proj2

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3/5/2022

Background

Part 1

a Total Population $(X_{1,1})$ ## ## The decimal point is 6 digit(s) to the right of the | ## ## ## ## 1 | 000000122233333444 1 | 55699 ## ## 2 | 1134 2 | 58 ## ## 3 | ## 3 I ## 4 | ## 4 | ## 5 | 1 ## 5 | ## 6 | 6 I ## 7 | ## 7 I 8 | ## 8 | 9 Land Area $(X_{1,2})$ ## ## The decimal point is 3 digit(s) to the right of the | ## ## ## 2 | 0001111466778 ## 3 | 3344688 ## ## 4 | 00122368 5 | 45 ## ## 6 I 023 7 | 29 ## ## 8 | 11 9 | 22 ## 10 I ## ## 11 | ## 12 | ## 13 | ## 14 | ## 15 | ## 16 | ## 17 | ## 18 | ## 19 |

```
## 20 | 1
```

```
Total Personal Income (X_{1,3}) and (X_{2,3})
```

```
##
##
    The decimal point is 4 digit(s) to the right of the |
##
##
     ##
     1 | 000000000001111111111222223333344444445555555567788888888999
     2 | 001111233344477788899
##
     3 | 0255678899
##
     4 | 19
##
     5 | 59
##
     6 I
##
##
     7 |
##
     8 |
##
     9 |
##
    10 |
##
    11 | 1
##
    12 |
##
    13 l
##
    14 |
##
    15 |
##
    16 |
    17 |
##
    18 | 4
Population Density (X_{2,1})
```

```
##
    The decimal point is 3 digit(s) to the right of the |
##
##
    ##
    2 | 00001112233456700111145
##
    4 | 05884
##
##
    6 | 2464
##
    8 | 19
    10 | 378
##
##
   12 |
   14 | 4
##
##
    16 I
##
    18 |
##
    20 |
##
   22 |
##
   24 |
##
   26 I
##
   28 |
##
   30 |
   32 | 4
##
```

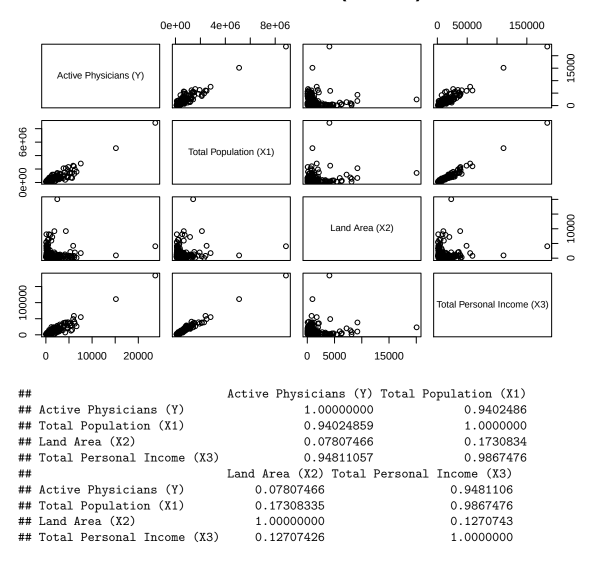
Total Population over 64 $(X_{2,2})$

```
##
##
   The decimal point is at the |
##
   2 | 0
##
##
   4 | 47890389
   6 | 1123455677990134566678899
##
##
   ##
   ##
   14 | 000011111112233344444555677889000000111122223455667778
##
##
   16 | 12556699901122345
   18 | 06778
##
##
   20 | 070
##
   22 | 018828
   24 | 47
##
##
   26 | 055
   28 | 1
##
##
   30 | 7
##
   32 | 138
```

Stem and Leaf plots tell us about the distribution of the data. In this case we can see that all of the predictor variable data is right-skewed. We see that Total Population, Land Area, Total Personal Income, and Population Density all have outliers.

