beta_i\neq 0$ for at least one i (not all β_i parameters are zero)

• Produce an ANOVA table for the overall multiple regression model (One combined regression SS source is sufficient).

```
## Analysis of Variance Table
##
## Response: survival
##
             Df Sum Sq Mean Sq F value
              1 1005152 1005152 18.5060 8.502e-05
## blood
## prognosis 1 1278496 1278496 23.5385 1.387e-05 ***
              1 3442172 3442172 63.3742 2.915e-10 ***
## enzyme
## liver
                  57862
                          57862
                                1.0653
                                           0.3073
## age
                  33032
                          33032
                                0.6082
                                           0.4394
                                0.0000
                                           0.9974
## gender
                      1
                              1
              1
## Residuals 47 2552807
                          54315
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

• Compute the F statistic for this test.

$$Full RegSS = RegSS_{blood} + RegSS_{prognosis|blood} + RegSS_{enzyme|blood\&prognosis} \\ + RegSS_{liver|blood\&prognosis\&enzyme} + RegSS_{age|blood\&prognosis\&enzyme\&liver} \\ + RegSS_{gender|blood\&prognosis\&enzyme\&liver\&age}$$

FullRegSS = 1005152 + 1278496 + 3442172 + 57862 + 33032 + 1 = 5816715

$$RegMS = \frac{RegSS}{k} = \frac{5816715}{6} = 969452.5$$

Test statistic:
$$F_{obs} = \frac{RegMS}{ResMS} = \frac{969452.5}{54315} = 17.84871$$

• State the Null distribution.

$$H_0: \beta_{blood} = \beta_{prognosis} = \beta_{enzyme} = \beta_{liver} = \beta_{age} = \beta_{gender} = 0;$$

 $H_1: \text{not all} \quad \beta_i = 0$

• Compute the P-Value

[1] 1.190218e-10

P-Value:
$$P(F_{6.47} >= 17.84871) = 1.190218e - 10 < 0.05$$

• State your conclusion (both statistical conclusion and contextual conclusion). P-value is 1.190218e-10. As P-value < 0.05, reject H_0 .

There is a significant linear relationship between survival and at least one of the five predictor variables.

d. Using model selection procedures discussed in the course, find the best multiple regression model that explains the data.

```
##
## lm(formula = survival ~ blood + prognosis + enzyme + liver +
##
       age + gender, data = surg.new)
##
## Residuals:
##
      Min
                1Q Median
                                3Q
                                       Max
##
  -388.25 -147.61
                     11.72 124.67
                                    954.44
##
## Coefficients:
##
                 Estimate Std. Error t value Pr(>|t|)
                            283.8232
                                     -4.155 0.000136 ***
## (Intercept) -1179.1889
## blood
                 86.6437
                             27.4920
                                       3.152 0.002825 **
## prognosis
                  8.5013
                              2.1601
                                       3.936 0.000273 ***
## enzyme
                  11.1246
                              1.9820
                                      5.613 1.03e-06 ***
## liver
                  38.5068
                             51.7967
                                      0.743 0.460926
                  -2.3409
                              3.0141 -0.777 0.441257
## age
                  -0.2201
                             67.5146 -0.003 0.997413
## gender
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 233.1 on 47 degrees of freedom
## Multiple R-squared: 0.695, Adjusted R-squared: 0.656
## F-statistic: 17.85 on 6 and 47 DF, p-value: 1.19e-10
```

Removing one none significant variable (if there are many none significant vars, pick the largest P-value). Remove \mathbf{gender} P-value = 0.997413

```
##
                   Estimate Std. Error
                                          t value
                                                      Pr(>|t|)
## (Intercept) -1179.366654 275.619347 -4.2789690 8.913076e-05
## blood
                  86.630445 26.904719 3.2198978 2.302423e-03
## prognosis
                  8.501113
                              2.137047 3.9779712 2.337301e-04
## enzyme
                  11.124165
                              1.957529
                                       5.6827582 7.623756e-07
## liver
                  38.553562 49.251408 0.7827911 4.375949e-01
                  -2.339958
                              2.969120 -0.7880981 4.345142e-01
## age
```

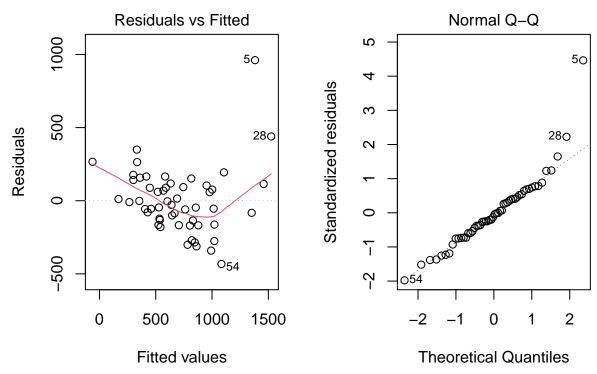
Removing one none significant variable (if there are many none significant vars, pick the largest P-value). Remove liver P-value = 0.437595

```
##
                   Estimate Std. Error
                                         t value
                                                     Pr(>|t|)
## (Intercept) -1246.654818 260.835337 -4.779471 1.640827e-05
## blood
                 100.659551 19.987172 5.036208 6.831794e-06
## prognosis
                  9.290889
                              1.876432
                                        4.951359 9.138843e-06
## enzyme
                                        8.058360 1.555971e-10
                  12.101482
                              1.501730
                              2.840741 -1.051209 2.983194e-01
## age
                  -2.986213
```

Removing one none significant variable (if there are many none significant vars, pick the largest P-value). Remove \mathbf{age} P-value = 0.298

```
##
## Call:
## lm(formula = survival ~ blood + prognosis + enzyme, data = surg.new)
##
## Residuals:
##
     Min
              1Q Median
                            3Q
                                   Max
## -432.4 -134.3 -19.1 111.9
                                961.1
##
## Coefficients:
##
                Estimate Std. Error t value Pr(>|t|)
## (Intercept) -1410.847
                            209.118 -6.747 1.50e-08 ***
## blood
                 101.054
                              20.005
                                      5.052 6.22e-06 ***
                                       5.000 7.43e-06 ***
## prognosis
                   9.382
                              1.876
## enzyme
                  12.128
                               1.503
                                       8.069 1.30e-10 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 229.9 on 50 degrees of freedom
## Multiple R-squared: 0.6841, Adjusted R-squared: 0.6652
## F-statistic: 36.1 on 3 and 50 DF, p-value: 1.469e-12
All predictors are significant and R^2 = 0.6652
 * Finallized fitted equation:
         sur\hat{v}ival = -1410.846901 + 101.053887blood + 9.381966prognosis + 12.127807enzyme
```

e. Validate your final model and comment why it is not appropriate to use the multiple regression model to explain the survival time.



From above picture show the residual vs fitted plot and the normal Q-Q plot for the final model of the survival response. The linearity of the points in the normal Q-Q plot suggests that the data are close to normally distributed. However, the residuals vs fitted plot has a pattern. This is the reason why this final model is not appropriate to use the multiple regression model to explain the survival time. In this case, In this transformation the response variable is needed.

- f. Re-fit the model using log(survival) as the new response variable. In your answer,
 - Use the model selection procedure discussed in the course starting with log(survival) as the response and start with all the predictors.

```
##
                   Estimate Std. Error
                                             t value
                                                         Pr(>|t|)
## (Intercept)
                4.100996997 0.302780511 13.54445497 7.665031e-18
## blood
                0.094858430 0.029328278
                                         3.23436749 2.233610e-03
                                         5.64980388 9.084175e-07
                0.013019518 0.002304419
## prognosis
## enzyme
                0.016245445 0.002114393
                                         7.68326550 7.593288e-10
## liver
               -0.003132326 0.055256339 -0.05668718 9.550347e-01
               -0.004863398 0.003215405 -1.51253027 1.370946e-01
## age
               -0.066139894 0.072024114 -0.91830208 3.631495e-01
## gender
```

Removing one none significant variable (if there are many none significant vars, pick the largest P-value). Remove liver P-value = 0.95503