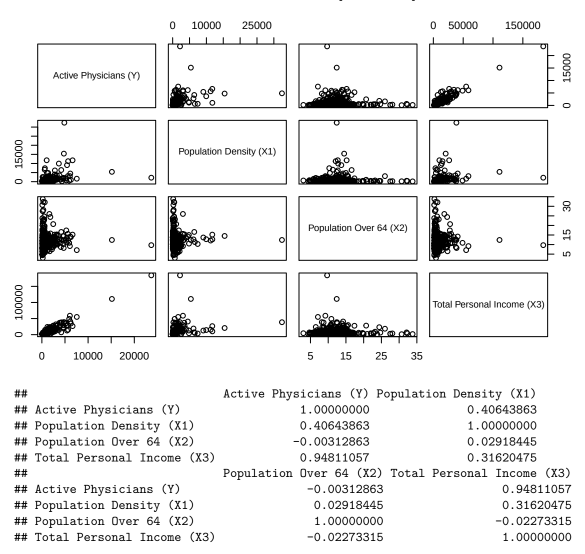
b

Scatter Plot Matrix (Model 1)



Total Population (X1) and Total Personal Income (X3) are highly correlated to The Number of Active Physicians (Y) with correlation coefficients of 0.940 and 0.948, respectively. This means there is a strong linear relationship between the predictor variables and number of active physicians. Land Area (X2) has a weak correlation with Active Physicians (Y) with a coefficient of 0.078. This means there is a weak linear relationship between land area and the number of active physicians. Based on the scatter plot, Total Population and Total Personal Income are positively correlated with Active Physicians, meaning that when they increase, the number of Active Physicians increases as well.

Scatter Plot Matrix (Model 2)



Of the three predictor variables, Total Personal Income (X3) has the strongest linear relationship with the number of Active Physicians (Y) with a correlation coefficient of 0.948. Then, Population Density (X1) has a weaker linear relationship with Active Physicians (Y) with a correlation coefficient of 0.406. Finally, Population Over 64 (X2) has the weakest linear relationship with Active Physicians (Y) with a correlation coefficient of -0.00312. X2 also has the only negative correlation coefficient. Based on the scatter plot, Population Density and Total Personal Income are positively correlated with Active Physicians (Y). Therefore, as those predictors increase, the number of active physicians will also increase.

 \mathbf{c}

 ${\bf Model}\ {\bf 1}$

$$Y = -13.316 + 0.001X_1 + -0.066X_2 + 0.094X_3$$

 $\bf Model~2$

$$Y = -170.574 + 0.096X_1 + 6.34X_2 + 0.127X_3$$

 \mathbf{d}

Model 1 R^2 : 0.9026432 Model 2 R^2 : 0.9117491

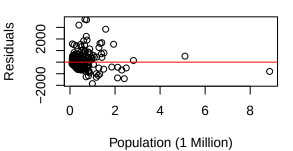
Based on the \mathbb{R}^2 value, Model 2 is a slightly better model, but they are close.

e: Model 1

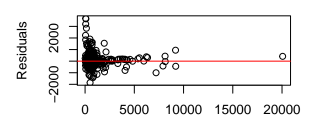
Model 1: Residuals ~ Fitted values

Sending Sendin

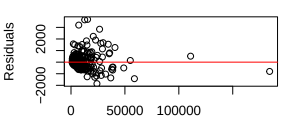
Model 1: Residuals ~ Population



Model 1: Residuals ~ Area

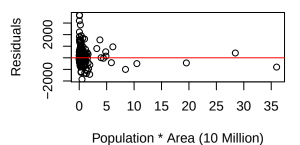


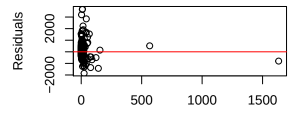
Model 1: Residuals ~ Income



Land Area Personal Income

Model 1: Residuals ~ Population * Are: Model 1: Residuals ~ Population * Incor

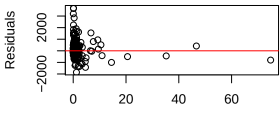




,

Population * Income (10 Million)

Model 1: Residuals ~ Area * Income



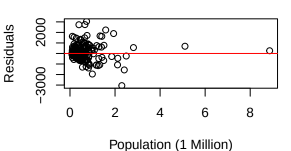
Area * Income (10 Million)

Model 2

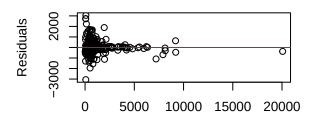
Model 2: Residuals ~ Fitted values

2000 Residuals -3000 15000 0 5000 Fitted Values

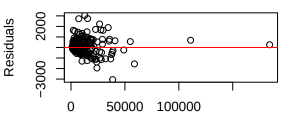
Model 2: Residuals ~ Population



Model 2: Residuals ~ Area

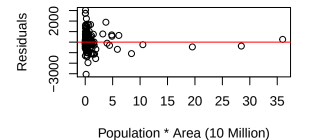


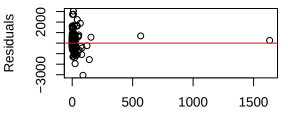
Model 2: Residuals ~ Income



Land Area

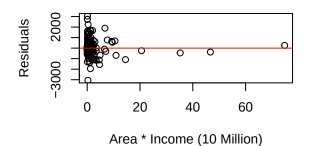
Personal Income Model 2: Residuals ~ Population * Are: Model 2: Residuals ~ Population * Incor