```
##
                   Estimate
                             Std. Error
                                            t value
                                                        Pr(>|t|)
                4.105132163 0.290794570 14.1169492 1.039336e-18
## (Intercept)
## blood
                0.093737869 0.021439405
                                         4.3722235 6.576975e-05
                0.012959509 0.002025519
                                         6.3981183 6.163295e-08
## prognosis
## enzyme
                0.016170082 0.001626984
                                         9.9386870 3.096659e-13
               -0.004810137 0.003042976 -1.5807348 1.205066e-01
## age
               -0.065009685 0.068487175 -0.9492242 3.472615e-01
## gender
```

Removing one none significant variable (if there are many none significant vars, pick the largest P-value). Remove \mathbf{gender} P-value = 0.3472615

```
## Estimate Std. Error t value Pr(>|t|)
## (Intercept) 4.028530718 0.279090496 14.434496 2.820187e-19
## blood 0.094845060 0.021386020 4.434909 5.202486e-05
## prognosis 0.013198656 0.002007759 6.573826 3.035041e-08
## enzyme 0.016402478 0.001606832 10.207960 1.012035e-13
## age -0.004766708 0.003039557 -1.568224 1.232646e-01
```

Removing one none significant variable (if there are many none significant vars, pick the largest P-value). Remove \mathbf{age} P-value = 0.1232646

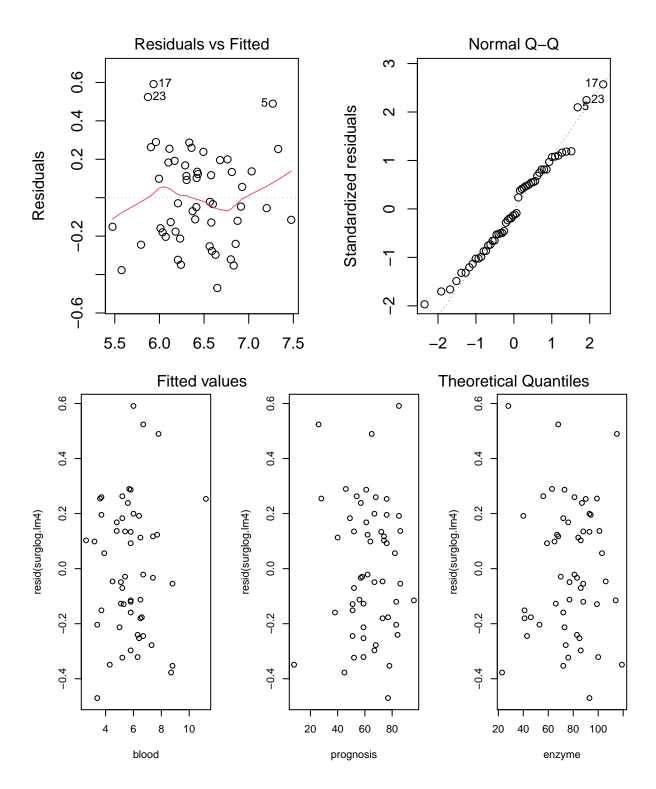
```
##
## Call:
## lm(formula = logsurvival ~ blood + prognosis + enzyme, data = surg.new)
##
## Residuals:
##
       Min
                  1Q
                      Median
                                    3Q
                                           Max
## -0.46994 -0.17938 -0.03116 0.17959 0.59105
##
## Coefficients:
##
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) 3.766441
                         0.226757 16.610 < 2e-16 ***
## blood
              0.095475
                         0.021692
                                    4.401 5.66e-05 ***
              0.013344
                         0.002035
                                    6.558 2.95e-08 ***
## prognosis
## enzyme
              0.016444
                         0.001630 10.089 1.19e-13 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.2493 on 50 degrees of freedom
## Multiple R-squared: 0.7572, Adjusted R-squared: 0.7427
## F-statistic: 51.99 on 3 and 50 DF, p-value: 2.137e-15
```

All predictors are significant and $R^2 = 0.7427$ which is better than the model with the survival response variable.