Population * Income (10 Million)

Model 1: Residuals ~ Area * Income



9

 \mathbf{f}

Model 1

$$Y_{1,2} = -84.914 + 0.001x_1 + -0.021x_2 + 0.089x_3 + -0.00000005x_1x_2$$
(1)

$$Y_{1,3} = -58.222 + 0.001x_1 + 0.093x_2 + -0.069x_3 + -0.00000001x_1x_3$$
(2)

$$Y_{2,3} = -94.964 + -0.021x_1 + 0.09x_2 + 0.001x_3 + -0.00000325x_2x_3$$
(3)

Model 2

$$Y_{1,2} = 133.388 + -0.476x_1 + -17.763x_2 + 0.128x_3 + 0.04428047x_1x_2 \tag{4}$$

$$Y_{1,3} = -252.911 + 0.192x_1 + 0.134x_2 + 6.371x_3 + -0.0000038x_1x_3$$
(5)

$$Y_{2,3} = -84.499 + -1.454x_1 + 0.113x_2 + 0.092x_3 + 0.00127165x_2x_3$$
(6)

 R^2 values of two factor interaction terms in Model 1

```
## (R2 of X1*X2) (R2 of X1*X3) (R2 of X2*X3)
## 0.9039154 0.9036285 0.9043730
```

 R^2 values of two factor interaction terms in Model 2

```
## (R2 of X1*X2) (R2 of X1*X3) (R2 of X2*X3)
## 0.9190910 0.9164615 0.9122407
```

For Model 1, the model with the $x_2 * x_3$ interaction term had the highest R^2 value of the other possible interaction models. Therefore, this is the preferable model, since a higher proportion of the variability in our response variable (number of active physicians) is explained by our predictors.

For Model 2, the model with the $x_1 * x_2$ interaction term had the highest R^2 value of the other possible interaction models. Therefore, this is the preferable model, since a higher proportion of the variability in our response variable (number of active physicians) is explained by our predictors.

Part 2

 \mathbf{a}

$$R_{Y,3|1,2}^2 = 0.0288$$

 $R_{Y,4|1,2}^2 = 0.0038$
 $R_{Y,5|1,2}^2 = 0.5538$

b

The coefficient of partial determination measures measures the proportionate reduction in variation of the response variable (number of active physicians) due to adding a new predictor $(x_3, x_4, \text{ or } x_5)$, given that x_1 (total population) and x_2 (total personal income) are already in the model. This means since the model including x_5 (number of hospital beds) had the higher proportion/coefficient of determination, 0.5538, this predictor variable is the best and most important to the model as it reduces the most variation in the number of active physicians being predicted. This higher proportion means that the sum of squares of the predictor being number of beds, is greater than the sum of squares for the other predictors, land area and proportion of population over 65, since the coefficient of partial determination is a ratio between the sum of squares of the predictor being considered and the predictors already in the model.

 \mathbf{c}

$$H_0: \beta_5 = 0$$

 $H_1: \beta_5 \neq 0$
 $F^* = 541.1801 > 6.6934$

reject H_0

Conclude that at the 1% significance level there is significant evidence that x_5 (number of hospital beds) is helpful in the regression model for predicting the response variable, number of active physicians, given that x_1 (total population) and x_2 (total personal income) are already in the model. Furthermore, we would not expect the F^* of the other 2 predictor variables, land area and percent of population older than 65, to be as high as the one for number of hospital beds. This is because the F^* value is calculated as the ratio between the sum of squares of the predictor in question given x_1 and x_2 are already in the model and the mean squared error of the whole model including the predictor in question. Since the sum of squares of the number of hospital beds is higher, this would mean this ratio and F^* value is greater and therefore given the null the outcome demonstrated by the data is more extreme as compared to the other possible predictor candidates.