```
log(\hat{survival}) = -3.76644097 + 0.09547451blood + 0.01334404prognosis + 0.01644450enzyme
```

g. Validate your final model with the log(survival) response. In particular, in your answer,

^{*} Finallized fitted equation:



- * The residuals vs fitted plot shows some pattern in but not significant.
- * There is slight curvature in the normal quantile plot of residuals, the data are close to normally distributed.
- * No sign of curvature in the residuals against predictors plots.
- Explain why the regression model with log(survival) response variable is superior to the model with the

survival response variable

From above picture show the residual vs fitted plot and the normal Q-Q plot for the log(survival) response. Comparing with the final model of the survival response, the normality assumption within the log(survival) model is better as there is some pattern but not significant in the residuals vs fitted plot and the normal Q-Q plot is close to normally distributed. It clarifies the reason why log(survival) is superior to the other response.

Question 2

A car manufacturer wants to study the fuel efficiency of a new car engine. It wishes to account for any differences between the driver and production variation. The manufacturer randomly selects 5 cars from the production line and recruits 4 different test drivers.

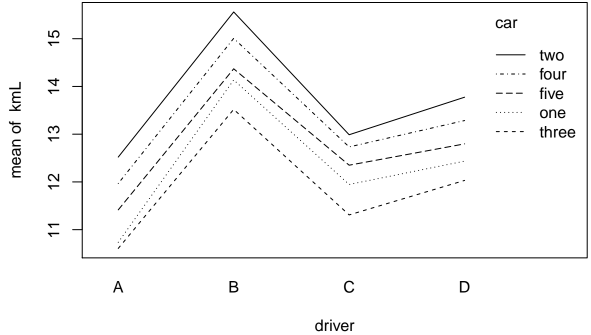
kmL	The observed efficiency of the car in km/L over a standard course
car	The specific car (labelled 1, 2, 3, 4 or 5)
driver	The driver of the car (labelled A, B, C, D)

a. For this study, is the design balanced or unbalanced? Explain why.

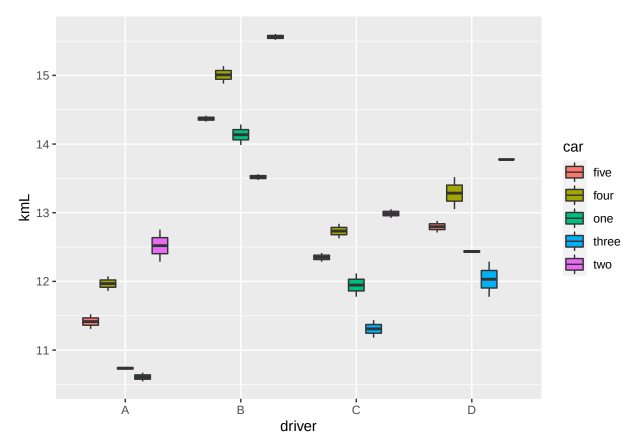
##	(
##	${\tt driver}$	five	four	one	${\tt three}$	two
##	A	2	2	2	2	2
##	В	2	2	2	2	2
##	C	2	2	2	2	2
##	D	2	2	2	2	2

This is a balanced design because there is the same no. of replicates for each treatment combinations.

b. Construct two different preliminary graphs that investigate different features of the data and comment.



As the lines are not parallel, interaction could be there.



Overall, there are different variation among each group.

- * Similar spread for drive A, C and D groups
- * Combination of drive B have higher kmL than others.
- * Groups of driver A and car-one and car-three have the lowest kmL
- c. Analyse the data, stating null and alternative hypothesis for each test, and check assumptions.

```
##
               Df Sum Sq Mean Sq F value
                                            Pr(>F)
## driver
                   50.66
                          16.887
                                   531.60 < 2e-16 ***
                   17.12
                            4.280
                                   134.73 3.66e-14 ***
## car
                4
               12
                    0.44
                            0.037
                                     1.16
                                              0.371
## driver:car
## Residuals
               20
                    0.64
                            0.032
                   0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Signif. codes:
```

Model: $Y = \mu + \alpha_i + \beta_j + \gamma_{ij} + \epsilon$

where ϵ_{ijk} are $N(0, \sigma^2)$ random variables

 μ : overall population mean

 α_i : main effect on driver

 β_i : main effect on car

 γ_{ij} : interaction effect between driver and car

 ϵ : error term

Hypotheses $H_0: \gamma_{ij} = 0$ against $H_1:$ at least one γ_{ij} non-zero

Because P-value = 0.371 > 0.05, γ_{ij} is not significant.

No evidence to suggest that the two factors (driver and car) are not independent.

As interaction is not significant, re-fit the model with main effects only.