 \mathbf{d}

$$\begin{split} R_{Y,3,4|1,2}^2 &= 0.0331 \\ R_{Y,3,5|1,2}^2 &= 0.5558 \\ R_{Y,4,5|1,2}^2 &= 0.5643 \end{split}$$

Based on these coefficient of partial determinations it is clear that the best pairing of predictors to use given that x_1 and x_2 are already in the model are x_4 (percent of seniors over 65) and x_5 (total number of hospital beds). By considering the pairing of these two predictors we can see that this pair reduces the greatest

proportion of variation (0.5643) in the total number active physicians compared to the other 2 possible pairings.

$$H_0: \beta_4 = \beta_5 = 0$$

$$H_1: \beta_4 \neq 0 \text{ or } \beta_5 \neq 0 \text{ or both } \neq 0$$

$$F^* = 122.7299 > 4.6543$$

reject H_0

Have significant evidence, that at the 1% significance level, the pairing of both or either of x_4 (proportion of population over 65) and x_5 (number of hospital beds) would be helpful and useful to our model given that x_1 (total population) and x_2 (total personal income) are already in the model.

Part 3

Appendix

```
knitr::opts chunk$set(echo = FALSE)
cdi_data = read.table("./CDI.txt")
phy = cdi_data$V8 # Number of Active Physicians
pop = cdi_data$V5 # Total Population
are = cdi_data$V4 # Land Area
inc = cdi_data$V16 # total personal income
den = pop / are # Population density
sen = cdi_data$V7
\# par(mfrow = c(1,2))
# stem(pop, width = 20)
# stem(are, width = 20)
# stem(inc, width = 20)
# stem(den, width = 20)
# stem(sen, width = 20)
stem(pop)
stem(are)
stem(inc)
stem(den)
stem(sen)
# Create Dataframes
model1 = data.frame(Y = phy, X1 = pop, X2 = are, X3 = inc)
colnames(model1) = c("Active Physicians (Y)", "Total Population (X1)", "Land Area (X2)", "Total Persona
model2 = data.frame(Y = phy, X1 = den, X2 = sen, X3 = inc)
colnames(model2) = c("Active Physicians (Y)", "Population Density (X1)", "Population Over 64 (X2)", "To
pairs(model1, main = "Scatter Plot Matrix (Model 1)")
cor(model1)
pairs(model2, main = "Scatter Plot Matrix (Model 2)")
cor(model2)
# Get models
fit1 = lm(model1)
fit2 = lm(model2)
f1s = summary(fit1)
f2s = summary(fit2)
betas1 = fit1$coefficients
betas2 = fit2$coefficients
slope = betas1[2] + betas1[3] + betas1[4]
int = betas1[1]
model1.R2 = f1s$r.squared
model2.R2 = f2s$r.squared
# Get residuals
m1.resid = f1s$residuals
m1.yhat = fit1$fitted.values
par(mfrow = c(2,2))
```

```
# Residual Plots
plot(m1.resid ~ m1.yhat,
     main = "Model 1: Residuals ~ Fitted values", xlab = "Fitted Values", ylab = "Residuals")
abline(0,0,col = "red")
# Predictors
plot(x = (pop)/1000000, y = m1.resid,
     main = "Model 1: Residuals ~ Population", xlab = "Population (1 Million)", ylab = "Residuals")
abline(0,0,col = "red")
plot(m1.resid ~ are,
     main = "Model 1: Residuals ~ Area", xlab = "Land Area", ylab = "Residuals")
abline(0,0,col = "red")
plot(m1.resid ~ inc,
     main = "Model 1: Residuals ~ Income", xlab = "Personal Income", ylab = "Residuals")
abline(0,0,col = "red")
par(mfrow = c(2,2))
# Two Factor
plot(x = (pop/100000) * (are/10000), y = m1.resid,
     main = "Model 1: Residuals ~ Population * Area", xlab = "Population * Area (10 Million)", ylab = "
abline(0,0,col = "red")
plot(x = (pop/100000) * (inc/10000), y = m1.resid,