```
##
              Df Sum Sq Mean Sq F value Pr(>F)
## driver
                  50.66
                         16.887
                                  501.5 <2e-16 ***
## car
                  17.12
                          4.280
                                  127.1 <2e-16 ***
               4
## Residuals
              32
                   1.08
                          0.034
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

Main Effects: Driver

```
Model: y_{ijk} = \mu + \alpha_i + \beta_j + \epsilon
```

Hypotheses: $H_0: \alpha_i = 0$ against $H_1:$ at least one α_i non-zero

P-Value <2e-16, less than 0.05, reject H0

Driver type is significant.

Main Effects: Car

```
Model: y_{ijk} = \mu + \alpha_i + \beta_j + \epsilon
```

 $\mbox{Hypotheses}: \quad H_0: \beta_j = 0 \quad \mbox{against} \quad H_1: \mbox{at least one} \quad \beta_j \mbox{ non-zero}$

P-Value <2e-16, less than 0.05, reject H0

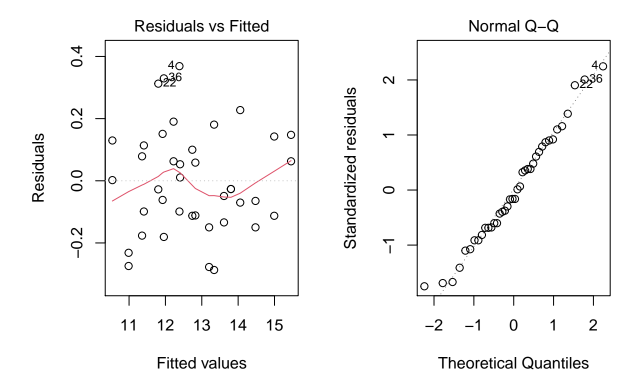
Car type is significant.

Both the driver and car effects are highly significant (P-Value < 0.001)

Coefficients table:

```
##
                 Estimate Std. Error
                                        t value
                                                    Pr(>|t|)
## (Intercept) 11.4076764 0.08206422 139.009140 4.363405e-46
## driverB
                3.0695397 0.08206422 37.404118 5.526204e-28
## driverC
                0.8162765 0.08206422
                                       9.946801 2.582755e-11
                1.4157295 0.08206422 17.251484 9.143265e-18
## driverD
## carfour
                0.5154871 0.09175059
                                       5.618352 3.285809e-06
               -0.4198297 0.09175059
                                      -4.575771 6.788252e-05
## carone
## carthree
               -0.8662309 0.09175059
                                      -9.441148 9.081856e-11
## cartwo
                0.9778312 0.09175059 10.657493 4.666698e-12
```

From the coefficients table, the effect of car-one and car-three are to reduce kml by 0.41983 and 0.86623 respectively, although the other variables are increase kml.



- Residuals vs Fitted plot shows a negligible pattern, variability among residuals vs fitted is not constant. There are several outliers, with residuals close to 0.4.
- The normal Q-Q plot of residuals follows a linear trend, residuals look close to normally distributed.
- d. State your conclusions about the effect of driver and car on the efficiency kmL. These conclusions are only required to be at the qualitative level and can be based off the outcomes of the hypothesis tests in
- e. and the preliminary plots in b. You do not need to statistically examine the multiple comparisons between contrasts and interactions.

From the above findings, we can say that both null hypotheses of the main effects (driver and car) are rejected. Moreover, the normal Q-Q plot follows a linear trend and residual plots have no pattern, suggesting the linear model adequately. From this, we can conclude that the efficiency of the car in km/L depend upon the factors test driver and cars.