     main = "Model 1: Residuals ~ Population * Income", xlab = "Population * Income (10 Million)", ylab
abline(0,0,col = "red")
plot(x = (are * inc)/10000000, y = m1.resid,
     main = "Model 1: Residuals ~ Area * Income", xlab = "Area * Income (10 Million)", ylab = "Residual
abline(0,0,col = "red")
# Get residuals
m2.resid = f2s$residuals
m2.yhat = fit2$fitted.values
par(mfrow = c(2,2))
# Residual Plots
plot(m2.resid ~ m2.yhat,
     main = "Model 2: Residuals ~ Fitted values", xlab = "Fitted Values", ylab = "Residuals")
abline(0,0,col = "red")
# Predictors
plot(x = (pop)/1000000, y = m2.resid,
     main = "Model 2: Residuals ~ Population", xlab = "Population (1 Million)", ylab = "Residuals")
abline(0,0,col = "red")
plot(m2.resid ~ are,
     main = "Model 2: Residuals ~ Area", xlab = "Land Area", ylab = "Residuals")
abline(0,0,col = "red")
```

```
plot(m2.resid ~ inc,
     main = "Model 2: Residuals ~ Income", xlab = "Personal Income", ylab = "Residuals")
abline(0,0,col = "red")
par(mfrow = c(2,2))
# Two Factor
plot(x = (pop/100000) * (are/10000), y = m2.resid,
     main = "Model 2: Residuals ~ Population * Area", xlab = "Population * Area (10 Million)", ylab = "
abline(0,0,col = "red")
plot(x = (pop/100000) * (inc/10000), y = m2.resid,
     main = "Model 2: Residuals ~ Population * Income", xlab = "Population * Income (10 Million)", ylab
abline(0,0,col = "red")
plot(x = (are * inc)/10000000, y = m2.resid,
     main = "Model 1: Residuals ~ Area * Income", xlab = "Area * Income (10 Million)", ylab = "Residual
abline(0,0,col = "red")
tf1_12 \leftarrow lm(phy \sim pop*are+inc)
tf1_13 \leftarrow lm(phy \sim pop*inc+are)
tf1_23 <- lm(phy ~ are*inc+pop)
r2_12 <- summary(tf1_12)$r.squared
r2_13 <- summary(tf1_13)$r.squared
r2_23 <- summary(tf1_23)$r.squared
res1 <- c(a = r2_{12}, b = r2_{13}, c = r2_{23})
names(res1) <- c("(R2 of X1*X2)", "(R2 of X1*X3)", "(R2 of X2*X3)")</pre>
tf2_{12} \leftarrow lm(phy \sim den*sen+inc)
tf2_13 \leftarrow lm(phy \sim den*inc+sen)
tf2_23 \leftarrow lm(phy \sim sen*inc+den)
r2_12 <- summary(tf2_12)$r.squared
r2_13 <- summary(tf2_13)$r.squared
r2_23 <- summary(tf2_23)$r.squared
res2 \leftarrow c(a = r2_{12}, b = r2_{13}, c = r2_{23})
names(res2) <- c("(R2 of X1*X2)", "(R2 of X1*X3)", "(R2 of X2*X3)")</pre>
tf1_b12 <- tf1_12$coefficients
tf1_b13 <- tf1_13$coefficients
tf1_b23 <- tf1_23$coefficients
tf2_b12 <- tf2_12$coefficients
tf2_b13 <- tf2_13$coefficients
tf2_b23 <- tf2_23$coefficients
res1
res2
# already have other predictors
bed <- cdi_data$V9</pre>
fit <- lm(phy \sim pop+inc)
fit3 <- lm(phy ~ pop+inc+are)</pre>
fit4 <- lm(phy ~ pop+inc+sen)
```

```
fit5 <- lm(phy ~ pop+inc+bed)</pre>
SSE <- function(fit) {</pre>
tail(anova(fit)$'Sum Sq', n = 1)
}
r2_3 = 1 - SSE(fit3)/SSE(fit)
r2_4 = 1 - SSE(fit4)/SSE(fit)
r2_5 = 1 - SSE(fit5)/SSE(fit)
# c(r2_3, r2_4, r2_5)
F5=anova(fit5)$Sum[3]/(SSE(fit5)/anova(fit5)$Df[4])
Fval=qf(1-0.01,1,436)
fit34=lm(phy~pop+inc+are+sen)
fit35=lm(phy~pop+inc+are+bed)
fit45=lm(phy~pop+inc+sen+bed)
r2_34=1-SSE(fit34)/SSE(fit)
r2_35=1-SSE(fit35)/SSE(fit)
r2_45=1-SSE(fit45)/SSE(fit)
SSE1245=SSE(fit45)
SSE12=SSE(fit)
SSR45=SSE12-SSE1245
F45=(SSR45/2)/(SSE12/435)
Fval45=qf(1-0.01,2,435)
```